Thermomechanical, transport and anodic properties of perovskite-type $(\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_1-x\text{Fe}_x\text{O}_3-\delta$

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**A B S T R A C T**

Effects of iron doping on the functional properties of $(\text{La},\text{Sr})\text{CrO}_3$-based electrode materials have been appraised in a range of conditions relevant for SOFCs and other electrochemical applications. Mössbauer spectroscopy of perovskite-type $(\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_1-x\text{Fe}_x\text{O}_3-\delta$ $(x=0.3–0.4)$, combined with thermogravimetry and X-ray diffraction, shows that the prevailing oxidation state of iron cations in both oxidizing and reducing atmospheres remains $\delta$. The redox behavior and transport properties are, therefore, essentially governed by $\text{Cr}^{3+/4+}$ couple, leading to dominant p-type electronic conduction in the oxygen partial pressure range from $0.5$ down to $10^{-20}$ atm at $973$–$1223$ K. The total conductivity and Seebeck coefficient variations indicate that the electronic transport decreases with iron additions as the hole concentration and mobility become lower. The partial ionic conductivity estimated from the steady-state oxygen permeability under air/H$_2$–H$_2$O–N$_2$ gradients is $0.05–0.085$ S cm$^{-1}$ at $1223$ K and $p(\text{O}_2)=10^{-17}$ atm, and increases with $x$ due to rising oxygen deficiency. A similar trend was observed for chemical expansion studied by the controlled-atmosphere dilatometry, whereas the average thermal expansion coefficients are almost independent of $x$ and vary from $11.1–11.3 \times 10^{-6}$ K$^{-1}$ in air down to $10.3–10.5 \times 10^{-6}$ K$^{-1}$ in CO–CO$_2$ at $350$–$1370$ K. The electrochemical activity of porous $(\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_1-x\text{Fe}_x\text{O}_3-\delta$ anodes with $\text{Ce}_0.8\text{Gd}_{0.2}\text{O}_2-\delta$ interlayers, applied onto LaGaO$_3$-based solid electrolyte, is higher compared to $(\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_1-x\text{Mn}_0.5\text{O}_3-\delta$ when no metallic current-carrying layers are used. Increasing iron content lowers the electrode performance in wet H$_2$-containing atmospheres, in correlation with electronic conduction.

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1. Introduction

Energy-related systems based on solid oxide fuel cells (SOFCs) and electrolysis cells (SOECs) offer important potential advantages due to a high efficiency, environmental safety, fuel flexibility including the prospects to directly utilize hydrocarbons and gasified coal, and possibility to recover exhaust heat [1–7]. Significant efforts are focused on the developments of oxide electrode materials stable in both oxidizing and reducing environments, in order to reduce costs of the electrochemical devices and to suppress degradation phenomena, such as coking and sulfur poisoning of the electrodes operating in fuel atmospheres. These problems are well known for the conventional SOFC anodes of Ni-containing cermets, which also degrade due to large volume changes on redox cycling and to sintering during SOFC operation [3–9]. At the same time, alternative oxide electrode compositions based on perovskite-related titanates and chromates stable in relatively wide ranges of the oxygen partial pressures, $p(\text{O}_2)$, possess usually a number of other functional disadvantages, primarily a worse electrochemical activity and lower electronic transport compared to the cermets. A very promising combination of properties was recently reported for perovskite-type $(\text{La}_{0.75}\text{Sr}_{0.25})_{1-\gamma}\text{Cr}_{1.05}\text{Mn}_{0.5}\text{O}_3-\delta$ $(\gamma=0–0.05)$ [3,4,10–12]. In these perovskites (LSCM), the presence of rigid CrO$_6$ octahedra in the crystal structure enables its stabilization down to low oxygen chemical potentials necessary for the SOFC anode operation, while moderate acceptor-type doping is necessary to improve electronic and ionic conduction. Nevertheless, the electrochemical performance of LSCM anodes without any additional components is essentially limited by the electronic transport, leading to rather high overpotentials; the incorporation of metals (e.g., Ni or Cu) and/or electrocatalytically active additives (e.g., CeO$_{2-\delta}$) decreases
electrode resistance [10–12]. Notice that introducing a ceria-based buffer layer between lanthanum gallate-based electrolyte and La$_{0.75}$Sr$_{0.25}$Cr$_{0.5}$Mn$_{0.5}$O$_3$–δ electrode enables also to suppress diffusion of Cr and Mn cations into the solid–electrolyte surface layers [11].

