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BaTa_{0.5}Li_{0.5}O_{2.5}: a new highly oxygen deficient perovskite oxide-ion conductor†

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High oxide-ion and proton conductivity in perovskite oxides has been well documented in the literature. Herein, we report a highly oxygen deficient perovskite oxide BaTa_{0.5}Li_{0.5}O_{2.5} (BTLO) as a new oxide-ion conductor. The material exhibits a simple primitive cubic perovskite structure (S.G.: Pm3m) with lattice parameter a = 4.1024(1) Å and completely disordered Ta and Li cations on the B-site. The ionic conduction is predominated by oxide-ion over a wide range of oxygen partial pressure and temperature, with an ionic conductivity comparable to the state-of-the-art yttria-stabilized zirconia. Ab initio molecular dynamics (AIMD) simulations reveal that oxide-ion diffusion follows three-dimensional pathways, involving oxygen vacancy hopping along the edges of (Ta/Li)O₄ tetrahedra and (Ta/Li)O₆ octahedra. This work may inspire future work to discover new oxide-ion conductors in similar systems.

1. Introduction

The oxide-ion electrolyte is a core functional component in solid oxide cells (SOCs) for high-efficiency and low-emission power/chemical production.¹⁻⁵ They not only define the chemistry of SOCs, but also determine the working temperature. The latter is of paramount importance to the commercialization of SOC technology because too high a working temperature would invoke unwanted chemical reactions with the adjacent components, causing performance degradation. The current benchmark SOCs based on 8 mol% Y2O3-stabilized ZrO2 (YSZ) oxideion electrolyte must have a thickness <10 µm in order to achieve reasonable performance even at ≥700 °C. However,

≥700 °C is still considered too high to reach commercially meaningful 40 000+ hours of operation for SOCs. Therefore, new oxide-ion electrolytes that can further lower SOC's working temperature are of high interest.

The magnitude, purity, and stability of oxide-ion conductivity in both oxidizing and reducing atmospheres are the top three criteria for the discovery of new oxide-ion conductors. In this regard, the early oxide-ion conductors were primarily found in solid oxides with cubic fluorite structures. Examples include stabilized ZrO2 and CeO2. In recent decades, the search for new oxide-ion conductors has been expanded to perovskite or perovskite-related oxides. 7,8 The well-known examples are Sr and Mg doped LaGaO₃ (LSGM), 9,10 Na_{0.5}Bi_{0.5}TiO₃, 11 hexagonal perovskites $Ba_3NbMO_{8.5}$ (M = Mo, W) and $Ba_7Nb_4MoO_{20}$, ^{12,13} Ruddlesden-Popper-type $La_{2-x}M_xBaIn_2O_{7-0.5x}$ (M = Ca, Sr, Ba), ¹⁴ Dion-Jacobson-type $CsBi_2Ti_2NbO_{10-\delta}$, ¹⁵ Aurivillius-type Bi₄Ti₃O₁₂, ¹⁶ and pyrochlore Sr₂Nb₂O₇. ¹⁷ Besides these perovskite or perovskite-related oxides with relatively low oxygen vacancy concentrations, is the brownmillerite structure with the general formula A₂B₂O₅, including the well-known Ba₂In₂O₅-based materials 18,19 and Sr₂ScGaO₅-based materials, 20,21 which can be viewed as highly oxygen-deficient perovskites with one-sixth of the oxygen sites vacant and ordered in the tetrahedral layers, which also show good oxide ion conduction properties, especially at temperatures above the oxygen vacancy order-disorder transition or when the disordered oxygen vacancies are stabilized at a low temperature by cation doping. While high oxideion conductivity is generally observed, these materials (except LSGM) do not have sufficient ionic purity and chemical stability in SOC operating environments. Therefore, direct use of these materials in SOCs still presents a challenge.

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Herein, we report on a primitive cubic structured perovskite BaTa_{0.5}Li_{0.5}O_{2.5} (BTLO) as a new oxide-ion conductor, the nominal chemical composition of which can be viewed as the same as the general formula of brownmillerite. We show that Ta and Li ions are in a complete disordered state on the B-site in BTLO and oxide-ion conductivity is nearly purely ionic under SOC operating conditions. We also discuss oxide-ion conduction mechanisms using ab initio molecular dynamic (AIMD) simulations.

