Oxygen Insertion Reactions within the One-Dimensional Channels of Phases Related-to FeSb2O4


To cite this version:

Benjamin P. De Laune, Gregory J. Rees, Mariana J. Whitaker, Hien-Yoong Hah, Charles E. Johnson, et al.. Oxygen Insertion Reactions within the One-Dimensional Channels of Phases Related-to FeSb2O4. Inorganic Chemistry, American Chemical Society, 2017, 56 (1), pp.594-607. <10.1021/acs.inorgchem.6b02466>. <hal-01691857>

HAL Id: hal-01691857
https://hal.archives-ouvertes.fr/hal-01691857

Submitted on 24 Jan 2018

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Oxygen Insertion Reactions within the One-Dimensional Channels of Phases Related to FeSb$_2$O$_4$


ABSTRACT: The structure of the mineral schafarzikite, FeSb$_2$O$_4$, has one-dimensional channels with walls comprising Sb$^{3+}$ cations; the channels are separated by edge-linked FeO$_6$ octahedra that form infinite chains parallel to the channels. Although this structure provides interest with respect to the magnetic and electrical properties associated with the chains and the possibility of chemistry that could occur within the channels, materials in this structural class have received very little attention. Here we show, for the first time, that heating selected phases in oxygen-rich atmospheres can result in relatively large oxygen uptakes (up to ~2% by mass) at low temperatures (ca. 350 °C) while retaining the parent structure. Using a variety of structural and spectroscopic techniques, it is shown that oxygen is inserted into the channels to provide a structure with the potential to show high one-dimensional oxide ion conductivity. This is the first report of oxygen-excess phases derived from this structure. The oxygen insertion is accompanied not only by oxidation of Fe$^{2+}$ to Fe$^{3+}$ within the octahedral chains but also Sb$^{3+}$ to Sb$^{5+}$ within the channel walls. The formation of a defect cluster comprising one 5-coordinate Sb$^{5+}$ ion (which is very rare in an oxide environment), two interstitial O$_2^-$ ions, and two 4-coordinate Sb$^{3+}$ ions is suggested and is consistent with all experimental observations. To the best of our knowledge, this is the first example of an oxidation process where the local energetics of the product dictate that simultaneous oxidation of two different cations must occur. This reaction, together with a wide range of cation substitutions that are possible on the transition metal sites, presents opportunities to explore the schafarzikite structure more extensively for a range of catalytic and electrocatalytic applications.

1. INTRODUCTION

The tetragonal mineral schafarzikite (FeSb$_2$O$_4$; P4$_2$/mnb c = 5.91 Å$^1$) is of significant interest because of its unusual one-dimensional (1-D) structural characteristics and its ability to display a high degree of compositional flexibility. The structure is related to that of rutile, TiO$_2$, but neighboring chains of edge-shared transition metal octahedra, MO$_6$, along [001] are now separated: in contrast to the rutile structure, the chains do not share anions with adjacent chains but are connected through O−Sb−O linkages (Figure 1). In FeSb$_2$O$_4$, the chains of octahedra contain Fe$^{2+}$ (A-site), and Sb$^{3+}$ (B-site) provides the link. The B-site has trigonal pyramidal coordination, SbO$_3$, which can also be regarded as pseudo-tetrahedral, SbO$_3$E, by including the stereochemically active lone pair of electrons (E). These electron pairs appear crucial for stabilizing the second key structural feature: channels that exist along [001], parallel to the chains of octahedra. Sb$^{3+}$ ions form the channel walls and their lone pairs, E, are directed into the channels.

The structure can accommodate a range of cations that exist along [001], parallel to the chains of octahedra. Sb$^{3+}$ ions form the channel walls and their lone pairs, E, are directed into the channels.
may result in simultaneous or preferential oxidation of Sb$^{3+}$, resulting in mixed phase products, we have recently achieved complete success for $M = \text{Fe}$, where controlled oxidation of Fe$^{2+}$ to Fe$^{3+}$ has been achieved.\textsuperscript{13}

Examination of the physical properties of materials related to FeSb$_2$O$_4$ has, to date, been largely limited to the magnetic properties of phases containing transition metal ions,\textsuperscript{7,12-15} for which the link between composition, structure, and magnetic order is now well understood.\textsuperscript{14} A complex interplay between three magnetic exchange pathways determines the magnetic ground state: intrachain direct M–M exchange, 90° super-exchange along a chain, and a weaker but significant interchain M–O–Sb–O–M exchange. All systems are antiferromagnetic except for CuAs$_2$O$_4$ which is ferromagnetic at low temperatures.\textsuperscript{10} The chemical reactivity of the materials has received even less attention. This is surprising since the structural channels display a feature which, to the best of our knowledge, is unique: cations (Sb$^{3+}$ ions) form the channel walls, whereas channels in oxide materials normally have walls of O$^{2-}$ ions such that the cavities are suitable hosts for cationic species. Channel walls in aluminosilicates, for example, are O$^{2-}$ ions from Al/SiO$_4$ tetrahedra. In schafarzikites, therefore, we could anticipate the chemical insertion of anions rather than cations, and here we demonstrate the first example of such a reaction where a unique oxidation mechanism occurs for phases with A-sites containing some Fe$^{2+}$ ions and B-sites containing some Sb$^{3+}$ ions. Oxygen insertion into the channels occurs at low temperatures (for solid state reactions) to provide new metastable phases in which simultaneous oxidation of both Fe$^{2+}$ and Sb$^{3+}$ has occurred; the proposed mechanism provides suitable chemical environments for both the Sb$^{3+}$ and Sb$^{5+}$ cations that bond to the interstitial O$^{2-}$ ions.

The low-temperature oxidation mechanism discussed here is thought to be unique and suggests high oxygen and/or oxide ion mobility within the channels. It is pertinent to note that low-dimensional structures containing partially filled oxide ion sites have recently gained interest for energy materials since they can provide an interstitial mechanism for oxide ion migration rather than the normal vacancy driven mechanism that occurs, for example, in the fluorite solid electrolyte YSZ, Zr$_{1−x}$Y$_x$O$_{2+δ}$.\textsuperscript{17} This interest also extends to cathode materials where mixed conductivity is required. A recent example is the study of the oxygen excess, mixed conducting, Ruddlesden–Popper phases (La,Pr)$_2$(Ni,Co)O$_{4+δ}$\textsuperscript{18} where the excess oxygen enters the structure between two adjacent rocksalt-related layers and can coordinate tetrahedrally to lanthanide ions. In this respect, materials with channels containing highly mobile interstitial oxide ions would be of special interest. Given that a range of transition metals can occupy the chains of octahedra, the focus of this study is therefore also highly relevant to the search for new catalysts and electrocatalysts.

