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Heterovalent and isovalent doping of bilayer protonconducting perovskite SrLa₂Sc₂O₇

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Perovskite or perovskite-related structural materials are widely studied for their many functional properties. They can be used as components of energy sources such as solid oxide fuel cells. Along with classical perovskites, layered perovskites can also carry out high-temperature proton transport and are promising materials for use in electrochemical power engineering. In this paper, the possibility of heterovalent and isovalent doping of La and Sc sublattices of bilayer perovskite SrLa₂Sc₂O₇ was made for the first time. It was shown that electrical conductivity increases in the row of bilayer perovskites SrLa₂Sc₁O₇ – SrLa₂Sc₂O₇ – BaLa₂In₂O₇ – BaNd₂In₂O₇.

keywords: layered perovskite; oxygen-ion conductivity; proton conductivity; hydrogen energy; Ruddlesden-Popper structure

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1. Introduction

Perovskite or perovskite-related structural materials are widely studied for their many functional properties. Some of these applications are solar cells [1-4], lightemitting diodes [5], semiconductor lasers [7], photovoltaic [8, 9] and optoelectronic applications [10], and solid oxide fuel cells and electrolyzers [11–15]. Solid oxide fuel cells are energy sources. They work by converting energy from chemical reactions into electricity. This is the field of hydrogen energy. It is very promising, both in terms of energy resource availability and the environment [16–18]. A solid oxide fuel cell consists of several components, including electrolyte, electrodes, interconnector and glass sealant. The electrolyte material can be an oxygen ion conductor or a proton conductor. Doped barium zirconate is the most studied electrolyte material for proton conducting solid oxide fuel cells [19]. It crystallises

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in the classic perovskite ABO₃ structure. However, there are a number of problems with the use of these materials, including low grain boundary conductivity, poor sinterability and low oxygen vacancy concentrations created by an accepter-doping mechanism [20–24]. Therefore, the search for new proton-conducting materials is necessary.

The ability of layered perovskites $AA'_nB_nO_{3n+1}$ to transport oxygen ions [25–32] and protons [33–38] was demonstrated a few years ago. The monolayer perovskites $AA'BO_4$ such as BaLalnO_4, SrLalnO_4, BaNdlnO_4, have a crystal lattice capable of doping both Ba/La and In sublattices [39]. The bilayer perovskites $AA'_2B_2O_7$ such as BaLa₂ln₂O₇ and BaNd₂ln₂O₇ allow predominantly doping only in the A or A' sublattices [40–43] (Figure I). Last year, the possibility of oxygen ion and proton transport was proved for the bilayer perovskite SrLa₂Sc₂O₇ [44]. This paper first explores the possibility of heterovalent and isovalent doping of La and Sc sublattices.

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Figure 1 The crystal structure of a bilayer perovskite with the general formula $AA'_2B_2O_7$ (a) and the scheme of doping of layered perovskites $BaLa_2In_2O_7$ and $BaNd_2In_2O_7$ (b).

2. Experimental

The compositions $SrLa_2Sc_2O_7$, $SrLa_{1.9}Ca_{0.1}Sc_2O_{6.95}$, $SrLa_{1.9}Sr_{0.1}Sc_2O_{6.95}$, $SrLa_{1.9}Ba_{0.1}Sc_2O_{6.95}$, $SrLa_2Sc_{1.9}In_{0.1}O_7$, $SrLa_2Sc_{1.9}Gd_{0.1}O_7$, $SrLa_2Sc_{1.9}Y_{0.1}O_7$, $SrLa_2Sc_{1.9}Ti_{0.1}O_7.05$ and $SrLa_2Sc_{1.9}Gd_{0.1}O_7$ were prepared by the solid state method. The powders of the starting reagents $BaCO_3$, $SrCO_3$, $CaCO_3$, Sc_2O_3 , La_2O_3 , In_2O_3 , Y_2O_3 , Gd_2O_3 , TiO_2 (all 99.99 % purity, REACHIM, Russia) were dried and used in stoichiometric amounts. An agate mortar was used for grinding. The compositions were heated after each grinding. The annealing was carried out in the temperature range of 80O - 130O °C with 10O °C step.

phase identification of The the obtained compositions was carried out using the Bruker Advance D8 Cu $K\alpha$ diffractometer. The thermogravimetry (TG) was made using STA 409 PC Netzsch Analyser. The heating of initially hydrated samples was made at the temperature range of 40 - 1100 °C with the rate of 10 °C/min under a flow of dry Ar. The preliminary hydrated samples were obtained by slow cooling from 1000 to 150 °C (1 °C/min) under a flow of wet Ar (99.999 % purity, $pH_2O = 2 \cdot 10^{-2}$ atm). The Ar atmosphere was used to avoid any carbonization of the sample.