2. Experimental procedure

2.1 Material synthesis

A traditional solid-state reaction method was employed to prepare BTLO samples with BaCO₃ (99.0%, Aladdin), Ta₂O₅ (99.99%, Macklin), and Li₂CO₃ (99.0%, Aladdin) as starting materials. After stoichiometric amounts of the starting materials were intimately mixed and pressed into pellets, they were first calcined at 800 °C for 12 h, then ground, pelletized, and sintered again at 1100 °C for 48 h. To compensate for the loss of lithium during hightemperature sintering, excess Li₂CO₃ (10 mol%) was added into the samples, and the pressed pellets were buried in the original powders.

2.2 Characterization

X-ray powder diffraction (XRD) was performed using a Panalytical X'pert Pro X-ray diffractometer with Cu Kα radiation. Topas-Academic software²² was used for Rietveld refinements. Atomic absorption analysis was performed using a PinAAcle 900T atomic absorption spectrometer to check the stoichiometry of the samples. Electron diffraction (ED) was performed on a JEOL JEM-2100F field-emission transmission electron microscope with an accelerating voltage of 200 kV. Time-of-flight (TOF) powder neutron diffraction (NPD) data were collected from a general-purpose powder diffractometer (GPPD) (90° bank) at the China spallation neutron source (CSNS).²³

The electrical properties such as ionic conductivity and ionic transport number determined from an oxygen concentration cell were investigated using a Bio-Logic VSP electrochemical system. For electrochemical impedance spectroscopy (EIS) measurements, the frequency was swept from 10⁶ to 10⁻¹ Hz with an AC amplitude of 10 mV. For all the cells subject to electrical characterization, electrodes were prepared by coating Pt paste on opposite sides of the pellets and cured at 750 °C for 1 hour to remove any organic components. Before the impedance measurements, the temperature was equilibrated at each setpoint for an hour. The measurements were conducted from 300 to 800 °C in various atmospheres.

2.3 Ab initio molecular dynamics (AIMD) simulations

To study the oxide-ion conduction mechanism, first-principles calculations are carried out within the density functional theory (DFT) framework as implemented in the Vienna Ab Initio Simulation Package (VASP),24 with the generalized gradient approximation (GGA-PBE)25 and a projector-augmented wave (PAW) approach.²⁶ The cutoff energy was set to 520 eV with a force criterion of 1.0×10^{-2} eV Å⁻¹ for the lattice optimization and an energy tolerance criterion of 10^{-4} eV. For modelling the targeted composition of BaTa_{0.5}Li_{0.5}O₃, a supercell 2 $\sqrt{2}$ × $2\sqrt{2}$ × 4 (144 atoms) is constructed. Moreover, we used a special quasi randoms structure to represent the cation ordering in the current structure by assuming all ions are randomly distributed.²⁷ For all calculations, a reciprocal space discretization of 25 K-points per \mathring{A}^{-1} was applied, and the convergence criteria were set as 10^{-6} eV for electronic loops and 0.02 eV Å⁻¹ for ionic loops. All the AIMD calculations are performed in an NVT ensemble with a time step of 2 fs and using a Nose-Hoover thermostat²⁸ for a period of 500 ps. To perform extrapolation following the Arrhenius relationship, three temperatures, e.g., 2000 K, 1800 K and 1600 K were used, at which the conductivity at 1073 K (800 °C) was extrapolated.

Results and discussion

3.1 Phase and structure analysis

Since there is a possibility of Li-loss during sintering, the chemical stoichiometry in the as-made BTLO sample was first chemically analyzed using an atomic absorption technique. The result indicates a Li content of 1.18 \pm 0.1 wt% νs . nominal 1.279 wt% in the sample, suggesting ~8% percent Lideficiency in the sample. After confirming the composition, the XRD pattern of Fig. 1(a) further suggests that BTLO possesses a primitive perovskite unit cell with space group $Pm\bar{3}m$; the latter is further verified by the SAED analysis shown in Fig. 1(b).