2. EXPERIMENTAL SECTION

A variety of compounds with the schafarzikite structure were synthesized by heating intimately ground stoichiometric mixtures of dried metal oxides and metals in evacuated, sealed quartz tubes at 600–750 °C for 6 h periods with intermittent grinding. Alumina inserts were used to prevent reaction with the quartz. Critical to the results presented in this paper is the amount of Fe$^{2+}$ per formula unit (ppm); this was controlled by either isovalent cation substitutions at the A-site or aliovalent substitution of Pb$^{2+}$ for Sb$^{3+}$ at the B-site, and compositions with particular relevance to this paper are given in Table 1. The compositions reported here were chosen to provide important control of the Fe$^{2+}$ content (given in Table 1); this was kept quite low to avoid large oxygen absorption, which was found to result in broadened diffraction peaks. Both Mg$^{2+}$ and Co$^{2+}$ substitutions for Fe$^{2+}$ were used to provide substantial differences to electronic conductivity within the chains of octahedra. The partial substitution of Pb$^{2+}$ for Sb$^{3+}$ not only reduced the Fe$^{2+}$ content but also enhanced the high temperature stability which was important for revealing that the oxidation involved two discrete steps. Oxidised phases were created by heating samples in air or oxygen. For $^{17}$O NMR analysis, samples of MgSb$_2$O$_4$ and Sb$_2$O$_4$ were exchanged with $^{17}$O-enriched H$_2$O hydrothermally, and FeSb$_2$O$_4$-related phases were exchanged with $^{17}$O-enriched oxygen gas (75% enrichment) at 350 °C for times of 12–66 min depending on the observed rate of oxygen absorption. Full experimental details are provided in Supporting Information.

3. RESULTS

3.1. Thermogravimetric Analysis (TGA).

Mass changes for three representative samples heated in oxygen gas are shown in Figure 2. All are unstable above 600 °C and form

![Figure 1. Schafarzikite structure viewed (a) approximately along [001] showing the empty channels and (b) along [110]. Elements are represented as colored spheres: Fe$^{2+}$ (A site; blue), Sb$^{3+}$ (B site; black), and O$^{2-}$ in the O2, equatorial, (red) and O1, apical, (yellow) positions. FeO$_6$ octahedra are gray.](https://example.com/Figure1.png)

![Figure 2. TGA plots for CoSb$_2$O$_4$ (green), Mg$_{0.25}$Fe$_{0.75}$Sb$_2$O$_4$ (brown), Co$_{0.25}$Fe$_{0.75}$Sb$_{1.75}$Pb$_{0.25}$O$_4$ (purple) heated in an O$_2$ atmosphere at 10 °C min$^{-1}$ to 800 °C.](https://example.com/Figure2.png)

Table 1. Important Phase Compositions and Their Fe$^{2+}$ Content

<table>
<thead>
<tr>
<th>composition</th>
<th>amount of Fe$^{2+}$ per formula unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co$<em>{0.25}$Fe$</em>{0.75}$Sb$_2$O$_4$</td>
<td>0.25</td>
</tr>
<tr>
<td>Mg$<em>{0.25}$Fe$</em>{0.75}$Sb$_2$O$_4$</td>
<td>0.25</td>
</tr>
<tr>
<td>FeSb$<em>{1.25}$Pb$</em>{0.25}$O$_4$</td>
<td>0.25Fe$^{2+}$ (+0.75Fe$^{3+}$)</td>
</tr>
<tr>
<td>Co$<em>{0.25}$Fe$</em>{0.75}$Sb$<em>{1.75}$Pb$</em>{0.25}$O$_4$</td>
<td>0.50Fe$^{2+}$ (+0.25Fe$^{3+}$)</td>
</tr>
</tbody>
</table>
Sb\textsuperscript{5+}-containing products; CoSb\textsubscript{2}O\textsubscript{6} for example, simply forms CoSb\textsubscript{2}O\textsubscript{8} with a mass increase of ca. 8% (expected 8.7%). However, a low-temperature plateau is also observed centered at \(\sim 400\) °C for all samples containing Fe\textsuperscript{2+} ions. Indeed, the sample Co\textsubscript{0.25}Fe\textsubscript{0.75}Sb\textsubscript{1.75}Pb\textsubscript{0.25}O\textsubscript{4} shows strong evidence that an additional process occurs at higher temperatures to provide a second plateau centered at \(\sim 550\) °C. It should be emphasized that the low-temperature plateau is not observed for CoSb\textsubscript{2}O\textsubscript{4} which also shows that the oxygen uptake at low Fe\textsuperscript{2+} contents is related to the amount of Fe\textsuperscript{2+} in the sample (see Figure 3). It is assumed that for all materials investigated, the substitution of Pb\textsuperscript{2+} for Sb\textsuperscript{3+} is balanced only by oxidation of Fe\textsuperscript{2+} to Fe\textsuperscript{3+}. However, a critical feature for all samples exhibiting the low temperature plateau is that the amount of oxygen absorbed is, in fact, significantly higher than that expected for oxidation of all Fe\textsuperscript{2+} to Fe\textsuperscript{3+}. This is clear in Figure 3 which also shows that the oxygen uptake at low Fe\textsuperscript{2+} contents (up to 0.50Fe\textsuperscript{2+} pfu) is, in fact, approximately twice that expected. Two possibilities were considered as plausible explanations: either other cations, in addition to Fe\textsuperscript{2+}, are oxidized (or higher oxidation states of Fe are formed), or oxygen enters as peroxide ions, O\textsuperscript{2−}, rather than O\textsuperscript{3−} ions. A structural study was therefore conducted to determine the precise nature of the structural changes that occur during oxidation.