Doping of acceptor-substituted LaCrO$_3$–δ with iron may further enhance electrode performance [4,5]. For example, a lower polarization resistance with respect to LSCM was reported for the electrodes made of its Fe-containing analog, La$_{0.75}$Sr$_{0.25}$Cr$_{0.5}$Fe$_{0.5}$O$_3$–δ [4]. The electrochemical cells with La$_{0.75}$Sr$_{0.25}$Cr$_{0.5}$Fe$_{0.5}$O$_3$–δ electrodes and (La,Sr)Ga$_2$MgO$_4$ (LSGM) solid electrolyte exhibit however a higher series resistance, presumably due to stronger reactivity of the materials in comparison with LSCM/LSGM cells [4]. An excellent performance was observed for the composite anodes made of perovskite-type LaSr$_2$Fe$_2$CrO$_8$ and gadolinia-doped ceria (CGO), with an additional CGO interlayer applied between the electrode and LSGM electrolyte [5]. On the other hand, literature data on phase stability of (La,Sr)Cr$_x$Fe$_{1-x}$O$_{3–δ}$ in reducing atmospheres are still quite contradictory [3–6,13], whereas information on thermomechanical and transport properties of these perovskites is very scarce.

The present work is focused on the evaluation of anode behavior and other functional properties of (La$_{0.75}$Sr$_{0.25}$)$_{1-x}$Fe$_x$O$_{3–δ}$ (x = 0.3–0.4). The concentration of iron cations in this model system was selected lower than that in La$_{0.75}$Sr$_{0.25}$Cr$_{0.5}$Fe$_{0.5}$O$_3$–δ [4] and LaSr$_2$Fe$_2$CrO$_8$–δ [5], thus making it possible to expect a moderate enlargement of the perovskite phase stability domain under reducing conditions due to higher fraction of iron–oxygen polyhedra surrounded exclusively by the CrO$_6$ octahedra. Emphasis is given to the properties important for practical application, namely thermal and chemical expansion, electronic conductivity, oxygen permeability, and electrode behavior. In order to assess phase stability and electronic transport mechanisms, Mössbauer spectroscopy and Seebeck coefficient measurements were also employed. The anodic overpotentials of porous (La$_{0.75}$Sr$_{0.25}$)$_{0.95}$Cr$_{0.5}$Fe$_{0.5}$O$_3$–δ electrodes applied onto LSGM with CGO interlayer are compared to those of LSCM and (La,Sr)/(Mn,Ti)O$_{2–δ}$, which were studied under similar conditions and partly reported elsewhere [12,14].

2. Experimental

Single-phase powders of (La$_{0.75}$Sr$_{0.25}$)$_{1-x}$Fe$_x$O$_{3–δ}$ were prepared by the glycine–nitrate process, a self-combustion technique utilizing nitrate of metal components as an oxidant and glycine as a fuel and chelating agent [15]. The obtained foam-like powders were annealed in air at 1273 K for 8 h in order to remove organic residuals, and then grinded. Gas-tight ceramic samples for the measurements of total conductivity, thermal expansion and oxygen permeability were uniaxially compacted at 50–150 MPa and sintered in air at 1973 K for 5 h, with subsequent slow cooling (2–4 K min$^{-1}$) to achieve equilibrium with atmospheric oxygen at low temperatures. The density of ceramic materials was in the range 92–95% of their theoretical density calculated from X-ray diffraction (XRD) results, Table 1. The powdered samples used for XRD, Mössbauer spectroscopy and thermogravimetric analysis (TGA), were obtained by grinding of sintered ceramics. Additional series of samples used for the studies of phase composition and iron states at elevated temperatures, were also annealed at 1223 K in air and in a flowing H$_2$–H$_2$O–N$_2$ mixture with P(O$_2$) = 10$^{-18}$ atm for 44 h, and then quenched.

Materials characterization included XRD (Rigaku D/Max-B and PAAnalytical X’Pert Pro diffractometers), scanning electron microscopy coupled with energy dispersive spectroscopy (SEM/EDS, Hitachi S-4100/UL-70), controlled-atmosphere dilatometry (Linseis L70/2001), TGA (Setaram SetSys 16/18), inductively coupled plasma (ICP) spectrometric analysis (Jobin Yvon 70 plus), measurements of total conductivity (four-probe DC) and Seebeck coefficient as a function of temperature and oxygen partial pressure, and determination of steady-state oxygen permeation fluxes under air/H$_2$–H$_2$O–N$_2$ gradients. Detailed description of the experimental procedures and equipment was published elsewhere [12,14,16–21]. For the measurements of electrical properties, the criteria for equilibration of a sample after a change in either oxygen activity or temperature included the relaxation rates of the conductivity and Seebeck coefficients less than 0.04% min$^{-1}$ and 0.001 μV K$^{-1}$ min$^{-1}$, respectively; the equilibration times varied from 2–4 h up to 80 h.