To further investigate the defect structure of BTLO, especially the exact Li content and oxygen vacancy concentration in the crystal structure, Rietveld structural refinements were carried out against the NPD data using a typical simple cubic perovskite structure as the model, in which Ba atoms were placed at the A site (1a), while Ta and Li were first mixed on the B site (1b), and O on the 3c site. For an initial refinement, the occupancies of both Ta and Li were simply set to be 0.5 and fixed without being refined according to the nominal composition and based on the experimental chemical analysis. This resulted in a poor fitting result with the reliability of $R_{\rm wp} \sim$ 8.23% and $R_p \sim 6.12\%$, see Fig. 2(a). To address this issue, we allow the sum of Ta and Li occupancies to be the unit, instead of a fixed value of 0.5. This significantly improves $R_{\rm wp}$ to $\sim 2.26\%$ and $R_{\rm p}$ to ~ 2.20 . The resultant occupancies are 0.592(1) and 0.408(1) for Ta and Li, respectively. In addition, the refinement yields O-occupancy of 0.858(4), implying a high concentration of oxygen vacancy, and Ba-occupancy of 1.013(4), implying a nearly full occupation on the A site. The final refined composition of BTLO is BaTa_{0.592(1)}Li_{0.408(1)}O_{2.574(5)}, slightly deviated from the nominal BaTa_{0.5}Li_{0.5}O_{2.5}. The Li concentration of 0.982 wt% by the refinement is a little bit lower than the chemically analyzed result of 1.18 \pm 0.1 wt%. The final refinement is shown in Fig. 2(b) and the corresponding structural parameters are summarized in Table 1.

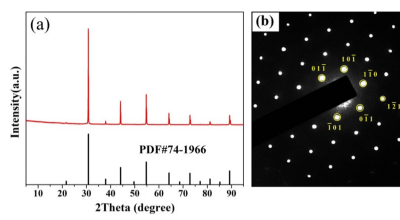
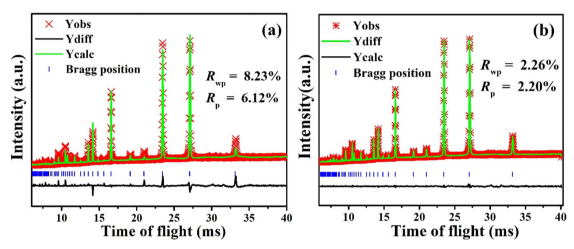


Fig. 1 (a) XRD pattern and (b) SAED pattern of BaTa_{0.5}Li_{0.5}O_{2.5}



(a) Refinement plots with Li and Ta occupancies fixed at 0.5; (b) the final refinement plots with the sum of Li and Ta occupancies fixed at the unit.

Table 1 The final refined structural parameters of BaTa_{0.5}Li_{0.5}O_{2.5}

Atoms	x	у	z	Occupancy	Beq (Ų)	BVS
Ва	0	0	0	1.0	1.36(3)	2.32
Ta/Li	0.5	0.5	0.5	0.5978(3)/0.4022(3)	0.46(2)	4.27/1.25
O	0.5	0.5	0	0.858(4)	1.75(3)	1.94

Lattice parameters: a = 4.0910(3) Å; space group, $Pm\bar{3}m$; interatomic distances (Ta/Li)-O (×6): 2.0455(2) Å; Ba-O (×12): 2.8928(2) Å. The refined composition: $BaTa_{0.592(1)}Li_{0.408(1)}O_{2.574(5)}$.

There have been several reports in the literature on perovskite oxides ABO₃ with B-site containing Ta/Nb and alkaline metals, which crystalize into a simple cubic perovskite or hexagonal eight-layered perovskite structure. Examples include $Ba_4(Ta/Nb)_3LiO_{12}$, $Ba_4(Ta/Nb)_3NaO_{12}$, $Sr_4(Ta/Nb)_3LiO_{12}$, and Sr₄Ta₃NaO₁₂.²⁹ For Ba₄(Ta/Nb)₃LiO₁₂, it forms a hexagonal eight-layered perovskite structure with 3:1 ordering of tantalum or niobium with lithium. However, the molar fractions of Li/Na on the B site (25 mol%) in these compounds are determined by the stoichiometry of oxygen. A higher molar fraction of Li/Na on the Ta/Nb-site would lead to oxygen deficiency in the structure, which has not been reported to the best of our

knowledge. The BTLO perovskite reported in this work is thus the first example of simple cubic perovskite oxides with random/disordered occupancy of Ta and Li and high oxygen deficiency. Usually, for an A(B'B")O₃-type perovskite compound, the B cation centered octahedron is distorted if the B" cation is substituted by a B' cation. If there is an insignificant difference in the shape and size between two stable B'O6 and B"O₆, the distribution of B' and B" cations may be random. If the distortion of BO₆ is large enough after substituting B' for B", then B' and B" will prefer to be in a certain way in order to stabilize the perovskite structure. The distortion of BO₆ caused by substitution depends not only on the differences in ionic radii and charges of B' and B", but also on their polarizabilities, because the polarizability of an ion represents the deformability of the ion under the action of the neighboring ions.