3.2. Structure of FeSb\textsubscript{1.25}Pb\textsubscript{0.75}O\textsubscript{4+x}. Powder X-ray diffraction (PXRD) analysis of samples obtained by cooling from the low temperature plateaus on the TGA plots revealed that the basic structure had been retained (space group \textit{P4}_\text{i}/ \textit{mnc}) but with changes to the unit cell dimensions: a reduction in \(a\) and increase in \(c\). This is similar to that observed when Pb\textsuperscript{2+} is substituted for Sb\textsuperscript{3+} in FeSb\textsubscript{2}O\textsubscript{4} where oxidation of A-site cations within the chains of octahedra results in increased intrachain Fe–Fe repulsions and consequently an expansion of \(c\). For samples with large Fe\textsuperscript{2+} contents, e.g., FeSb\textsubscript{2}O\textsubscript{4}, the large amount of oxygen incorporation resulted in a degree of structure degradation; structure determination was therefore focused on the oxidation product of FeSb\textsubscript{1.25}Pb\textsubscript{0.75}O\textsubscript{4}, which initially contained 25% Fe\textsuperscript{2+} and 75% Fe\textsuperscript{3+}. Figure 3 shows that a significant oxygen uptake occurs for this composition, and a sample for NPD study (\(\sim 4\) g) was prepared by heating in air at 400 °C for 12 min. TGA suggested that the final composition was FeSb\textsubscript{1.42}Pb\textsubscript{0.58}O\textsubscript{4.29} compared to FeSb\textsubscript{1.25}Pb\textsubscript{0.75}O\textsubscript{4.125}, which is predicted if oxygen insertion as O\textsuperscript{2−} ions is balanced only by oxidation of Fe\textsuperscript{2+} to Fe\textsuperscript{3+}.

NPD data (HRPT at PSI, 300 K, \(\lambda = 1.8852\) Å) were collected on FeSb\textsubscript{1.25}Pb\textsubscript{0.75}O\textsubscript{4.29} at ambient temperature. Because of the relatively small change in composition, structure refinement based on the structure of the stoichiometric, unoxidized material gave a good fit to the data. However, difference Fourier maps suggested additional scattering density within the channels, as shown in Figure 4; inclusion of an oxide ion into the refinement at the suggested position resulted in a small, but statistically significant, improvement to the fit (\(\chi^2\) decreased from 2.6 to 2.2). The profiles for the refinement are given in Figure 5, and the refined structural parameters are given in Table 2 where O3 represents the incorporated interstitial oxygen, O1 the apical and O2 the equatorial positions (see Figure 1). The disorder, from both the mixed Sb/Pb on the 8h-site and the interstitial oxygen, resulted in

![Figure 3. Increase in oxygen content per formula unit (pfu) as a function of initial Fe\textsuperscript{2+} content pfu from TGA data for the three series indicated. The results are compared to those expected assuming only oxidation of Fe\textsuperscript{2+} to Fe\textsuperscript{3+}.](image)

![Figure 4. Difference Fourier map for FeSb\textsubscript{1.25}Pb\textsubscript{0.75}O\textsubscript{4} showing additional neutron scattering near the center of the channel running along [001] and located at (0, 0, z); see Figure 1. The position corresponding to the O3 site of Table 2 is marked.](image)

![Figure 5. Rietveld refinement profiles for FeSb\textsubscript{1.25}Pb\textsubscript{0.75}O\textsubscript{4} showing raw data (red crosses), fitted profile (green) and difference profile (mauve). Reflection positions are marked as vertical bars.](image)
Table 2. Structural Parameters for FeSb1.25Pb0.75O4.24 from NPD Data

<table>
<thead>
<tr>
<th>atom</th>
<th>position</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>100 × Uiso (Å²)</th>
<th>occupancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>4d</td>
<td>0.00</td>
<td>0.5</td>
<td>0.25</td>
<td>2.19(6)</td>
<td>1.0</td>
</tr>
<tr>
<td>Sb/Pb</td>
<td>8h</td>
<td>0.1542(3)</td>
<td>0.1557(3)</td>
<td>0.0</td>
<td>2.82(6)</td>
<td>0.625/0.375</td>
</tr>
<tr>
<td>O1</td>
<td>8g</td>
<td>0.6701(2)</td>
<td>0.1701(2)</td>
<td>0.25</td>
<td>3.90*</td>
<td>1.0</td>
</tr>
<tr>
<td>O2</td>
<td>8h</td>
<td>0.0981(3)</td>
<td>0.6278(3)</td>
<td>0.0</td>
<td>2.64(8)</td>
<td>1.0</td>
</tr>
<tr>
<td>O3</td>
<td>16i</td>
<td>0.540(5)</td>
<td>0.417(3)</td>
<td>0.254(7)</td>
<td>3.1(1)</td>
<td>0.059(4)</td>
</tr>
</tbody>
</table>

*Anisotropic parameters (all × 100/Å²): U11 4.5(1), U22 4.5(1), U33 2.7(2); U12 = -2.4(1). U13 = -0.8(1) U12 = -0.8(1). P4/mmbc; a = 8.4063(3) Å, b = 8.4063(3) Å, c = 6.0443(3) Å. χ² = 2.24; Rwp = 0.051.

Table 3. Selected Bond Distances and Bond Valence Sums (Using the r₀ Parameter for Fe³⁺) for FeSb1.25Pb0.75O4.24

<table>
<thead>
<tr>
<th>bond distances/Å</th>
<th>BVS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe−O1</td>
<td>2.023(3) [× 2]</td>
</tr>
<tr>
<td>Fe−O2</td>
<td>2.029(2) [× 4]</td>
</tr>
<tr>
<td>Sb/Pb−O1</td>
<td>2.108(2)</td>
</tr>
<tr>
<td>Sb/Pb−O2</td>
<td>2.095(4) [× 2]</td>
</tr>
<tr>
<td>Sb/Pb−O3</td>
<td>1.91(4), 2.30(5)</td>
</tr>
<tr>
<td>O3−O3</td>
<td>1.09(3), 1.54(5)</td>
</tr>
</tbody>
</table>

(BVS)³⁺ are given in Table 3. The shortest O3−O3 distances are 1.09(3) Å and 1.54(5) Å, the latter being across the center of the channel as shown in Figure 6. Although these distances are too short for fully occupied sites, it should be noted that only 6% of the 16i-sites are actually occupied, so the model is entirely satisfactory for such an average structure. Table 3 gives the BVS for Fe (2.91) based on the r₀ parameter for Fe³⁺, 1.759.¹⁰ This methodology has previously been found to yield excellent agreement between the Fe BVS and the Fe oxidation state for FeSb₁₋ₓPbₓO₄ with x > 0.4.¹³ For the Fe oxidation states of 2.45+, 2.50+, and 2.70+, the bond distances in ref 13 give BVS values of 2.54, 2.54, and 2.64. The BVS of 2.91 (Table 3) therefore provides strong evidence that the oxygen incorporation has resulted in complete oxidation of the Fe²⁺ present to Fe³⁺. However, the increase in oxygen content determined by the O3 occupancy is approximately twice that expected for a model in which insertion of O₂⁻ ions is accompanied by oxidation of all Fe²⁺ to Fe³⁺. The O3−O3 separation of 1.54 Å across the center of the channel is therefore of particular interest since it is similar to the O−O separation in simple peroxides, e.g., 1.49 Å in Na₂O₂.²⁰ The existence of peroxide (or even superoxide) links within the structure could explain why the observed mass increase is higher than that expected for simple oxidation of Fe²⁺ to Fe³⁺.