Mössbauer transmission spectra were collected at room temperature and 4 K using a conventional constant-acceleration spectrometer and a 25 mCi $^{57}$Co source in a Rh matrix. The velocity scale was calibrated using α-Fe foil. The absorbers were obtained by pressing the powdered samples into perspex holders. Isomer shifts (IS, Table 2) are related to metallic α-Fe at room temperature. The 4 K spectra were collected using a JANIS bath cryostat, model SIV-400, with the sample immersed in liquid He. The spectra were fitted to Lorentzian lines using a non-linear least-squares method [20]. In the course of refinement procedure, the relative areas and line widths of both peaks in a quadrupole doublet and of the symmetrical peaks constituting a magnetic sextet (i.e., peaks 1–6, 2–5 and 3–4) were constrained to remain equal. The distribution of magnetic splittings was fitted according to the histogram method [22].

The steady-state anode polarization experiments were performed by the 3-electrode technique using the electrochemical cells comprising symmetrical working (WE) and counter (CE) electrodes made of porous (La,Sr)(Cr,Fe)O$_{3–δ}$ and Pt, respectively, Ce$_{0.8}$Ga$_{0.2}$O$_{2–δ}$ interlayers, (La$_{0.9}$Sr$_{0.1}$)$_{0.98}$Ga$_{0.8}$Mg$_{0.2}$O$_{3–δ}$ solid electrolyte membranes, porous Pt reference electrodes (REs), and Pt gauze current collectors. The cell configuration, fabrication conditions and measuring procedures were all identical to those described in Ref. [12]: processing and functional properties of (La$_{0.9}$Sr$_{0.1}$)$_{0.98}$Ga$_{0.8}$Mg$_{0.2}$O$_{3–δ}$ (LSGM) electrolyte ceramics were reported in Ref. [19]. Prior to the WE deposition, as-prepared (La$_{0.75}$Sr$_{0.25}$)$_{1-x}$Fe$_x$O$_{3–δ}$ powders were annealed in air at 1273 K.
Table 2 Parameters\textsuperscript{*} estimated from the Mössbauer spectra of (La\textsubscript{0.75}Sr\textsubscript{0.25})\textsubscript{0.95}Cr\textsubscript{1−x}Fe\textsubscript{x}O\textsubscript{3−δ} equilibrated in air and in reduced atmosphere.

<table>
<thead>
<tr>
<th>(x)</th>
<th>Equilibration conditions</th>
<th>CN</th>
<th>Measurement temperature, K</th>
<th>(I) ((\text{mm} \text{s}^{-1}))</th>
<th>(QS, 2\nu) ((\text{mm} \text{s}^{-1}))</th>
<th>(R_{yy}) (T)</th>
<th>(I(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.3</td>
<td>(p(\text{O}_2)) = 0.21 atm, 295 K</td>
<td>6</td>
<td>295</td>
<td>0.35</td>
<td>0.34</td>
<td>(\sim) 100</td>
<td>100</td>
</tr>
<tr>
<td>0.4</td>
<td>(p(\text{O}_2)) = 0.21 atm, 295 K</td>
<td>6</td>
<td>295</td>
<td>0.46</td>
<td>(-0.02)</td>
<td>46.0</td>
<td>100</td>
</tr>
<tr>
<td>0.3</td>
<td>(p(\text{O}_2)) = 10\textsuperscript{-18} atm, 1223 K</td>
<td>5</td>
<td>295</td>
<td>0.35</td>
<td>0.31</td>
<td>100</td>
<td>29</td>
</tr>
<tr>
<td>0.4</td>
<td>(p(\text{O}_2)) = 10\textsuperscript{-18} atm, 1223 K</td>
<td>5</td>
<td>295</td>
<td>0.27</td>
<td>1.07</td>
<td>71</td>
<td>42</td>
</tr>
</tbody>
</table>

\textsuperscript{*}CN, IS, QS, 2\nu, \(R_{yy}\), and \(I\) are the coordination number of trivalent iron, isomer shift relative to metallic \(\alpha\)-Fe at 295 K, quadrupole shift, magnetic hyperfine field and relative area, respectively. The average \(R_{yy}\) values of magnetic sextet distributions are given. Estimated errors are \(\pm 0.2\text{ mm s}^{-1}\) for IS, \(\pm 0.2\%\) for \(I\), and \(\pm 0.02\text{ mm s}^{-1}\) for the other parameters. CN is the coordination number of iron cations in the perovskite lattice.