To predict the order or disorder of the distribution of B' and B" in A(B'B")O₃-type perovskite compounds, some rules and methods have been proposed.30-33 Among which, a simple method proposed by Liu et al.30 has been proven to work well for the compounds with the ratio of B'/B'' = 1:1 and 1:2(or 2:1). In this method, an order-disorder phase diagram is drawn into three regions corresponding to ordered, disordered,

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and order-disorder transition regions. For this diagram, the coordinate Y is defined as $Y = \ln(100 \times |\Delta\alpha|/(N_A \times n_A))$, where $\Delta\alpha$ is the difference in the polarizabilities of B' and B" cations in $A(B'B'')O_3$ compounds, N_A and n_A are the largest principal quantum number and the number of total electrons in the outer electronic shells of the A atom, respectively. For the abscissa X, it is defined as $X = \ln(|Z_{B'}/r_{B'} - Z_{B''}/r_{B''}|)$ where $Z_{\rm B'}/r_{\rm B'}$ and $Z_{\rm B''}/r_{\rm B''}$ are the ratios of charges to ionic radii of B' and B" cations, respectively. According to this diagram, a compound with X > 1.8 would be located in the ordered region regardless of the Y value. Here in our work, the calculated X value for BaTa_{0.5}Li_{0.5}O_{2.5} is 1.87, greater than 1.8, and thus expected to result in an ordered distribution of Ta and Li ions in the structure, which is contrary to our experimental results. This might be caused by the high concentration of oxygen vacancy in the structure; as pointed out by Liu et al. 30 their method could lead to a contradictory conclusion if there is significant change in oxygen stoichiometry.

In addition, it is important to highlight that BTLO should be perceived as a cubic perovskite not Brownmillerite Ba₂TaLiO₅ due to the fact that Ta and Li are randomly distributed, and oxygen vacancies are disordered; the latter is the key for oxideion conduction. It is well known that highly acceptor-doped ABO₃ can transform into an orthorhombic A₂B₂O₅ brownmillerite structure, in which the two B cations have distinct positions and 1/6 of the O-site becomes vacant and ordered at lower temperatures. Above the characteristic temperature, the highly ordered orthorhombic brownmillerite structure can transform into a cubic perovskite structure by oxygen vacancies disordering, which is accompanied by an order of magnitude increase in oxide-ion conductivity such as that observed in Sr₂Fe₂O₅³⁴ and Ba₂In₂O₅, ^{35,36} just to name a few. Interestingly, a cubic oxygen disordered single crystal was reported to be synthesized for the Sr₂ScGaO₅ material, ²⁰ although the polycrystalline sample for this material also showed an orthorhombic ordered phase at ambient temperature. The cubic phase of this single crystal Sr₂ScGaO₅ material can be maintained until 1400 °C where it then transforms into an orthorhombic phase and is then transferred to the cubic phase again at 1600 °C. Here for our BTLO material, the VT-XRD study revealed that the

pure cubic phase of BTLO can be maintained at least to 900 $^{\circ}$ C (higher temperatures were not investigated due to the limitation for the instrument in our lab), as shown in Fig. 3(a). As can be seen, within the studied temperature region, no peaks from the secondary phase were detected, and the refined lattice parameters showed a linear increase with temperature (Fig. 3(b)), from which the thermal expansion coefficient was estimated to be $\sim 9.71 \times 10^{-6} \, \mathrm{K}^{-1}$. Therefore, it is expected that simple cubic perovskite BaTa_{0.5}Li_{0.5}O_{2.5} with disordered oxygen vacancies exhibits good oxide-ion conductivity over a broad temperature range.

3.2 Electrical properties

The conductivity of BTLO was measured using AC impedance spectroscopy. Fig. 4(a) shows an example of a Nyquist plot at 300 °C, which features a semicircle that can be modeled by a parallel R-C circuit in the high frequency range and Warburg impedance with characteristic capacitance higher than 10^{-7} F cm⁻¹ that can be modeled by semi-infinite ionic diffusion at the electrolyte/electrode interface in the low freguency range.³⁷ The intercept of the semicircular arc at low frequency represents the sum of resistance contributed from the grain and grain boundary $(R_g + R_{gb})$, from which the total conductivity of the material can be calculated out. At higher temperatures, e.g. 700 °C, the R-C semicircular arc becomes less pronounced due to reduced $R_g + R_{gb}$ and increased double layer capacitance, which makes the electrode response (R_e) to frequency dominate the spectrum, see Fig. 4(b). To preliminarily determine the nature of charge carriers responsible for the ionic conduction, impedance spectra were also measured under N2, air and O2 atmospheres with different oxygen partial pressures (pO_2) at 750 °C; the results are shown in Fig. 4(b). Two observations are noted: (1) the electrode response is very sensitive to pO_2 , *i.e.*, the higher the pO_2 , the lower the electrode resistance (R_e). The strong dependence of R_e on pO_2 infers the nature of active oxygen electrocatalysis $(O_2 + 4e^- \leftrightarrow 2O^{2-})$ at the BTLO/Pt interface, which is commonly observed in the oxide-ion conductors; 11,38 (2) $R_{\rm g}$ + $R_{\rm gb}$ is insensitive to $p{\rm O}_2$, implying pure ionic conduction. 11,38 Furthermore, Fig. 4(c) shows that the independence of $R_g + R_{gb}$ occurs over a wide