A possible O−O linkage is marked in Figure 6, where the O3 species are bonded to Sb/Pb at a distance of 1.91(4) Å (Table 3).

3.3. Structural Changes during Oxidation. In order to obtain more detailed information on the structural changes occurring during the oxidation, and hence the mechanism involved, an in situ NPD study was conducted (D20 at ILL, Grenoble, λ = 1.5430 Å). The sample chosen for investigation was Co₀.₂₅Fe₀.₇₅Sb₁.₇₅Pb₀.₂₅O₄ since it has a high final decomposition temperature and a double plateau at low temperatures (Figure 2). Essential experimental details are given in Supporting Information for the oxidation under flowing oxygen with structural data collected up to 500 °C. Initially, data were collected for 5 min (5 °C) intervals up to 350 °C, conditions which were expected to yield a composition on the first low temperature plateau. Data were then collected for several extended periods (15 min) at 350 °C before temperature ramping was continued up to 500 °C. To minimize the effects of correlation between the O3 site occupancy and its isotropic displacement parameter (IDP), the IDPs for O1, O2, and O3 were constrained to be equal; in addition, Fe/Co site occupancy was constrained to 0.75/0.25, allowing oxygen with structural data collected up to 500 °C. Although these distances are too short for fully occupied sites, it should be noted that only 6% of the 16i-sites are actually occupied, so the model is entirely satisfactory for such an average structure. Table 3 gives the BVS for Fe (2.91) based on the r₀ parameter for Fe³⁺, 1.759.¹⁰ This methodology has previously been found to yield excellent agreement between the Fe BVS and the Fe oxidation state for FeSb₁₋ₓPbₓO₄ with x > 0.4.¹³ For the Fe oxidation states of 2.45+, 2.50+, and 2.70+, the bond distances in ref 13 give BVS values of 2.54, 2.54, and 2.64. The BVS of 2.91 (Table 3) therefore provides strong evidence that the oxygen incorporation has resulted in complete oxidation of the Fe²⁺ present to Fe³⁺. However, the increase in oxygen content determined by the O3 occupancy is approximately twice that expected for a model in which insertion of O₂⁻ ions is accompanied by oxidation of all Fe²⁺ to Fe³⁺. The O3−O3 separation of 1.54 Å across the center of the channel is therefore of particular interest since it is similar to the O−O separation in simple peroxides, e.g., 1.49 Å in Na₂O₂.²⁰ The existence of peroxide (or even superoxide) links within the structure could explain why the observed mass increase is higher than that expected for simple oxidation of Fe²⁺ to Fe³⁺.

A possible O−O linkage is marked in Figure 6, where the O3 species are bonded to Sb/Pb at a distance of 1.91(4) Å (Table 3).
between 350 and 500 °C, but the slope for the uptake is reduced in this higher temperature region. It is important to note that all structural changes recorded in Figure 8 show a distinct discontinuity at approximately 350 °C, which suggests a change in the oxidation mechanism at about this temperature. This was confirmed by other techniques, and details are presented later (sections 3.6 and 4).

Figure 8b,c shows changes to the unit cell parameters during the main oxidation (200 °C < T < 350 °C): a decreases, c increases, and the volume decreases. This behavior is very similar to that observed for increasing x in FeSb2−xPbO4 where the increase in c was attributed to enhanced cation–cation repulsion within the chains of octahedra resulting from the oxidation of Fe2+ to Fe3+.13 For T > 350 °C, some oxidation continues (Figure 8a), but the main changes to the unit cell appear to be thermal in nature and the volume increases with temperature. These data support the importance of oxidation of Fe2+ to Fe3+ at temperatures up to 350 °C. Confirmation is provided by the decrease in the average Fe/Co−O bond distance within the octahedra, Figure 8d. Simple bond valence sum calculations,19 based on oxidation of all the Fe2+ present to Fe3+, show that the changes observed are compatible with such a process. Since no low-temperature oxidation of CoSb2O4 occurs, Figure 2, we conclude that facile oxidation of Co3+ does not occur.

Figure 8e,f shows data with direct relevance to oxygen entering the channels. The change in distance between two Sb/Pb cations immediately across a given channel is seen in Figure 8e and shows a substantial decrease of 7% (from 4.00 to 3.73 Å); this is largely responsible for the contraction in the unit cell a parameter (Figure 8b). The decrease in channel diameter is also consistent with the insertion of negative O2− ions which can bond to the positive ions that form the channel walls, in this case Sb3+ and Pb2+. Since oxidation of Pb-free schafarzikites display the same oxidation characteristics, lead cannot be actively involved in the oxidation mechanism, and the formation of Sb–O bonds must occur. This bond formation is consistent with the change in the average bond length for the three Sb/Pb–O1/O2 bonds, Figure 8f. In marked contrast to the Fe/Co–O distances (Figure 8d), these bonds increase with the level of oxidation and cannot simply relate to oxidation of Sb3+ to Sb5+. Although BVS estimates are not straightforward because of the partial substitution of Pb on the Sb sites, Figure 8f shows the predicted average bond distances derived from BVS calculations for this site when coordinated to both three and four oxygens. The calculations are weighted to include the contribution from Pb2+ ions. The figure supports the hypothesis that oxygen enters the channels and bonds to Sb to form 4-coordinate Sb3+. However, it must be remembered that not all Sb3+ ions will be bonded to an extra O2− ion, so the apparent quantitative agreement of the data in Figure 8f is partly coincidental.