1473 K during 10 h in order to avoid local inhomogeneities, phase separation and excessive sintering of the porous layers; these conditions were selected on the basis of preliminary tests involving various thermal treatments, XRD and SEM/EDS. Then the \((\text{LaSr})_y\text{CrFe}_x\text{O}_{3−\delta}\) electrodes were applied onto CGO/LSGM half-cells and sintered at 1473 K for 10 h in air. After fabrication of CE and RE, assembling of the setup and sealing, the anodes were equilibrated at 1073 K in 10\%\textsubscript{H}_2−90\%\textsubscript{N}_2 atmosphere (flow rate of 3 L h\textsuperscript{-1}) during 15–25 s before starting the measurements. The polarization tests were carried out in the galvanostatic mode using a PGSTAT302 Autolab instrument at 873–1073 K in flowing wet 10\%\textsubscript{H}_2−\textsubscript{N}_2 gas mixture (3 L h\textsuperscript{-1}) where the oxygen chemical potential was continuously controlled by an electrochemical oxygen sensor. The ohmic (\(R\)) and polarization (\(R_{yy}\)) resistances were determined from the impedance spectra collected in the frequency range from 10 mHz to 1 MHz (AC signal amplitude of 3 mA). The relaxation times after a change in the working electrode potential or temperature were 1–8 h. The as-prepared and tested electrodes were examined by XRD and SEM/EDS in order to reveal possible structural and microstructural alterations.

3. Results and discussion

3.1. Structural and microstructural characterization

XRD analysis confirmed that as-prepared \((\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_{1−x}\text{Fe}_x\text{O}_{3−\delta}\) powders and ceramics are single-phase, as expected (Fig. 1). The crystal structure under oxidizing conditions was identified as rhombohedrally distorted perovskite (space group \(R\text{3c}\)); the unit cell parameters are listed in Table 1. Note that the rhombohedral symmetry was earlier reported for \(\text{LaSr}_x\text{FeCr}_y\text{O}_{8+\delta}\) \[23\], whereas \((\text{La}_{0.75}\text{Sr}_{0.25})\text{Cr}_x\text{Fe}_y\text{O}_3\) is orthorhombic \[24\]. Compared to the latter compound, the higher symmetry of oxidized \((\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_{1−x}\text{Fe}_x\text{O}_3\) \((x=0.3–0.4)\) originates from the higher concentration of \(\text{Cr}^{3+}/\text{Cr}^{4+}\) cations having smaller radii with respect to \(\text{Fe}^{3+}/\text{Fe}^{4+}\) \[25\], thus increasing the perovskite tolerance factor. Moreover, in accordance with the data on electronic transport and Mössbauer spectra discussed below, the average oxidation state of chromium cations is higher than that of iron, again decreasing the average size of B-site cations in the ABO\(_3\) perovskite structure. Consequently, reduction in H\(_2\)-containing atmospheres leads to a larger radius of these cations and to a lower symmetry. Rietveld refinement of the XRD patterns of reduced \((\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_{1−x}\text{Fe}_x\text{O}_{3−\delta}\) (Fig. 1) showed the formation of orthorhombic perovskite structure, space group \(P\text{bnm}\). At the same time, no traces of metallic iron or other secondary phases were detected, in agreement with Mössbauer spectroscopy. This confirms phase stability of the title perovskites in a wide \(p(\text{O}_2)\) range necessary for the SOFC anode applications, although the “rhombohedral ↔ orthorhombic” transition may contribute to chemically induced strains in redox cycling.

EDS/SEM inspection of the dense ceramics and porous electrodes revealed an absence of compositional inhomogeneities and quite uniform grain size. The cation composition of selected samples was also verified by the ICP spectroscopy. Typical microstructures are shown in Fig. 2. The electrodes were found to possess a sufficient porosity and a relatively high surface area, ensuring gas penetration and facile surface-exchange processes. The average grain diameter in the electrode layers varied in the range 0.5–2 \(\mu\)m. No exfoliation or visible reaction traces between \((\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_{1−x}\text{Fe}_x\text{O}_{3−\delta}\) and CGO interlayer were observed (Fig. 2D). The electrochemical tests during over 150 h did not result in significant changes of the electrode grain size.


