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Investigation of the Relationship between the Structure and Conductivity of the Novel Oxide Ionic Conductor Ba3MoNbO8.5

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ABSTRACT: A variable temperature neutron diffraction study of the novel oxide ion conductor $Ba_3MoNbO_{8.5}$ has been performed between 25 °C and 600 °C. Non-monotonic behaviour of the cell parameters, bond lengths and angles are observed indicating a structural rearrangement above 300 °C. The oxygen/vacancy distribution changes as the temperature increases so that the ratio of $(Mo/Nb)O₄$ tetrahedra to $(Mo/Nb)O₆$ octahedra increases upon heating above 300 °C. A strong correlation between the oxide ionic conductivity and the number of $(Mo/Nb)O₄$ tetrahedra within the average structure of Ba₃MoNbO_{8.5} is observed. The increase in the number of $(Mo/Nb)O_4$ tetrahedra upon heating from 300 – 600 \degree C most likely offers more low energy transition paths for transport of the O²⁻ ions enhancing the conductivity. The unusual structural rearrangement also results in relaxation of Mo(1)/Nb(1) and Ba(2) away from the mobile oxygen, enhancing the ionic conductivity. The second order Jahn-Teller effect most likely further enhances the distortion of the MO_{4}/MO_{6} polyhedra as distortions created by both electronic and structural effects are mutually supportive.

INTRODUCTION

Solid oxide electrolytes are the basis of two important energy technologies, Solid Oxide Fuel Cells (SOFCs) and Solid Oxide Electrolyser Cells (SOECs). SOFCs offer an efficient alternative to combustion technology for electricity production. The advantages of the high operating temperature for SOFCs include the possibility of running directly on practical hydrocarbon fuels without the need for complex and expensive external fuel reformer and purification systems. The high quality exhaust heat released during operation can be used as a valuable energy source, either to drive a gas turbine when pressurized or for combined heat and power (CHP) applications, so further increasing system efficiency.

To date, oxide ion conductivity has been reported in a number of structural families including fluorite-like systems¹, silicon and germanium apatites², La₂Mo₂O₉ (LAMOX) materials ³, $Bi_4V_2O_{11}$ derivatives (BIMEVOX)⁴ and complex oxides with $GaO₄$ tetrahedral moieties $5.$ To simplify certain materials issues, such as sealing and to enable the use of cheaper steel interconnects, it is desirable to lower the SOFCs' operating temperature from 800 ^oC, to an intermediate range of 400 °C – 600 °C ⁶. So far, materials that exhibit high conductivity at low temperature, such as aliovalent doped ceria or isovalent cation stabilised bismuth oxide, are less stable under the reducing fuel environment. The oxide ion conductivity of a material is strongly dependent on the crystal structure,

hence in order to reach the objective of an intermediate temperature fuel cell it is important to discover new structural families of oxide ion conducting materials.

The perovskite structural family offers great potential for the discovery of new oxide ion conductors due to its structural adaptability. Several perovskite materials with high oxide ion conductivity have been reported including strontium and magnesium-doped lanthanum gallates (LSGM) $⁷$ and more recently the well-known ferroelectric</sup> material $\text{Na}_{0.5}\text{Bi}_{0.5}\text{TiO}_3$ ⁸ and the perovskite derivative NdBaInO₄⁹. We have recently identified oxide ion conductivity in $Ba_3MoNbO_{8.5}$ ¹⁰ which exhibits a bulk conductivity of 2.2 x 10⁻³ S cm⁻¹ at 600 °C. Ba₃MoNbO_{8.5} crystallises in a hybrid of the 9R hexagonal perovskite and palmierite structures, which is a novel and so far unique crystal structure. Units of $(Mo/Nb)O₄$ and $(Mo/Nb)O₆$ coexist within the structure, forming a disordered arrangement of Mo/Nb tetrahedra and octahedra. Ba₃MoNbO_{8.5} exhibits oxygen transport numbers of 0.99 in $air/O₂$ and 0.92 in air/5% H_2 in Ar at 600 °C, suggesting that $Ba_3MoNbO_{8.5}$ is an oxide ion conductor with negligible electronic conductivity in \ar{air}/O_2 and that a small amount of electronic conduction is observed in air/5% H_2 in Ar. The phase is also surprisingly stable in 5% H_2/N_2 ¹⁰. Ba₃MoNbO_{8.5} is the first hexagonal perovskite to display such promising transport numbers / oxide ion transport at low temperature. All other hexagonal systems previously reported exhibit mixed oxide ion and electronic and/or proton conductivi-