The in situ structural examination gives significant information concerning the nature of the oxidation of Co0.25Fe0.75Sb1.75Pb0.25O4 and hence other related phases. It is clear that oxidation of cations within the chains of octahedra occurs, but the level of interstitial oxygen is higher than this mechanism can support; in this respect, the data endorse the results from TGA analysis. At this stage, two mechanisms...
remained plausible to explain the enhanced oxygen content for the materials oxidized without phase decomposition:

(i) Peroxide links occur across the channels, e.g., Sb–O–O–Sb as shown in Figure 6;
(ii) Another oxidation process accompanies the oxidation of Fe$^{2+}$ to Fe$^{3+}$.

In order to resolve this uncertainty, additional information was sought from Raman spectroscopy, neutron PDF analysis, Mössbauer spectroscopy, NMR spectroscopy, and X-ray photoelectron spectroscopy (XPS); the results are discussed below and in Supporting Information.

3.4. Raman Spectroscopy. The best method to probe the existence of O–O–O bonds is Raman spectroscopy for which the O–O vibration is characterized by a band in the Raman shift range 750–850 cm$^{-1}$.

The parent material, FeSb$_2$O$_4$, has previously been studied, and the strongest band at 668 cm$^{-1}$ was assigned to a Sb–O vibration with symmetry $A_{2g}$. We have examined a range of materials oxidized by heating in oxygen as previously described (vide supra) but have not found any evidence for a vibrational band to support the presence of peroxide (or superoxide). Figure 9, for example, shows spectra for Co$_{0.75}$Fe$_{0.25}$Sb$_2$O$_4$ before and after oxidation. The spectrum of the initial material agrees well with that previously described for FeSb$_2$O$_4$, whereas the oxidized material has broadened bands which often show shoulders. The principal Sb–O vibration shows such features which are to be expected because of the change in coordination of some of the Sb$^{3+}$ ions following oxidation. However, no band in the O$_2^-$ region (or at

Figure 8. Variation with temperature of various structural parameters for Co$_{0.75}$Fe$_{0.25}$Sb$_{1.75}$Pb$_{0.25}$O$_4$ on heating in oxygen: (a) O3 site occupancy; (b) unit cell parameters; (c) unit cell volume; (d) average bond distance within the MO$_6$ octahedra showing values predicted for oxidation of Fe$^{2+}$ to Fe$^{3+}$; (e) cation separation across the channel (Sb/Pb–Sb/Pb); (f) average Sb/Pb–O (excluding interstitial O3). Error bars are included ($\pm$1esd) for all plots.
higher shifts compatible with O$_2^-$ is seen which precludes the existence of such species in the oxidized compounds.

3.5. Mössbauer Spectroscopy. Since no low-temperature oxidation of CoSb$_2$O$_4$ occurs, Figure 2, Co$^{2+}$ oxidation cannot occur under the low temperature oxidation conditions employed. The preferential oxidation of Sb$^{3+}$ rather than Co$^{2+}$ is supported by the formation of the trirutile CoSb$_2$O$_6$ (containing Co$^{2+}$ and Sb$^{5+}$).$^{23}$ Similarly, the synthesis of FePbBiO$_4$ (containing Fe$^{3+}$ and Pb$^{2+}$ ions)$^{24}$ and Pb$_2$Sb$_2$O$_7$ (containing Sb$^{5+}$ and Pb$^{2+}$ ions)$^{25}$ under oxidizing conditions illustrates that Pb$^{2+}$ ions are thermodynamically stable in the presence of Fe$^{3+}$ and Sb$^{5+}$ ions and will also remain unoxidized in the reactions under current discussion. The formation of Fe$^{4+}$ ions in the presence of Sb$^{3+}$ was considered highly unlikely, and the most plausible mechanism to explain the experimental obervations was one involving the simultaneous oxidation of Fe$^{2+}$ (to Fe$^{3+}$) and Sb$^{3+}$ (to Sb$^{5+}$) during the low temperature oxidation reaction. Since Mössbauer spectroscopy is the most powerful probe to determine Fe and Sb oxidation states,$^{57}$ Fe and $^{121}$Sb Mössbauer spectra were collected to test this hypothesis. Although data were collected on a variety of materials, here we focus only on Co$_{0.25}$Fe$_{0.75}$Sb$_{1.75}$Pb$_{0.25}$O$_4$ and its oxidized product for which we are able to compare results with those from the detailed in situ NPD data.

Figures 10 and 11 show $^{57}$Fe and $^{121}$Sb Mössbauer spectra for the initial compound and the sample after oxidation in air at 350 $°$C for 3 h, respectively. It can be seen that the absorption peaks from Sb$^{3+}$ are suggestive of more than one component, which may relate to various factors including quadrupolar effects and the presence of more than one Sb$^{3+}$ site postoxidation (vide infra). These are under investigation and will be reported later; for the present analysis, since the Fe and Sb oxidation states are of primary concern, the Sb$^{3+}$ absorption peaks have been fitted to a single site, but quadrupole splitting effects have been included. Table 4 gives the detailed Mössbauer parameters. The $^{57}$Fe spectrum for the initial material, Figure 10a, suggests 58% Fe$^{2+}$ and 42% Fe$^{3+}$ based on the respective peak areas (Table 4). Correction of these ambient temperature data for Fe$^{2+}$ and Fe$^{3+}$ Debye–Waller factor effects (using thermal factors for FeSb$_2$O$_4$ $^{13}$ and FePbBiO$_4$ $^{26}$) results in the composition 62% Fe$^{2+}$ and 38% Fe$^{3+}$, in reasonable agreement with the values expected for an initial composition of 0.50 Fe$^{2+}$ and 0.25 Fe$^{3+}$pfu (67% Fe$^{2+}$ and 33% Fe$^{3+}$).