Fig. 1. XRD patterns of \((\text{La}_{0.75}\text{Sr}_{0.25})_{0.95}\text{Cr}_{1−x}\text{Fe}_x\text{O}_{3−\delta}\) \((x=0.3, 0.4)\) after sintering in air at 1973 K, annealing in air at 1473 K and subsequent reduction in H\(_2−\text{H}_2\text{O−N}_2\) atmosphere at 1223 K.
Fig. 2. SEM micrographs of fractured (La0.75Sr0.25)0.95Cr0.6Fe0.4O3−δ ceramics (A), top-view of as-prepared porous electrodes of (La0.75Sr0.25)0.95Cr0.7Fe0.3O3−δ (B) and (La0.75Sr0.25)0.95Cr0.6Fe0.4O3−δ (C), and fractured (La0.75Sr0.25)0.95Cr0.6Fe0.4O3−δ electrode applied onto CGO/LSGM after electrochemical measurements in H2-containing atmosphere (D).

3.2. Defect formation in oxidized (La0.75Sr0.25)0.95Cr1−xFexO3−δ

The Mössbauer spectra of (La0.75Sr0.25)0.95Cr1−xFexO3−δ equilibrated with atmospheric oxygen down to room temperature, are shown in Figs. 3 and 4. The room-temperature spectra (Fig. 4, top) can be adequately described by a quadrupole doublet with isomer shifts typical for octahedrally coordinated Fe3+ in oxide phases, Table 2 (e.g., [26–28] and references cited). At 4.2 K, below the magnetic transition point, a distribution of magnetic sextets is observed (Fig. 3). The estimated IS and the average magnetic hyperfine field (Bhf) indicate that six-fold coordinated Fe3+ is the predominant state of iron cations. The distribution originates hence from the so-called chemical disorder, i.e., the presence of different cations in the nearest coordination spheres of iron (Cr3+, Cr4+ and Fe3+ in the B sublattice, and La3+ and Sr2+ in the A sites).

Fig. 3. Mössbauer spectra of oxidized (La0.75Sr0.25)0.95Cr1−xFexO3−δ, collected at 4 K. The lines plotted over the experimental points are the distributions of magnetic sextets. Insets show the probability distribution of the magnetic hyperfine fields.
Fig. 5. Relative weight changes in (La0.75Sr0.25)0.95Cr1-xFe2O3-δ, measured by TGA on reduction in flowing dry 1\(\text{H}_2\)–9\(\text{N}_2\) gas mixture at 1223 K (A) and 1073 K (B). Inset shows the weight changes in atmospheric air with respect to room temperature. Dashed vertical lines correspond to the atmosphere changes from air to flowing Ar and then to 10\(\text{H}_2\)–9\(\text{N}_2\).

different configurations in the second coordination sphere around Fe\(^{3+}\) give rise to slightly different, unresolved magnetic hyperfine interactions. Note that the IS remains essentially constant for all the sextets in the distribution, thus confirming that all Fe\(^{3+}\) cations have the same octahedral coordination. No Fe\(^{4+}\) states are detected in the Mössbauer spectra.

In combination with the absence of pentacoordinated Fe\(^{3+}\), the latter observation shows that the charge compensation of S\(^{2-}\) and A-site cation deficiency in oxidized (La0.75Sr0.25)0.95Cr1-xFe2O3-δ occurs via formation of tetravalent chromium cations. The concentration of oxygen vacancies under oxidizing conditions is very low since no traces of iron–oxygen pyramids, easily detectable after reduction (see below), are found. It should be mentioned that Cr\(^{4+}\) is frequently observed in Cr-containing perovskite phases [29–31], where surface oxidation and hydration at low temperatures leads even to the formation of Cr\(^{5+}\) impurity states. Indeed, thermo-gravimetric analysis of (La0.75Sr0.25)0.95Cr1-xFe2O3-δ revealed very minor weight losses on heating in air (inset in Fig. 5). These variations may originate from surface desorption processes in the intermediate-temperature range and an appearance of modest oxygen deficiency in the bulk at high temperatures. Decreasing chromium concentration leads to smaller mass changes, thus suggesting that the oxygen vacancy formation, if any, is governed by the Cr\(^{3+}\)/Cr\(^{4+}\) redox couple. However, even neglecting any desorption from (La0.75Sr0.25)0.95Cr1-xFe2O3-δ surface, the resultant oxygen nonstoichiometry variations are extremely low, (6–10) \(\times\) 10\(^{-4}\) atoms per formula unit (f.u.). In other words, the crystal lattice of (La0.75Sr0.25)0.95Cr1-xFe2O3 under oxidizing conditions is oxygen-stoichiometric within the limits of experimental uncertainty. The same conclusion can be drawn from the behavior of Seebeck coefficient in oxidizing atmospheres (Fig. 6). As discussed below, the Seebeck coefficient is proportional to the reciprocal electron–hole concentration and reflects, therefore, any variations in the oxygen deficiency due to the crystal electroneutrality condition. The facts that the thermopower is p(O\(_2\))-independent within the limits of experimental error and displays only a very weak increase with temperature, confirm that the oxygen stoichiometry under oxidizing conditions is essentially constant. Finally, one should mention that the observed trends are in excellent agreement with numerous literature data showing that the average oxidation state of chromium cations in Fe- and Cr-containing perovskite solid solutions is always higher than that of iron (e.g., [30,31] and references cited).