ty with low conductivities and low transport numbers \mathbf{u}, \mathbf{u} . In order to further investigate the correlation between the structural and the transport properties in $Ba_3MoNbO_{8.5}$ we have performed a variable temperature neutron diffraction study between 25 to 600 °C. A structural rearrangement occurs above 300 $^{\circ}$ C so that the ratio of $(Mo/Nb)O₄$ tetrahedra to $(Mo/Nb)O₆$ octahedra increases upon heating between 300 °C and 600 °C. The results demonstrate that there is a strong correlation between the number of tetrahedra within the average structure and the observed ionic conductivity.

EXPERIMENTAL

Stoichiometric amounts of BaCO₃ (99.999%, Aldrich), $MoO₃$ (99.5+%, Aldrich) and $Nb₂O₅$ (99.99%, Aldrich) were ground, pressed into 13 mm pellets and calcined in an alumina crucible at 900 \degree C for 10 hours. The pellets were subsequently reground, pelleted and heated at 1100 °C for 48 hours and then cooled to room temperature at 5 °C/min. The latter heating step was repeated until phase pure products were obtained.

Variable temperature neutron diffraction patterns in the temperature range 25 to 600 °C were recorded on the high-resolution powder diffractometer D2B at the Institut Laue-Langevin (ILL) in Grenoble, France. A sample of 5 grams of $Ba_3MoNbO_{8.5}$ was inserted in an open quartz tube and heated up to the desired temperatures. Data were collected at λ = 1.59432 Å with a total collection time of 2.5 hours for each temperature step.

For the impedance spectroscopy measurements, a pellet of ∼ 10 mm diameter and ∼ 1.5 mm thickness was prepared from a powder sample of $Ba₃MoNbO_{8.5}$ and sintered at 1100 °C for 48 hours (to achieve > 95 % of the theoretical density). Pt electrodes were painted on both sides of the pellet using a Pt-paste (Metalor 6082). Impedance spectra were recorded with a Solartron 1260 impedance analyser in the frequency range 0.1 Hz $-$ 1 MHz with an applied alternating voltage of 0.1 V. Data were recorded upon cooling from 600 \degree C to 300 \degree C in a sealed tube furnace under the flow of dry air, measuring every 15 \degree C and allowing 2 hours of equilibration at each temperature step. The obtained data were corrected by the geometrical factor of the sample and treated with the ZView software (Scribner Associates, Inc.). The data were fitted using the model described previously¹⁰.

RESULTS AND DISCUSSION

 Neutron diffraction data at all temperatures were fitted by the Rietveld method using the GSAS/EXPGUI program ^{13, 14}. Modelling of the background was performed by the shifted Chebyschev polynomial function and the peak shapes were fitted using a Pseudo-Voigt function. The hybrid 9R polytype – palmierite model previously reported ¹⁰ (Figure 1) was used as a starting model for the variable temperature refinements. This model is formed by a disordered arrangement of Mo and Nb cations in mixed tetrahedral – octahedral coordination, which is created by the occupation of both the 9*e* (9R polytype) and 6*c* (palmierite) oxygen positions. Oxygen atoms are present at three different Wyckoff sites: O(1) in 18*h*, O(2) in 9*e* and O(3) in 36*i*; the latter position is employed instead of 6*c* to take into account the dynamic disorder evidenced by the large U_{11} , U_{22} and U_{12} values for $O(3)$ found in the previous structural refinement¹⁰. Mo/Nb cations (M) are in 6*c* and 3*b*, while the Ba atoms occupy the 6*c* and 3*a* sites.

An excellent fit to the neutron data was obtained with the hybrid model described above for all temperatures (the refined data recorded at 25° C and 600 °C are displayed in Figure 2, while the Rietveld fits to the data collected at the remaining temperatures are presented in the Supporting Information).