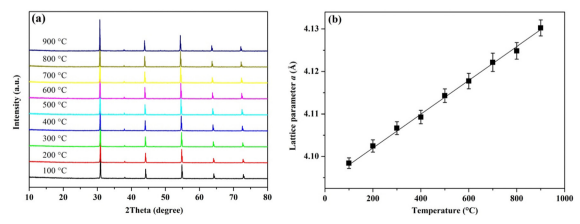


Fig. 3 (a) VT-XRD patterns and (b) corresponding lattice parameters of BTLO.

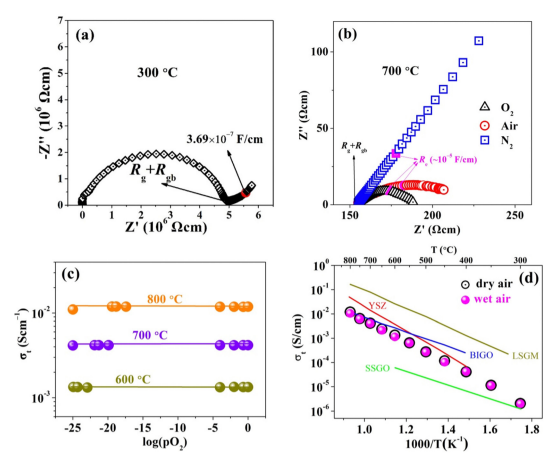


Fig. 4 (a) Nyquist impedance plot of BTLO at 300 °C; (b) impedance spectra under oxygen, air and nitrogen atmospheres at 750 °C; (c) pO₂ dependence of the total conductivity for BTLO; (d) Arrhenius plot of the total conductivity of BTLO in both dry and wet air, compared with yttria-stabilized zirconia (YSZ), ⁴⁴ gallium-doped cubic Ba₂In_{1.5}Ga_{0.5}O₅ (BIGO), ⁴³ cubic single crystal $SrSc_{0.5}Ga_{0.5}O_{2.5}$ (SSGO), ³⁸ and Sr- and Mg-doped cubic La_{0.8} $Sr_{0.2}$ -Ga_{0.83}Mg_{0.17}O_{2.815} (LSGM).¹⁰

pO₂ range from highly reducing to highly oxidizing atmospheres and a temperature range of 600–800 $^{\circ}\text{C}.$ It is evident that the total conductivity is invariable over $pO_2 = 1$ to 10^{-25} atm from 600 to 800 °C, showing no sign of n-type electronic conduction even at low pO2. It should be stressed here that the three low pO_2 at the 600 °C studied temperature were achieved by bubbling the 10%H₂-90Ar% gas, 5%H₂-95Ar% gas, and 1%H₂-99% Ar gas through room temperature water, respectively. This also means that the conductivities under wet atmospheres did not show obvious enhancements compared to that measured under dry gas (four different gases were used in this work: pure Ar with pO_2 of $\sim 10^{-4}$ atm; 1% O_2 -99%Ar gas with pO_2 of 10^{-2} atm; air gas with pO_2 of 0.21 atm; pure O_2 with pO_2 of 1.0 atm). Therefore, this strongly indicates that the proton conduction in the BTLO material, if any, is small. For the low pO₂ at 700 and 800 °C, they were achieved by mixing the dry CO and CO2 gases. Moreover, the conductivities of BTLO under both dry and wet air atmospheres from 300 to 800 °C were also closed to each other at a given temperature, as shown in Fig. 4(d), which again supported the conclusion that the proton conduction in BTLO can be negligible. This is unlike in many other oxygen deficient perovskite materials where both oxide ion conduction and proton conduction are usually