Full oxidation of Fe$^{2+}$ to Fe$^{3+}$ is indicated in the product of oxidation at 350 $°$C, Figure 11a, but no evidence for higher oxidation states of Fe has been observed; the formation of Fe$^{4+}$ was therefore eliminated from the oxidation mechanism. The partial oxidation of Sb$^{3+}$ to Sb$^{5+}$ indicated in the $^{121}$Sb spectrum in Figure 11b is therefore of major significance. The peak areas in the $^{121}$Sb Mössbauer data recorded at 90 K, Table 4, are necessary to provide a reliable estimate for the Sb$^{5+}$ concentration (18(5)%). The difference between this value and that deduced from the peak areas in the ambient temperature data (22(5)%) is expected because of the significantly higher Debye–Waller factor for the lower valent Sb$^{3+}$ ion (compared with Sb$^{5+}$) and the resulting lower recoil-free fraction at 300 K for Sb$^{3+}$ compared with Sb$^{5+}$. At 90 K, the recoil-free fractions for both oxidation states are similar. For 18(5)% conversion of Sb$^{3+}$ to Sb$^{5+}$ (Table 4) and all Fe being present as Fe$^{3+}$, the final composition would be Co$_{0.25}$Fe$_{0.75}$Sb$_{1.75}$Pb$_{0.25}$O$_{4.57(9)}$, which is in excellent agreement with the NPD data in Figure 8a which suggests an oxygen content of 4.50 pfu. The critical conclusion is that the oxidation process at 350 $°$C does indeed relate to the oxidation of both Fe$^{2+}$ to Fe$^{3+}$ and Sb$^{3+}$ to Sb$^{5+}$. We note that two Fe$^{2+}$ ions (0.50 pfu) are oxidized for one Sb$^{3+}$ (14.3% Sb corresponds to 0.25 Sb pfu), and this is discussed in detail later.
oxidation processes relate to synchronous oxidation of Fe\(^{2+}\) to Fe\(^{3+}\) and subsequent further oxidation of Sb\(^{3+}\) to Sb\(^{5+}\) at higher temperatures.

3.6. \(^{17}\)O NMR Spectroscopy. In order to obtain additional information concerning the nature of oxygen absorption, especially with respect to possible peroxide formation, \(^{17}\)O NMR spectra were recorded for Fe\(^{2+}\)-containing samples after oxidation in \(^{17}\)O-enriched oxygen gas. To assist the interpretation of the spectra, the \(^{17}\)O and \(^{25}\)Mg MAS NMR data of the diamagnetic MgSb\(_2\)O\(_4\) and Sb\(_2\)O\(_3\) systems were also obtained and are shown in Figure 13; the measured NMR parameters and the associated GIPAW DFT calculated parameters are summarized in Table 5. The variable \(B_0\) field \(^{17}\)O MAS NMR data of Figure 13a shows the presence of two resonances associated with the apical and equatorial positions in the MgSb\(_2\)O\(_4\) lattice; these are denoted as sites 1 and 2, respectively, and correspond to O1 and O2 in Figure 1 and Table 3. In addition, channel H\(_2\)O and surface OH resonances associated with the apical and equatorial positions are observed. The presence of all four O species is corroborated by the \(^{17}\)O 2D 3QMAS data of Figure 1b. A comparison of the \(\delta_{iso}\) and \(P_Q\) values elucidated from the variable \(B_0\) field analysis of the data characterizing sites 1 and 2 (see Figure 13c) with the corresponding calculated GIPAW DFT values demonstrates that only one of the O positions exhibits a good agreement between measurement and calculation. Furthermore, the \(^{25}\)Mg \(\delta_{iso}\) and \(P_Q\) parameters measured from the data of Figure 13d are in excellent agreement with the GIPAW DFT calculated values, especially after geometry optimization. Hence, it can be inferred that the residual H\(_2\)O and OH\(^-\) speciation in the channels selectively influence only one of the framework O positions. Similar evidence of this selective influence emanates from the \(^{17}\)O MAS NMR data from Sb\(_2\)O\(_3\) shown in Figure 13e; although the apical and equatorial O positions of MgSb\(_2\)O\(_4\) are comparable, possessing similar arrangements with the Sb positions, only the apical site 1 exhibits \(\delta_{iso}\) and \(P_Q\) values.
that are in agreement with the Sb₂O₅ unit, thus suggesting that the interaction with channel H₂O and surface OH⁻ species influences the equatorial site 2 represented by the upfield resonance.

Solid state ¹⁷O NMR studies on paramagnetic materials are uncommon due to O typically being directly bonded (or in close spatial proximity) to paramagnetic species.³⁰ The ¹⁷O MAS NMR data of the paramagnetic schafarzikite-related systems shown in Figure 14 are hyperfine shifted, presumably by contributions from both the pseudocontact shift and the Fermi contact interactions. These spectra exhibit broad characteristic anisotropies which are represented as manifolds of spinning sidebands that are usually associated with the pseudocontact shift contribution. Through the implementation of variable MAS frequencies the isotropic position of each sideband manifold has been determined. It is proposed that the narrow ¹⁷O resonances at δiso ~ 2150 and ~1850 ppm represent the high symmetry axial and equatorial oxygen environments of the framework which corroborates the NPD results. A substantial amount of labeled O is located in the channels and is responsible for the broad underlying component (span Δδ ≈ 0–4,000 ppm) that resides underneath the better defined spinning sideband manifolds (see Figure 14b–e). This observation is also supported by the NPD refinement, with the broad nature of this resonance indicating a clear lack of order in these channel O environments. A direct comparison of the ¹⁷O MAS NMR data measured from diamagnetic MgSb₂O₄ (see Figure 14a) with the paramagnetic schafarzikite-related systems shown in Figure 14b–e indicates that the magnitude of the paramagnetic shifts and anisotropies are dependent on the nominal composition and the relative amounts of the paramagnetic Co²⁺, Fe²⁺, and Fe³⁺ defining each system.

The NMR data therefore show significant ¹⁷O substitution at the lattice O1 (apical) and O2 (equatorial) sites following short oxygen incorporation reactions. This indicates that oxide ion diffusion is occurring within the structural framework in addition to that within the channels. The broad signals from the channel sites prevent definitive characterization of their environment, but do not support the presence of peroxide bond formation.

### 3.7. Electronic Conductivity

Conductivity data were collected on pellets of Coₐ₋₀.₅₀Fe₀.₅₀Sb₂O₄ under three conditions: ramping up to 500 °C in dry nitrogen; ramping up to 500 °C in air and then cooling; ramping to 350 °C under dry nitrogen and then switching the atmosphere to dry oxygen under isothermal conditions. The results are shown in Figure 15. Figure 15a shows, in blue, the variation of the conductivity of the starting material with temperature (the heating is in nitrogen gas). The black lines/symbols show the major effects of heating in air: the oxidation begins at ca. 260 °C and results in an increase in conductivity; the conductivity peaks at 330 °C and then falls to a value below that of the starting material heated in nitrogen. This behavior reflects the increased electronic conductivity caused by the presence of both Fe²⁺ and Fe³⁺ ions and was seen previously for the system FeSb₂O₄, where the conductivity was highest for equal Fe²⁺ and Fe³⁺ concentrations. The same data plotted as log(conductivity) against 1000/T are shown in Figure 15b where the red lines are linear fits to the data prior to oxidation and on cooling of the oxidized product. The good fits indicate thermally promoted conduction with activation energies of 19.5(2) kJ mol⁻¹ and 21.3(2) kJ mol⁻¹ for the unoxidized and oxidized materials, respectively. The conductivity for Co₀.₅₀Fe₀.₅₀Sb₂O₄ is approximately an order of magnitude lower than that reported for FeSb₂O₄, which suggests that electron migration primarily involves Fe³⁺. Figure 15b shows that the conductivity of the oxidized phase, Co₀.₅₀Fe₀.₅₀Sb₂O₄, is lower than that for Co₀.₅₀Fe₀.₅₀Sb₂O₄ by a factor of about 5. Since the conduction will be dominated by the chains of octahedral transition metal ions, this is probably linked to the expansion along [001], the chain direction, during oxidation and the resulting effect on Fe–Fe separation. It should be noted that the small discrepancy between the measurement in nitrogen and the low temperature measurement in air can be traced to a significant difference in pellet densities: 50% for the pellet in air cf. 77% for the pellet in nitrogen.