Figure 1. Crystal structure of Ba3MoNbO8.5. The hybrid structural model formed by the superimposition of the 9R-polytype and the palmierite sub-units representing the average structure of the system. Colours indicate: light green Ba(1), dark green Ba(2), blue M (1), cyan $M(z)$, red $O(1)$, orange $O(2)$ and yellow $O(3)$.

There is no evidence of peak splitting or superstructure peaks and the *R*3*m* H symmetry space group is observed over the whole temperature range. The refined atomic positions and agreement indices are displayed in Table S1. The atomic displacement parameters, U_{ii} , were refined anisotropically for all atoms except $O(3)$, for which an U_{iso} parameter was used as previously reported¹⁰. Attempts to independently refine the $M(1)$ and $M(2)$ U_{ii} parameters generated unrealistic U_{33} values; therefore the U_{ij} parameters of the M atoms on the 6*c* and 3*b* sites were constrained to be the same. The displacement parameters for all atoms are reported in Table S₁. The atomic U_{ii} anisotropic values generally increase with the temperature as expected. Overall, the obtained thermal displacement values are within the range expected for this type of material, with similar values being found for α - and β-La₂Mo₂O₉¹⁵, γ -Bi₄V₂O₁₁¹⁶, La_{1.54}Sr_{0.46}Ga₃O_{7.27}⁵, and LSGM materials 17 . The large U_{33} values indicate that the M cations exhibit motion mainly parallel to the *c*-axis (Figure S2). Ba(1) presents $U_{11} = U_{22} > U_{33}$, indicating preferential thermal motion on the *ab* plane. On the contrary, the Ba(2) atom shows larger U_{33} values, evidencing aniso-

tropic motion along the *c*-axis. Ba(1) is encircled by a belt of $O(2)$ and $O(3)$ sites, while Ba(2) and M(1) are directly above (or below) the $O(2)$ and $O(3)$ positions. The distribution of occupied/unoccupied $O(z)$ and $O(3)$ sites creates different coordination environments for the metal cations, thus inducing motion on the *ab* plane (Ba(1)) or along the *c*-axis ($Ba(2)$ and $M(1)$). The $O(1)$ thermal ellipsoids are oriented perpendicularly to the M–O bonds. The O(2) ellipsoids are nearly spherical in the whole temperature range and $O(3)$ exhibits large U_{iso} values as previously reported¹⁰.

Figure 2. Rietveld refinement fit to the variable temperature neutron diffraction data of Ba3MoNbO8.5. Rietveld refinement fits of neutron diffraction data collected at 25 °C and 600 °C are shown. Black dots indicate the observed data, the red line the Rietveld fit, the blue line the difference between the observed and the calculated patterns, the green line the background function and the pink bars are the reflection positions.

The temperature dependency of the *a* and *c* cell parameters are shown in Figure 3 and non-monotonic behaviour is observed. A clear change in slope of both *a* and *c* is observed above 300 °C suggesting a possible structural modification. The fractional occupancies of the atoms in the structural model obtained previously¹⁰ were used as starting values for the refinement of the variable temperature neutron diffraction data. The Ba and $O(1)$ fractional occupancies refined to within \pm 1% of full occupancy and were fixed at 1.0. The occupancies of the other atoms were refined from their initial values. The refined fractional occupancies are presented in Table S1. Figure 4 shows the temperature dependency of the fractional occupancies of $M(1)$, $M(2)$, $O(2)$ and $O(3)$.

Figure 3. Unit cell dimensions of Ba3MoNbO8.5. Thermal dependency of the *a***- and** *c***-axis of the unit cell of Ba3MoNbO8.5. The linear fits evidence the change in slope above 300 °C.**

Figure 4. Variation of fractional occupancies with the temperature. Temperature dependency of the fractional occupancies of M_1 and M_2 and O_2 and O_3 . Con**necting lines are guides for the eye only.**

There is little change in the fractional occupancies between 25 °C – 300 °C. Above this temperature a sudden increase in the $M(1)$ fractional occupancy, with a concomitant decrease in the $M(z)$ fractional occupancy, is evidenced. The same behaviour is observed for the $O(z)$ and $O(3)$ fractional occupancies, which respectively decrease and increase above 300 °C. Above 500 °C there is little variation in the fractional occupancies of $M(1)$, $M(2)$, $O(2)$ and $O(3)$. An overall oxygen stoichiometry of 8.5 was obtained at all temperatures.