3.3. Stability and redox behavior in reducing atmospheres

The room-temperature Mössbauer spectra of reduced (La0.75Sr0.25)0.95Cr1-xFe2O3-δ, annealed at 1223 K and oxygen partial pressure of 10\(^{-18}\) atm (Fig. 4), display three resolved peaks. One of them, at the highest Doppler velocity, has a pronounced shoulder. These spectra were fitted with two quadrupole doublets; introducing additional subspectra did not improve the fitting quality. The IS and quadrupole splitting (QS) values of both doublets (Table 2) indicate that most iron cations still exist in the trivalent state, but oxygen-vacancy formation on reduction leads to the presence of two major coordination numbers, 6 and 5. For the doublet with higher IS, the estimated IS and QS are equal to those of the oxidized phases within the limits of experimental error. This doublet corresponds hence to octahedrally coordinated Fe\(^{3+}\). The lower IS of another subspectrum is characteristic of a lower coordination. The corresponding isomer shifts are significantly higher compared to the range known for tetrahedral coordination in the ferrite-based phases with perovskite-related or brownmillerite structures (0.14–0.21 mm s\(^{-1}\)), but are typical for pentacoordinated Fe\(^{3+}\) in perovskites [20,26–28,32,33]. Furthermore, this doublet assigned to pentacoordinated Fe\(^{3+}\) has higher QS (Table 2), as expected for the sites with more distorted local coordination than octahedral. Hence, the predominant oxidation state of iron in (La0.75Sr0.25)0.95Cr1-xFe2O3-δ remains 3+ irrespective of the oxygen partial pressure, while oxygen coordination of Fe\(^{3+}\) cations decreases on reduction.
Fig. 7 shows the relative variations of oxygen deficiency on reduction in a dry 10%H₂–90%N₂ flow, calculated from the TGA results (Fig. 5), with respect to the stoichiometry in atmospheric air. The latter reference point was selected as no complete reduction into phase mixtures with definite oxidation states of iron and chromium cations (e.g., metallic Fe) was achieved during TGA experiments up to 1373 K. The results indicate that, although the reduction leads to significant nonstoichiometry values, the overall level of oxygen deficiency can still be tolerated by the perovskite lattice. Taking into account that iron cations are trivalent, and assuming δ = 0 in air for the sake of comparison, the estimated average oxidation state of chromium in dry 10%H₂–90%N₂ tends to asymptotically approach +3.10 for x = 0.3 and +3.01 for x = 0.4 at 1223 K. At 1073 K, these values are +3.16 and +3.14, respectively. Consequently, one may conclude that the redox behavior and electronic conductivity of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋ᵣ should be primarily governed by Cr³⁺/⁴⁺ states in the entire p(O₂) range relevant for the electrode applications. In particular, a higher concentration of p-type electronic charge carriers (Cr⁴⁺) and higher hole conductivity can be expected for the composition with x = 0.3, whereas the Seebeck coefficient should remain positive as for lanthanum–strontium chromites [31]. These trends are indeed observed experimentally (Fig. 8).