Figure 15c reports data for subsequent isothermal oxidation at 350 °C of the pellet that had previously been heated in nitrogen to provide the data (in blue) shown in Figure 15a,b. The data are recorded against time after the gas was switched from nitrogen to oxygen. After an initial delay corresponding to Fe separation. It should be noted that the small discrepancy between the measurement in nitrogen and the low temperature measurement in air can be traced to a significant difference in pellet densities: 50% for the pellet in air cf. 77% for the pellet in nitrogen.
4. DISCUSSION

The NPD and TGA data clearly indicate that low-temperature oxidation of schafarzikite materials that contain both Fe$^{2+}$ and Sb$^{3+}$ is associated with oxygen entering the channels and bonding to Sb$^{3+}$ ions. However, the level of oxygen incorporation is greater than that corresponding to oxidation of Fe$^{2+}$ to Fe$^{3+}$ within the octahedral chains. Moreover, the data consistently suggest that for Fe$^{2+}$ concentrations of <0.5 pfu in the unoxidised samples, the oxygen uptake is ca. twice that expected. Mössbauer spectra reveal that charge balance associated with the extra oxygen content is provided by oxidation of Sb$^{3+}$ to Sb$^{5+}$. To examine the change in oxidation states of all four cations in Co$_{0.25}$Fe$_{0.75}$Sb$_{1.75}$Pb$_{0.25}$O$_4$, XPS data were collected but the changes observed were insufficient to resolve the details of the redox reactions. However, oxidation of neither Co$^{2+}$ nor Pb$^{2+}$ is plausible as discussed in Section 3.6.

With respect to the formation of Sb$^{5+}$ ions, it is relevant to note that an attempt to partially substitute Sb$^{3+}$ by Pb$^{2+}$ in MnSb$_2$O$_4$ resulted in oxidation of Sb$^{5+}$ rather than the desired Mn$^{2+}$, and the resulting Sb$^{5+}$ ions entered the octahedral sites by replacing some of the Mn$^{2+}$ ions. This possibility was rejected for the oxidations studied here for two reasons: (1) substitution of Sb$^{5+}$ ions in the chains of octahedra would result in significant impurity phases; (2) PXRD and NPD are both sensitive to such mixing and both confirm that no detectable substitution of Sb$^{5+}$ ions within the chains occurs. The Sb$^{5+}$ ions therefore remain on the channel walls, and it is therefore important to consider how the insertion of oxygen ions into the O3 sites can satisfy the very different stereochemical preferences of Sb$^{3+}$ and Sb$^{5+}$ ions.

For the oxidation of Co$_{0.25}$Fe$_{0.75}$Sb$_{1.75}$Pb$_{0.25}$O$_4$, the first low temperature plateau corresponds closely to oxidation of all Fe$^{2+}$ to Fe$^{3+}$ (0.50Fe$^{2+}$ forming 0.50Fe$^{3+}$) and the simultaneous oxidation of 0.25Sb$^{3+}$ to 0.25Sb$^{5+}$; this would correspond to an oxygen content of 4.50 pfu which is close to that determined by TGA, NPD, and Mössbauer spectroscopy. To consider the likely environment for Sb$^{5+}$, it is informative to explore in more detail the local environment of the O3 site. As shown in Table 3, each O3 is close to two adjacent Sb sites with Sb$^{5+}$-O distances of 1.91 and 2.30 Å. BVS considerations require an average Sb$^{5+}$-O distance of 1.97 Å in the unoxidized phase, less than the observed value of 2.10 Å (Figure 8f) because the larger Pb$^{2+}$ ion occupies the same site. An additional bond at 2.30 Å

### Table 5. Experimentally Determined and GIPAW DFT Calculated NMR Parameters ($\delta_{\text{iso}}$, $C_Q$, $\eta$, and $P_Q$) from the Diamagnetic MgO, Sb$_2$O$_3$, and MgSb$_2$O$_4$ Systems