 $Ba(1)$, $Ba(2)$ and $O(1)$ form the rigid part of the Ba₃MoNbO_{8.5} structure, while M(1), M(2), O(2) and O(3) are responsible for the variable tetrahedral/octahedral coordination. Therefore, changes in the occupancies of the latter atoms indicate that the structure is able to adjust the ratio of tetrahedra to octahedra and, consequently, oxygen/vacancy distributions as the temperature increases. The results demonstrate that the number of $(Mo/Nb)O₄$ tetrahedra increases upon heating above 300 $^{\circ}C.$

The variation of the fractional occupancy of $O(3)$ and the Arrhenius plot of the bulk conductivity of $Ba₃MoNbO_{8.5}$ is presented in Figure 5. The bulk conductivity is 2.2 x 10⁻³ S cm⁻¹ at 600 °C with an activation energy of 1.211 (6) eV for temperatures between 300 - 500 °C. There is a clear correlation between the change in the crystal lattice and the electrical properties at temperatures between 300 °C and 600 °C. The rearrangement of $M(1)$, $M(2)$, $O(2)$ and $O(3)$ within the crystal structure results in an increase of the ratio of $M(1)O₄$ tetrahedra to $M(1)O_6$ octahedra with concomitant changes in the bond lengths and angles. At 300 °C there are ∼ 49.9% tetrahedra within the average structure which increases to ∼

63.5% at 500 °C. Above 500 °C the rate of increase in the number of tetrahedra within the crystal structure of Ba₃MoNbO_{8.5} decreases so that at 600 °C there are ∼ 64.8% tetrahedra. At the same time a change in slope is evidenced in the Arrhenius plot. The results clearly demonstrate that the magnitude of ionic conductivity observed in $Ba_3MoNbO_{8.5}$ is strongly related to the number of $Mo/NbO₄$ tetrahedra within the average crystal structure.

Increasing the number of $(Mo/Nb)O₄$ tetrahedra most likely offers more low energy transition paths for transport of the $O²$ ions, further enhancing the conductivity. The preference for a tetrahedral environment gives a large number of possible oxygen positions which are important for collective mechanisms involving strongly correlated motions of sets of tetrahedra as previously observed for the brownmillerite $Ba_2In_2O_5^{-18}$.

The self-rearrangement of the Mo/Nb and oxygen occupancies is striking evidence of the ability of the metal lattice to support variable coordination environments. It has been demonstrated that the presence of flexible *d*metal cations is essential for the ionic transport of various oxide ion conductors. The high oxygen conduction of $β$ - $La₂Mo₂O₉$ is thought to be facilitated by the capacity of the molybdenum atom to assume co-ordinations between 4 and $6^{3,19}$. In the BIMEVOX materials the oxygen motion proceeds through the decomposition and reformation of the V–O units on the perovskite layers. An oxygen atom is transferred from a $VO₆$ octahedron to a vacancy contained in a nearby $VO₄$ tetrahedron, forming a shortliving VO_5VO_5 complex. The complex then separates into a VO₄ tetrahedron and a V[']O₆ octahedron, completing the oxygen migration ²⁰. In the same way, $Ba_3MoNbO_{8.5}$ possesses a flexible lattice in which the M atoms are readily able to change coordination, creating a potential avenue for conduction as it is possible for an oxide ion to migrate from one $M(1)O_X$ unit to the other. The change in the oxygen fractional occupancies with temperature further supports the hypothesis that the oxide ion migration occurs via the partially occupied $O(z)$ and $O(3)$ sites¹⁰.