The electrical conductivity was measured with an impedance spectrometer Z-1000P, Elins, RF. The investigations were carried out from 1000 to 200 °C with a cooling rate of 1 °/min under dry air or dry Ar. The dry gas (air or Ar) was prepared by circulating the gas through P_2O_5 ($pH_2O = 3.5 \cdot 10^{-5}$ atm). The wet gas (air or Ar) was obtained by bubbling the gas at room temperature first through distilled water and then

through a saturated solution of KBr $(pH_2O = 2 \cdot 10^{-2} \text{ atm})$.

3. Results and discussions

In this work, the heterovalent (acceptor) doping of the lanthanum sublattice $SrLa_{1.9}M_{0.1}Sc_2O_{6.95}$ (M = Ca, Sr, Ba) and the heterovalent (donor) doping SrLa2Sc1.9Tio.1O7.05 and the isovalent doping $SrLa_2Sc_{1.9}M'_{0.1}O_7$ (M' = In, Gd, Y) and $SrLa_2ScInO_7$ of the scandium sublattice were provided. The XRD-analysis showed that only two compositions SrLa₂Sc₂O₇ and SrLa₂SclnO₇ were obtained as single phases. Figure 2 represents the analysis of XRD-data for SrLa₂ScInO₇ composition. Both undoped SrLa₂ScInO₇ and doped SrLa₂ScInO₇ compositions are indexed in the ortorombic symmetry (space group *Fmmm*). The lattice parameters are presented in Table 1.



Figure 2 XRD-patterns for the composition SrLa₂SclnO₇.

Table 1 – Unit cell parameters and volume of SrLa₂SclnO₇ and SrLa₂SclnO₇ compositions.

Composition	a, Å	<i>b</i> , Å	ĢÅ	V _{cell} , Å ³
SrLa ₂ Sc ₂ O ₇	5.7811	5.7381	20.5342	681.171
SrLa ₂ ScInO ₇	5.8172	5.8139	20.6772	699.345

 Table 2 – Phase composition of obtained samples.

Nominal composition	Phase composition		
SrLa1.9Ca0.1Sc2O6.95	SrLa1.9Cao.1Sc2O6.95, La2O3		
SrLa1.9Sr0.1Sc2O6.95	SrLa1.9Sr0.1Sc2O6.95, La2O3		
SrLa _{1.9} Ba _{0.1} Sc ₂ O _{6.95}	SrLa _{1.9} Ba _{0.1} Sc ₂ O _{6.95} , La ₂ O ₃		
SrLa2Sc1.9Ino.1O7	SrLa2Sc1.9In0.1O7, La2O3		
SrLa2Sc1.9Gd0.1O7	SrLa2Sc1.9Gd0.1O7, La2O3		
SrLa ₂ Sc _{1.9} Y _{0.1} O ₇	SrLa ₂ Sc _{1.9} Y _{0.1} O ₇ , La ₂ O ₃		
SrLa2Sc1.9Tio.1O7.05	SrLa2Sc1.9Tio.1O7.05, LaScO3, La2O3, Sc2O3		

As can be seen, the introduction of indium ions with bigger ionic radius ($r_{Sc^{3+}} = 0.745$ Å, $r_{In^{3+}} = 0.80$ Å [45]) leads to the increase in the lattice parameters and unit cell volume. The phase composition of other samples is presented in Table 2.

The graphical representation for the samples with nominal compositions SrLa₂Sc_{1.9}Gd_{0.1}O₇ and SrLa₂Sc_{1.9}Tio.1O7.05</sub> is shown in Figure 3. As previously shown for the bilayer perovskites BaLa₂In₂O₇ and BaNd₂In₂O₇ [42], the donor doping of the indium sublattice is almost non-existent, in contrast to monolayer perovskites. The crystal lattice of bilayer perovskite contains a linkage of two layers of octahedrons in one perovskite layer and is more rigid compared to the monolayer perovskite layered structure where the layers of the perovskite block are one layer of octahedrons. Accordingly, donor doping, which involves the formation of interstitial oxygen in the interoctahedral space, is not realized. It is obvious that the same reason is an explanation for non-single phase of donor-doped compositions based on SrLa₂Sc₂O₇. At the same time, the lattice parameters and unit cell volume for SrLa₂Sc₂O₇ composition are much smaller than for BaLa₂In₂O₇ and BaNd₂In₂O₇ compositions (Table 3). We can assume that this decrease can prevent acceptor doping in the lanthanum sublattice, which is inevitably accompanied by a change in the local structure and the formation of point defects.



Figure 3 The phase composition (weight ratio) for the samples "SrLa₂Sc_{1.9}Gd_{0.1}O7" (1 - SrLa₂Sc_{1.9}Gd_{0.1}O7, 2 - La₂O₃) and "SrLa₂Sc_{1.9}Ti_{0.1}O7.05" (1 - SrLa₂Sc_{1.9}Ti_{0.1}O7.05, 2 - LaScO₃, 3 - La₂O₃, 4 - Sc₂O₃).