Although increasing iron content in (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋ᵣ increases reducibility of the materials (Fig. 7), the Mössbauer spectroscopy and thermogravimetric analyses showed no evidence of phase decomposition in mildly reducing atmospheres, as for XRD (Fig. 1). In particular, no traces of metallic iron were detected in the Mössbauer spectra after reduction at 1223 K and p(O₂) = 10⁻¹⁸ atm. Taking into account that the low-p(O₂) stability limits of iron-containing phases usually exhibit van’t Hoff type dependencies on reciprocal temperature [1,2,34,35], an acceptable thermodynamic stability under anodic conditions in the intermediate-temperature SOFCs can be

![Fig. 7. Variations of the oxygen nonstoichiometry of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋ᵣ on reduction in dry 10%H₂–N₂ gas mixture at 1223 K (A) and 1073 K (B), calculated from thermogravimetric data. The reference points correspond to atmospheric air at a given temperature. The dashed vertical lines show atmospheric changes.](image)

![Fig. 8. Oxygen partial pressure dependencies of the total conductivity (A and C) and Seebeck coefficient (B and D) of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋ᵣ ceramics. The data were collected in the isothermal oxidation regime (see text), with equilibration during 2–80 h for each data point. Solid lines are a guide for the eye.](image)
expected. Nonetheless, long-term testing is necessary to access reducibility in pure H₂ at 770–970 K.

Another conclusion can be drawn regarding the distribution of oxygen vacancies in the crystal lattice. In a hypothetical state when all chromium cations in reduced \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) are trivalent as iron, the oxygen vacancy concentration should correspond to approximately 0.19 sites per formula unit (δ ≈ 0.19). Assuming a random distribution of the vacancies, approximately 28% penta- and 5% tetra-coordinated B-sites might be predicted. The estimated fractions of pentacoordinated \(Fe^{3+}\) (Table 2) are, however, substantially higher, thus indicating a greater energetic favorability for vacancy location near \(Fe^{3+}\) in comparison with \(Cr^{3+}\). Moreover, the calculated concentration of pentacoordinated \(Fe^{3+}\), 0.21–0.23 cations/f.u., is almost independent of \(x\) and is close to the vacancy concentration. As the oxygen nonstoichiometry is relatively low and statistical distribution of \(Cr/Fe\) cations in the B sublattice was confirmed by Rietveld refinement of the XRD patterns, the absence of tetrahedral \(Fe^{3+}\) may originate from coulombic repulsion between the randomly distributed oxygen vacancies. No tendency to anion vacancy ordering is hence observed on reduction, in contrast to the perovskite-related systems with higher oxygen deficiency [20,26,27,32]. In the latter cases, extensive vacancy ordering under reducing conditions similar to those used in this work, leads always to an appearance of brownmillerite-like domains and/or to the phase transitions into brownmillerite-type structures. This factor, along with the high concentration of chromium–oxygen polyhedra limiting the chains of iron cations and oxygen vacancies in the perovskite structure, seems responsible for the improved thermodynamic stability of \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\).

### 3.4. Thermal and chemical expansion

**Fig. 9** presents typical dilatometric curves of \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) ceramics at atmospheric \(p(O_2)\) and in flowing CO–CO₂ mixture where the oxygen partial pressure, continuously controlled by an electrochemical oxygen sensor, was \(3 \times 10^{-20}\ atm\) at 923 K and \(2 \times 10^{-12}\ atm\) at 1223 K. In both cases the thermal expansion is essentially linear up to 1373 K. When pre-oxidized \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) is heated in CO–CO₂ atmosphere, a significant chemical contribution to the apparent thermal expansion is observed at 970–1220 K due to progressive oxygen losses from the perovskite lattice (Fig. 9B). This effect becomes negligible on cooling back in the same atmosphere. The average TECs of \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) are relatively low, \(10.3–11.3 \times 10^{-6}\ K^{-1}\), and tend to slightly decrease on reduction (Table 1). These TECs are substantially lower compared to other Fe-containing perovskites (e.g., [36]) due to the stabilization of \(Fe^{3+}\) state, validated by the Mössbauer spectroscopy. The moderate thermal expansion provides an acceptable thermomechanical compatibility of \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) and common solid electrolytes, such as yttria-stabilized zirconia (YSZ), CGO and LSCM [7].

Despite the \(Fe^{3+}\) stabilization, the isothermal chemical strains of \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) measured by dilatometry (Table 3) are 3–5 times higher compared to \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Mn_{0.5}O_{3-δ}\) under similar conditions [12]. The overall level of chemical expansion observed for \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) is intermediate between the perovskite-type chromites [37,38] and ferrites [39,40]. This fact, and increasing volume variations when \(x\) increases (Table 2), are in agreement with the above conclusion that iron additions promote reducibility of \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\). Note that the apparent lattice expansivity may also be contributed by the "rhombohedral → orthorhombic" phase transition on reduction. Whatever the microscopic mechanisms, the thermomechanical stability of the title materials on redox cycling is worse with respect to LSCM.