Selected bond lengths and angles for $Ba₃MoNbO_{8.5}$ at the various temperatures are displayed in Table S2. Information on how the M polyhedra respond to the oxygen/vacancy rearrangement can be obtained by following some key bond lengths and angles. Figure 6 (b) shows a superimposition of the coordination of the $M(1)$ - $O(1)$ ₃ $O(3)$ tetrahedra and the M(1)– $O(1)$ ₃ $O(2)$ ₃ octahedra. $M(1) - O(1)$ ₃O(3) and $M(1) - O(1)$ ₃O(2)₃ share the same $M(1)O(1)$ ₃ unit, which presents three equal $M(1)-O(1)$ bond lengths and three equal $O(1)-M(1)-O(1)$ angles (α). $M(1)-O(1)$ ₃O(3) is then defined by the $M(1)-O(3)$ bond length and the $O(1)$ – $M(1)$ – $O(3)$ angle (β), obtained by the average of the three possible angles given by the $O(3)$ split position. The $M(1)-O(1)$ ₃ $O(2)$ ₃ octahedron is defined by the M(1)–O(1) bond length, α, the M(1)–O(2) bond lengths and the $O(1)$ – $M(1)$ – $O(2)$ and $O(2)$ – $M(1)$ – $O(2)$ angles (γ and δ respectively). The temperature dependencies of selected distances and angles defined above are displayed in Figure 6 (a). All of the bond lengths and angles

exhibit little variation with temperature below 300 °C but evidence a sizeable change above 300 °C. Above 300 °C there is an increase in α and decrease in γ, β and δ . The $M(1)-O(1)$ bond length contracts whilst both $M(1)-O(3)$ and $M(1)-O(2)$ expand (Table S2, Figure 5 (a) and (b)).

The thermal modifications of the $M(1)$ polyhedron are reflected to the neighbouring $M(z)O(1)$ ₆ octahedron where the $M(2)-O(1)$ bond lengths and the $O(1)-M(2)$ -O(1) angles both increase with the temperature resulting in a net expansion of the $M(z)O(1)_6$ octahedra.

Figure 6. Thermal relaxation of the M(1) polyhedra. Colours indicate: blue M(1), cyan M(2), red O(1), orange O(2) and yellow O(3) (a) Temperature dependency of selected bond distances and angles for the M(1) polyhedra. (b) Representation of the M-O lattice relaxation; black arrows show the modifications of the bond lengths and angles at 600 °C **from the "equilibrium" values at 25 °C. The red arrow represents the average M(1) displacement. (c) Variation of the M(1) displacement with the temperature. In (a) and (c) lines are guides for the eyes only; in the inset in (c) the red line is the linear fit to the data and the numbers are the calculated activation energies (in eV).**

 $Ba₃MoNbO_{8.5}$ presents an irregular thermal variation of the cell parameters, bond lengths and angles. These quantities exhibit nonlinear behaviour, with abrupt changes above 300 °C. The trends in the variation of the cell parameters, bond lengths and angles with temperature mimic the thermal variation of the conductivity (Fig. 5) where the largest changes arise between 300 °C and 500 °C. The Ba₃MoNbO_{8.5} structure relaxes in response to the rearrangement of the oxygen/vacancy distribution. The M(1) polyhedra are already distorted at 25 °C ¹⁰ (Table S2), due to the effect of the second order Jahn-Teller distortion²¹. Examination of the variation of the M(1)-O distances and angles with temperature indicates further displacement of the $M(1)$ atom away from the $O(2)/O(3)$ sites, towards the $[O(1)-O(1)-O(1)]$ face of the M(1)-O(1)₃ unit (Figure ζ (a) and (b)). The magnitude of the displacement (*D*) can be obtained by calculation of the distance between $M(1)$ and the $[O(1)-O(1)-O(1)]$ face. This is given by D = $-M(1)-O(1)cos(β)$ and the displacement can be evaluated by the difference between the $M(1)$ –[O(1)- $O(1)-O(1)$] distance at temperature, T, and the M(1)–[$O(1)$ – $O(1)-O(1)$] distance at 25 °C. The M(1) displacement is plotted against the temperature in Figure 6(c). The results clearly demonstrate that the variation of the oxygen/vacancy distribution above 300 °C strongly affects the relaxation of the M–O framework. We note that the second order Jahn-Teller effect previously reported for $Ba₃MoNbO_{8.5}$ ¹⁰ may further enhance the distortion of the MO_{4}/MO_{6} polyhedra, as distortions created by both electronic and structural effects are mutually supportive 21 . The Ba lattice relaxes in a similar way, with $Ba(z)$ displacing away from $O(2)/O(3)$ and consequent relaxation of the $Ba(1)-O$ framework (Figure S_3).