Table 3 – The lattice parameters and unit cell volumes of bilayer perovskites AA'₂B₂O₇.

Composition	<i>a,</i> Å	<i>b</i> , Å	ςÅ	V_{cell} , Å ³
SrLa2Sc2O7	5.7811	5.7381	20.5342	681.171
BaLa ₂ In ₂ O ₇	5.9149	5.9149	20.8465	729.336
BaNd2In2O7	5.8916	5.8916	20.4960	710.520

The possibility of interaction with water vapours was investigated using thermogravimetry (TG) and mass spectrometry (MS) measurements. Mass loss occurs at the temperatures below 500 °C (TG curve) and is solely due to the release of water (MS(H₂O) curve). The water uptake for composition SrLa₂ScInO₇ is about 0.06 mol H₂O per mol complex oxide (Figure 4), which is slightly more than for composition SrLa₂Sc₂O₇ (0.05 mol H₂O [44]). The compositions $BaLa_2In_2O_7$ and $BaNd_2In_2O_7$ are characterised by the water uptake 0.17 and 0.15 mol H₂O per mol complex oxide correspondingly. For the layered perovskites, the possibility of water uptake is due to the presence of enough space between perovskite blocks and rock-salt layers [40]. The increase in the size of this space leads to the increase in the water uptake. Accordingly, the changes in unit cell volume and the changes in this space should lead to changes in water uptake in the same way.

The temperature dependencies of the conductivity values for the compositions $SrLa_2SclnO_7$ and $SrLa_2Sc_2O_7$ under dry (filled symbols) and wet (open symbols) air are presented in Figure 5.



Figure 4 TG- and MS (H₂O)-curves for the hydrated composition SrLa₂ScInO₇.



Figure 5 Temperature dependencies of conductivity for the compositions SrLa₂SclnO₇ (blue symbols) and SrLa₂Sc₂O₇ (red symbols) under dry (filled symbols) and wet (open symbols) air.



Figure 6 Temperature dependencies of conductivity under dry air for the compositions SrLa₂SclnO₇ (1), SrLa₂Sc₂O₇ (2), BaLa₂ln₂O₇ [41] (3), BaNd₂ln₂O₇ [43] (4).

As can be seen, the electrical conductivity for the composition with two different ions in the B-sublattice of the bilayer perovskite AA'₂B₂O₇ (SrLa₂ScInO₇) is lower than for the compositions with one type of ions (SrLa₂Sc₂O₇). We can assume that the presence of ions of different nature in the same sublattice can lead to a local distortion of the crystal lattice, which in turn can prevent the transfer of ions despite an increase in the lattice parameters. It should be noted that difference in electrical conductivity values decreases with decreasing temperature. The values of activation energy (0.9 eV for SrLa₂Sc₂O₇ and 0.8 eV for SrLa₂ScInO₇) correlate well with this. The effect of the air humidity starts lower 500 °C which agrees well with TG-data. This allows to say that the proton conductivity contribution occurs in a humid atmosphere at low temperatures. However, the conductivity values are not high. Among bilayer perovskites AA'₂B₂O₇ electrical conductivity values increase in the row SrLa₂ScInO₇ - SrLa₂Sc₂O₇ - BaLa₂In₂O₇ - BaNd₂In₂O₇ (Figure 6).

Thus, it can be said that the bilayer perovskite $SrLa_2Sc_2O_7$ is not a suitable matrix to modify and obtain new promising proton-conducting materials.

4. Conclusions

In this paper, the possibility of heterovalent and isovalent doping of La and Sc sublattices of bilayer perovskite SrLa₂Sc₂O₇ was made for the first time. It was shown that both heterovalent (acceptor and donor) and isovalent doping of the indium sublattice is almost nonexistent. The compositions $SrLa_{1.9}M_{0.1}Sc_2O_{6.95}$ (M = Ca, Sr, Ba), SrLa₂Sc_{1.9}M'_{0.1}O₇ (M' = In,Gd, Y) and SrLa₂Sc_{1.9}Tio.1O_{7.05} were non-single phase. The possibility of interaction with water vapours and electrical conductivity were investigated for the single phase composition SrLa₂ScInO₇. It was shown that composition SrLa₂SclnO₇ introduces a small amount of water (0.06 mol H₂O per mol complex oxide). The proton conductivity contribution occurs in a humid atmosphere at low temperatures. The conductivity values for composition SrLa₂SclnO₇ are not high. They increase in the row SrLa₂SclnO₇ – SrLa₂Sc₂O₇ – BaLa₂ln₂O₇ – BaNd₂ln₂O₇. The bilayer perovskite SrLa₂Sc₂O₇ is not a suitable matrix to modify and obtain new promising proton-conducting materials.

Supplementary materials

No supplementary materials are available.

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None.

Author contributions

Nataliia Tarasova: Conceptualization; Data curation; Writing – original draft; Writing – review & editing.

Conflict of interest

The authors declare no conflict of interest.

Additional information

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