observed under wet atmospheres, such as in typical proton conducting Ba(Zr/Ce)O3-based materials39 and recently reported SrSc_{0.5-x}Zn_xGa_{0.5}O_{2.5-0.5}-based materials.²¹ As reported in Kreuer et al.'s work, 40 for the Y- and Sc-doped BaZrO3 materials, the former (Y-doped) have a lower activation energy and higher proton conductivity than that of the latter (Sc-doped), which was ascribed to the difference in the basicity of the oxygen ions coordinated to Y and Sc: the electron densities on the oxygen ions coordinated to Y are almost the same with the electron densities of oxygen coordinated to Zr, whereas the oxygen ions coordinated to Sc had significantly higher electron densities and therefore higher basicities. The more basic oxygen ions coordinated to Sc have therefore more strong bonds with protons, leading to lower proton mobility and higher activation energy. In the present work for the BaTa_{0.5}Li_{0.5}O_{2.5} material, almost half of the B-sites were occupied by the Li ions, and the oxygen ions coordinated to these Li ions were supposed to have much higher electron densities and basicity. Therefore, the protons bonding to these oxygen ions would have much lower mobility and higher activation energy. This may be a possible reason for the negligible proton conduction in the BaTa_{0.5}Li_{0.5}O_{2.5} material.

Fig. 4(d) also compares BTLO's total conductivity with YSZ, Ba₂In_{1.5}Ga_{0.5}O₅ (BIGO), and SrSc_{0.5}Ga_{0.5}O_{2.5} in an Arrhenius plot.

As shown, the conductivities of BTLO are comparable to YSZ at temperatures below 400 °C and close to BIGO at temperatures above 700 °C. However, the conductivities of BTLO here are apparently higher than that of the SrSc_{0.5}Ga_{0.5}O_{2.5} material which has the same oxygen defect level and cubic perovskite structure. As Sammells et al. 41,42 suggested, the activation energies of oxide ions in perovskite oxides have a strong relationship with the radius of the saddle point (bottleneck) which was formed by two A-site cations and one B-site cation in the perovskite structure. The radius of this kind of saddle point $(r_{critical})$ can be calculated according to the following equation

$$r_{\text{critical}} = \frac{-r_{\text{A}}^2 + \frac{3}{4}a^2 - \sqrt{2}ar_{\text{B}} + r_{\text{B}}^2}{2r_{\text{A}} + \sqrt{2}a - 2r_{\text{B}}}$$
(1)

where r_A and r_B are the Shanon's ionic radius of the A-site cation and B-site cation, respectively, and a is the lattice parameter. Usually, a larger saddle point radius is expected to result in a lower activation energy and high mobility of oxide ion. As reported in Yao et al.'s work⁴³ for the Ga-doped cubic Ba₂(In_{1-x}Ga_x)₂O₅ (x = 0.25, 0.3, 0.35, 0.4, 0.45), the $r_{critical}$ decreased with Ga-content from 0.887 for x = 0.25 to 0.882 for x = 0.45 and resulted in a decrease in conductivity. According to the above equation, the $r_{\rm critical}$ calculated for BTLO here is ~ 0.872 which is slightly lower than that (0.887) of Ba₂(In_{0.75}Ga_{0.25})₂O₅, but obviously higher than that (0.818) of SSGO. Given the almost identical concentration of oxygen vacancy in these three materials, the radius of the saddle point seems to be the main factor that affects the activation energy and mobility of oxide ions and therefore the oxide ion conductivity. This may the reason (structural origins) here why the conductivity of BTLO is lower than BIGO but apparently higher than SSGO.

In addition, we can see in Fig. 4(d) that the conductivity of BTLO is more than one order of magnitude lower than that of the Sr and Mg co-doped LaGaO3 (LSGM) with the composition of La_{0.8}Sr_{0.2}Ga_{0.83}Mg_{0.17}. As has been reported for LSGM materials, 10 that their conductivities increase with the increase of Sr/Mg concentration at first, after reaching a maximum, and the oxide ion conductivities of LSGM decrease with the increase

of Sr/Mg concentration (or oxygen vacancy concentration). The composition of La_{0.8}Sr_{0.2}Ga_{0.83}Mg_{0.17}O_{2.815} was reported to exhibit the highest conductivity of 0.166 S cm⁻¹ at 800 °C. Moreover, the activation energy of oxide ion conduction in LSGM was reported to increase with the oxygen vacancy concentration. Here for the BTLO, although it has apparently higher oxygen vacancy concentration than that in LSGM, the oxide ion conduction was significantly lower than that of LSGM. Here, as the radius of the saddle point for oxide ion migration in $La_{0.8}Sr_{0.2}Ga_{0.83}Mg_{0.17}O_{2.815}$ was calculated to be ~0.948, which is apparently higher than that of BTLO (0.872), and thus was expected from this point of view to have a much higher conductivity than BTLO.