Displacement of the metal atoms of the cationic lattice away from the mobile oxide ions has been reported in several different oxide ion conductors. In doped LaGaO₂ perovskites, La and B-site cations shift away from the mobile oxygen atom 22 . Similarly, the Si atoms in the $La_{9.33}Si₆O₂₆$ apatite move away from the channels in which migration of the interstitial oxygen atoms occurs 23 . In $Ba₃MoNbO_{8.5}$, the average M(1) displacement is an effect of the rearrangement of the oxygen/vacancy distribution at the $O(2)$ and $O(3)$ sites but can also be correlated to the energetics of the ionic conduction as displacement of *d*-metal cations from oxygen vacancies in the lattice is thought to lower the motional enthalpy required for the mobility of the oxide ions 24 . The average structural relaxation, obtained from the neutron diffraction data, is therefore most likely a consequence of both the structural rearrangement and the dynamic and local structural reorganisation induced by the oxygen hopping.

The oxide ionic conductivity is thought to arise in the $Ba(1)O(2,3)_{2.5}$ layer. Oxygen mobility at the partially occupied $O(2)$ and $O(3)$ sites is evidenced by the change in the fractional occupancies with temperature. In addition the short distance between the 9*e* and 36*i* positions (< 2.2 Å) excludes simultaneous occupancy of adjacent $O(z)$ and $O(3)$ sites. This leads to a random distribution of vacancies on the $Ba(1)O(2,3)_{2.5}$ layer, which are available for the

ionic migration. The partially occupied $O(2)$ and $O(3)$ sites on the Ba(1)O(2,3)_{2.5} layers are distributed in a zigzag fashion inside channels created by the rigid $Ba(1)$, $Ba(2)$ and $O(1)$ framework (Figure S₄). These channels propagate in the [100], [010] and [110] directions and might provide possible conduction pathways. It is most likely that the movement of oxide ions is via octahedral and tetrahedral interchange and suggests a cooperative motion such as an interstitialcy mechanism.¹⁰. This process is assisted by the ability of the system to locally and dynamically reorganise its oxygen/vacancy distribution. Above 300 °C it is proposed that the increase in the number of tetrahedra, coupled with the displacement of $M(1)$ and $Ba(2)$ away from $O(2)$ and $O(3)$ enhances the electrical properties, so that a conductivity comparable to other leading solid oxide ion conductors is observed.

CONCLUSIONS

In summary this variable temperature neutron diffraction study demonstrates an unusual structural rearrangement in Ba₃MoNbO_{8.5} above 300 °C so that the ratio of tetrahedra to octahedra increases and, consequently the oxygen/vacancy distribution changes as the temperature increases. We propose that the conductivity is enhanced by the increase in the proportion of MO ₄ tetrahedra between 300 \degree C and that substitution of Nb⁵⁺ with an aliovalent cation with preferential tetrahedral coordination may be a way to further optimise the conductivity at low temperature in $Ba_3MoNbO_{8.5}$. The change in vacancy distribution combined with the oxide ion migration results in displacement of the $M(1)$ and $Ba(2)$ sites away from the mobile oxygen, lowering the motional enthalpy required for the mobility of the oxide ions and enhancing the conductivity.

ASSOCIATED CONTENT

Supporting Information is available free of charge via the Internet at http://pubs.acs.org. The supporting information includes tables and figures of refined atomic positions and bond lengths and angles from Rietveld refinement of neutron diffraction data, figures of the Rietveld refinement fits to the neutron diffraction data at all temperatures and an alternative view of the $Ba_3MoNbO_{8.5}$ structure showing the rigid Ba- $O(1)$ and the flexible $M(1,2)-O(2,3)$ frameworks.

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Author Contributions

The manuscript was written through contributions of all authors.

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Chemistry of Materials