<table>
<thead>
<tr>
<th>sample</th>
<th>site</th>
<th>isotope</th>
<th>$\delta_{\text{iso}}$ (ppm)</th>
<th>$\delta_{\text{iso}}$ error (ppm)</th>
<th>$C_Q$ (MHz)</th>
<th>$\eta$</th>
<th>$P_Q$ error (MHz)</th>
<th>$P_Q$ (MHz)</th>
<th>CASTEP</th>
<th>geom. opt.</th>
<th>$\sigma$ (ppm)</th>
<th>$\delta_{\text{iso}}$ (ppm)</th>
<th>$C_Q$ (MHz)</th>
<th>$\eta$</th>
<th>$P_Q$ error (MHz)</th>
<th>$P_Q$ (MHz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MgO$^{25}$</td>
<td>1</td>
<td>$^{25}$Mg</td>
<td>26</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>no</td>
<td>535.2</td>
<td>26$^{a}$</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sb$_2$O$_3$</td>
<td>1</td>
<td>$^{17}$O</td>
<td>167</td>
<td>5.2</td>
<td></td>
<td>0.99</td>
<td></td>
<td></td>
<td>no</td>
<td>70.8</td>
<td>167$^{b}$</td>
<td>5.93</td>
<td>0.93</td>
<td>6.73</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MgSb$_2$O$_4$</td>
<td>1</td>
<td>$^{17}$O</td>
<td>0</td>
<td></td>
<td>4.9</td>
<td></td>
<td></td>
<td></td>
<td>no</td>
<td>50.2</td>
<td>146.1</td>
<td>4.47</td>
<td>0.00</td>
<td>4.47</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$^{17}$O</td>
<td>139</td>
<td></td>
<td>4.9</td>
<td>0.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$^{25}$Mg</td>
<td>19</td>
<td></td>
<td>6.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$^{17}$O</td>
<td>0</td>
<td></td>
<td>4.9</td>
<td>0.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$^{17}$O</td>
<td>139</td>
<td></td>
<td>4.9</td>
<td>0.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$^{25}$Mg</td>
<td>19</td>
<td></td>
<td>6.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$^a$σ$^{ref}$ from Dupree and Smith. $^b$Measured σ$^{ref}$ given in Figure 13e.
would require an expansion of the three lattice bonds by 0.06 Å to 2.03 Å to retain the same BVS value. Figure 8f shows that this should readily be achieved and suggests that the larger Sb−O3 distance corresponds to 4-coordinate Sb3+. The bond at 1.91 Å, however, would require a corresponding expansion of 0.19 Å to 2.16 Å. This appears less likely, and this bond distance is more indicative of Sb5+: expected average Sb5+−O distances of 1.86, 1.94, and 2.01 Å for coordination numbers of 4, 5, and 6, respectively. It therefore seems highly likely that the Sb5+ ions, which we have shown to be essential to explain the observed oxygen contents, are adjacent to a 4-coordinate Sb3+ ion. The oxidation of Sb3+ to Sb5+ ion is accompanied by the introduction of an extra oxide ion into the structure; this can coordinate to the Sb5+ ion (Sb5+−O bond distance of 1.91 Å) to provide a 5-coordinate species without a lone pair and also provide a link to a second 4-coordinate Sb3+ ion via a longer bond (Sb3+−O bond distance of 2.30 Å). We therefore suggest the presence of defect clusters which contain two interstitial O2− ions, two 4-coordinate Sb3+ ions, and one 5-coordinate Sb5+ ion. This arrangement is shown in Figure 16 and is based on the average atom coordinates given in Table 2 from the high resolution NPD data for FeSb1.25Pb0.75O4+y. Figure 16 shows that the spatial coordination of O3 retains space for the lone pair on the 4-coordinate Sb3+ ion whose stereochemistry can be related to a trigonal bipyramid, SbO4E, where the lone pair (E) occupies an equatorial position. Sb5+, on the other hand, is 5-coordinate with a structure that can be described as either distorted square pyramidal or distorted trigonal bipyramidal. Sb5+ ions are normally octahedral in extended oxide structures, but a similar distorted geometry has previously been suggested.
clusters will be restricted by the random distribution of Pb\textsuperscript{2+} on sample after extended heating at 500°C. Site occupancy corresponds to an oxygen content of 4.5 pfu. Here, however, the changes seen in Figure 8 for temperatures above can fully explain the oxidation up to 350°C. Moreover, the structural data shown in Figure 16 could be oxidized to Sb\textsuperscript{5+} and be bonded to only one Sb\textsuperscript{3+} (probability 12.5% Pb) since it can form only two bonds at 1.91 Å. Overall, the probability that the O3 ion can form the chain shown in Figure 16 is 86%. Of course, as more clusters form, the probability of creating new clusters will gradually decrease.

The electronic conductivity data in Figure 15 provide a means of monitoring the oxidation of Fe\textsuperscript{2+} ions to Fe\textsuperscript{3+}, since the conductivity is a maximum when the amounts of these species are equal. Unfortunately, the three-dimensional conductivity of these materials is not high, and it would be very useful to produce single crystals to allow determination of the anisotropic nature of the conductivity. It would also be interesting to synthesize and characterize materials containing some second/thirdrow transition metal ions into the chains of octahedra in an attempt to enhance the conductivity.

The temperature at which oxidation is rapid suggests that oxygen diffusion within the channels of oxidized schafarzikite phases is high at low temperatures, but it is not straightforward to quantify this because of the electronic component and the difficulty in achieving high quality sintered ceramic samples. Examination of cations such as Ru\textsuperscript{4+} and Pt\textsuperscript{4+} would be of particular interest because of the possible enhancement to electronic conductivity. The materials studied here may also have high oxide ion conductivity, but their high relative molecular mass and toxicity are a limiting factor for their use in many energy applications.

5. CONCLUSIONS

This study has revealed that phases related to FeSb\textsubscript{2}O\textsubscript{4} undergo a unique low temperature oxidation in air or oxygen: simultaneous oxidation of Fe\textsuperscript{2+} to Fe\textsuperscript{3+} and Sb\textsuperscript{3+} to Sb\textsuperscript{5+} occurs, and charge balance is achieved by oxide ions entering the structural channels and bonding to Sb in the channel walls. The resulting Sb\textsuperscript{5+} ions remain in the channel wall sites to provide a mixture of Sb\textsuperscript{3+} and Sb\textsuperscript{5+}. It is not simple for the interstitial oxide ion site to satisfy the bonding requirements of both Sb\textsuperscript{3+} and Sb\textsuperscript{5+}.
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorg-chem.6b02466.

Full experimental details: synthesis, thermal characterization, structural characterization (PXRD, NPD), Raman, Mössbauer, and NMR spectroscopy, and conductivity measurements. XPS spectra (PDF)

**ACKNOWLEDGMENTS**

The authors declare no competing financial interest.

**AUTHOR INFORMATION**

**Corresponding Author**
*E-mail: c.greaves@bham.ac.uk.*

**ORCID**
Colin Greaves: 0000-0002-9568-3697

**Notes**
The authors declare no competing financial interest.

**REFERENCES**


(5) Stähli, S. The crystal structure of ZnSb2O4 and isomorphous compounds. *Arkiv Kemi, Min. och Geol. 1943*, 17B, 1–7.


(14) Camby, J.; de Laune, B. P.; Greaves, C. The Structures and Magnetic Properties of FeCo1−xSb2O4 and MnCo1−xSb2O4, 0 ≤ x ≤ 1. *J. Mater. Chem. C 2016*, 4, 201–208.


(22) Kharbish, S. Raman spectra of minerals containing interconnected As(Sb)O3 pyramids: trippkeite and schafarzikite. *J. Geosci. 2012*, 57, 53–62.


(26) Sridivastava, A.; Chauhan, M.; Padogaonker, R. Theoretical investigation of high-pressure phase transitions in Mg1−xSrO. *Phase Transitions 2014*, 87, 126–135.