Based on the above results, we can draw a general conclusion that the oxide-ion conduction is likely dominating in BTLO with negligible electronic conductivity, although the possibility for lithium-ion conduction could not be ruled out yet. The dominant oxide-ion conduction is further verified by the electromotive force (EMF) measurement using an oxygen concentration cell consisting of air BTLO O2, which shows an EMF value of ~ 33.5 mV at 750 °C, which is 97% of the theoretical EMF calculated from the Nernst equation. To further obtain partial oxide-ion conductivity, we employed a modified fourprobe Hebb-Wagner cell, with which the Li⁺ ion conductivity, if any, can be obtained by subtracting the partial oxide-ion conductivity from the total conductivity. In this modified four-probe Hebb-Wagner cell, the La_{1.54}Sr_{0.46}Ga₃O_{7.27} (denoted as LSGO in the following) material, which has been welldocumented to be a pure oxide ion conductor, 38 was used as an electron as well as a Li+ blocking layer. Besides two reversible porous Pt electrodes, there is an additional porous Pt layer situated between BTLO and LSGO. The cell configuration in this work can thus be described as (+) Pt||LSGO/Pt/BTLO||Pt (-); a schematic of this cell is shown in Fig. 5(a). The additional porous Pt layer between BTLO and LSGO was applied to mainly overcome the high BTLO/LSGO interfacial resistance, which is usually observed in a traditional Hebb-Wagner cell. The cell was placed in a quartz tube furnace filled with pure O2.

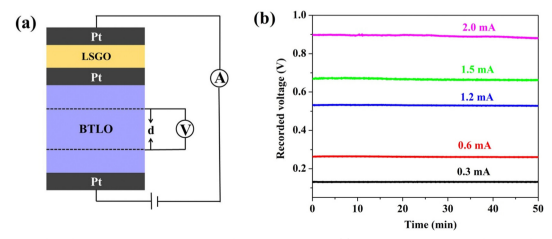


Fig. 5 (a) Schematic of the four-probe Hebb-Wagner cell configuration in this work; (b) the recorded voltages as a function of time using various currents

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Table 2 Oxygen transport number of BTLO at 800 °C

I (mA)	$\sigma_{ m o}$	σ_{t}	$t_{ m o}$
0.3	0.01172	0.01195	0.981
0.6	0.01168	0.01195	0.977
1.2	0.01151	0.01195	0.963
1.5	0.01144	0.01195	0.957
2.0	0.01132	0.01195	0.947

The measurements were carried out at 800 °C to ensure a sufficient rate of the oxygen redox reactions (oxidation/ reduction) on the porous Pt electrodes. Under a constant current (I), the voltage drop (V) across a given distance of the specimen was measured. Oxide-ion conductivity (σ_0) is then calculated by Ohm's law with the known sample dimensions. It is important to mention that the electrical measurements by the Hebb-Wagner polarization method should be carried out under steady-state conditions, see Fig. 5(b), where only the oxide-ion current should be observed in BTLO. The constant current I was kept in a range of 0.3-2.0 mA in the present study to ensure a steady-state voltage. The results are summarized in Table 2 and compared with the total conductivities obtained using an AC impedance technique. The results reveal clearly that the oxide-ion conduction dominates the conductivities of BTLO, with an average oxide-ion transport number (t_0) of ~ 0.965 , suggesting almost negligible electronic and lithiumion conductions.

Thus, it is safe to say that the measured total conductivity of BTLO under air conditions shown in Fig. 4(d) is largely attributed to oxide-ion conduction. The same magnitude of ionic conductivity in BTLO as that of the conventional YSZ and BIGO implies the same level of mobile oxygen vacancy concentration. Given the fact that the oxide-ion conductivity is independent of pO2, BTLO is a new electrolyte candidate for SOFCs. Like LSGM or other perovskite oxide-ion conductors, one distinct advantage of BTLO is the direct use of highly active perovskite cathodes without a barrier layer such as Gd-doped CeO2 that must be used in ZrO2-based SOFCs.

3.3 Discussion on the oxide-ion conduction mechanism

As have been reported for other perovskite materials, such as LaBO₃ (B = Cr, Mn, Fe, Co), 45 La_{0.6}Sr_{0.4}CoO_{3- δ}, 46 (La_{0.8}Sr_{0.2})- $(Ga_{0.8}Mg_{0.15}Co_{0.05})O_{2.8}$, and so on, the oxygen mobility is mainly mediated by hopping of oxygen vacancies along the octahedral edge of BO6. In the present work, we performed AIMD simulations on ionic diffusion. The diffusivities at three selected temperatures (2000 K, 1800 K and 1600 K) are obtained from the mean square displacements (MSD) as shown in Fig. S1a (ESI†). Fig. S1b-d (ESI†) present the MSD for Ba, Ta, and Li ions, respectively. One can clearly see that the Ta and Ba ions are not mobile at all the simulated temperatures (1600 K, 1800 K, and 2000 K). While for the Li ions, the MSD values (below 1 Å² at 300 ps) at 1600 and 1800 K are significantly lower than that of the oxygen ions ($\sim 14 \text{ Å}^2$ and $\sim 27 \text{ Å}^2$ at 1600 and

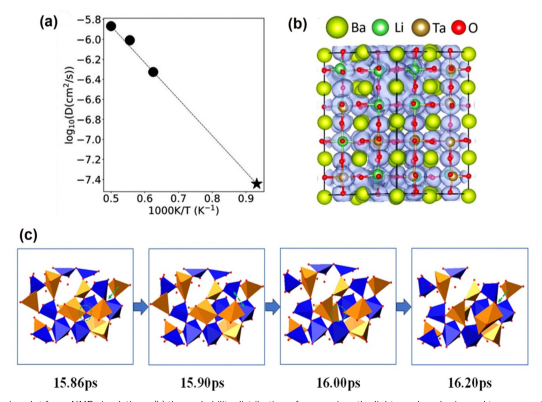


Fig. 6 (a) Arrhenius plot from AIMD simulations; (b) the probability distribution of oxygen ion; the light purple color is used to represent the probability isosurface of oxygen ions. (c) Selected snapshots of MD simulation for oxygen vacancy migration in (Ta/Li)On polyhedral; the blue and orange polyhedral denotes the TaO_n and LiO_n respectively

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1800 K, respectively) at 300 ps. However, at a higher simulated temperature of 2000 K, the Li ions seem to have an even larger MSD value than that of oxygen ions at the same temperature. This may suggest that at a sufficiently high temperature, the Li ions in the BTLO would have a long-range displacement and then contribute to the ionic conductivity. However, in our work the experimentally investigated temperatures are believed to be lower than this trigger temperature for Li ions' long-range migration.

Furthermore, the simulation data at these three temperatures fit the Arrhenius relationship well as shown in Fig. 6(a). Extrapolated to 800 °C, the ionic conductivity at 800 °C is $3.014 \times 10^{-2} \text{ S cm}^{-1}$ with an activation energy of 0.734 eV, which are close to our experimental data. Fig. 6(b) clearly displays three-dimensional migrating pathways, while Fig. 6(c) shows some selected snapshots for demonstrating the role of the (Ta/Li)O_n polyhedral for oxygen transport, which is akin to that observed in other simple cubic perovskite oxideion conductors. 48 More specifically, the simulations revealed that Ta ions are mainly coordinated by 5 or 6 oxygen atoms. Whereas the Li ions are coordinated by 3 or 4 oxygen atoms within most of the simulated time, sporadically 5, 6 and even 2 oxygen coordinated Li ions were also observed.

4. Conclusion

In this work, we report a new simple cubic perovskite material BaTa_{0.5}Li_{0.5}O_{2.5} with disordered oxygen vacancies. A high and pure oxide ion conductivity of $\sim 1.2 \times 10^{-2} \, \mathrm{S \ cm^{-1}}$ at 800 °C is observed, which is comparable to that of the widely used YSZ material. In addition, the oxide-ion conduction mechanism in BaTa_{0.5}Li_{0.5}O_{2.5} is supported by ab initio molecular dynamic simulations, revealing 3D migration pathways for oxygen vacancies. This work may inspire future work to discover new oxideion conductors in similar systems.

Author contributions

Jungu Xu, conceptualization, data curation, funding acquisition, writing - review & editing; Qin Cao, investigation, data curation; writing - original draft; Lin Wang, formal analysis; Bin Ouyang, formal analysis, funding acquisition, writing review & editing; Tianjie Wei, investigation, data curation; Jiazheng Hao, investigation, data curation; funding acquisition; Jie Chen, investigation, data curation; Lunhua He, supervision, funding acquisition, writing - review & editing; Laijun Liu, resources, writing - review & editing; Kevin Huang, supervision, writing - review & editing.

Conflicts of interest

There are no conflicts of interest to declare.

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