### 3.5. Oxygen permeability and ionic conductivity

**Fig. 10** displays the steady-state oxygen permeation fluxes \(j\) through dense \((La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}Fe_{x}O_{3-δ}\) membranes of different thickness \(d\) under air/H₂–H₂O–N₂ gradients, and the specific oxygen permeability \(J(O_2)\) related to these quantities as ([14,21,31] and references therein):

\[
j = \frac{J(O_2)}{d} = \frac{\ln \left( \frac{p_2}{p_1} \right)}{d}
\]

where \(p_2\) and \(p_1\) are the feed- and permeate-side oxygen partial pressures, respectively. When the oxygen chemical potential gradient is high enough, increasing membrane thickness leads to lower oxygen fluxes, whereas the specific permeability proportional to \(j \times d\) becomes thickness-independent within the limits of experimental error. In combination with relatively high total conductivity
(Fig. 8), this behavior unambiguously indicates that the permeation is limited by the bulk ion diffusion, and makes it possible to estimate the partial ionic conductivity \( \sigma_0 \) in reducing atmospheres [2]:

\[
\sigma_0(p_1) = \frac{16F^2d}{RT} \left( \frac{\partial j}{\partial \ln p_1} \right)_{p_2}
\]

where \( F, R \) and \( T \) are the Faraday constant, universal gas constant and absolute temperature, respectively. When the oxygen chemical potential gradient decreases, the permeability starts to depend on the membrane thickness, thus indicating an increasing role of the surface exchange and/or gas diffusion limitations. The values of ionic conductivity and ion transference numbers \( (t_0 = \sigma_0/\sigma) \) estimated by Eq. (2) at 1223 K and \( p_1 = 10^{-17} \) atm, are presented in Table 4. As for the oxygen permeation fluxes (Fig. 10), the bulk ionic transport in the title perovskites increases with \( x \) due to increasing oxygen vacancy concentration in reducing atmospheres and weakening of the metal–oxygen bonds. Due to the same reasons, the ionic conductivity of \( (La_{0.75}Sr_{0.25})_xCr_{1-x}FeO_{3-x} \) is higher with respect to acceptor-doped lanthanum chromites [41] and LSCM [12], but is slightly lower compared to \( (La, Sr)FeO_3 \) with similar strontium concentrations [42].

3.6. p-Type electronic conductivity and Seebeck coefficient

Under oxidizing conditions the total conductivity of \( (La_{0.75}Sr_{0.25})_{0.95}Cr_{1-x}FeO_{3-x} \) (Fig. 11) is comparable to that of other oxide electrode materials considered for the “symmetrical” SOFC/SOEC concept where the cathode and anode are made of similar compositions, such as LSCM [3,7]. The thermopower sign and oxygen permeation data (Figs. 6, 8 and 10) show that the conductivity is predominantly p-type electronic in the entire \( p(O_2) \) range studied in this work, as for most acceptor-doped phases based on LaCrO\(_3\) [31]. Increasing iron content leads to lower hole conduction, primarily due to lowering charge-carrier concentration. As the oxygen nonstoichiometry variations in oxidizing atmospheres are very minor (Fig. 5), the conductivity and Seebeck coefficient in these conditions are essentially independent of the oxygen pressure (Figs. 6 and 11B). Reducing \( p(O_2) \) results in progressive oxygen losses from the perovskite lattice and in decreasing \( Cr^{4+} \) concentration, which is reflected by lower conductivity and higher thermopower (Fig. 8). This mechanism implies a small-polaron hopping mechanism as for other perovskite-type chromites [31]. Although exact calculations of the charge-carrier concentration \( p \) and mobility \( \mu_p \) are impossible in the present case due to a variety of likely spin-degeneracy and site-exclusion effects influencing the number of states \( N \), their temperature dependencies in a narrow \( p(O_2) \) range can be evaluated from the Seebeck coefficient and conductivity data by using the modified Heikes equation [35,43]:

\[
\alpha = k \frac{\partial N - p}{p} \ln \frac{\beta N}{p} \frac{p}{N_{fu}}
\]

\[
\sigma = \mu_p e \frac{N_{fu}}{V_{uc}}
\]

where \( N \) and \( p \) are related to unit formula, \( N_{fu} \) is the number of formula units per unit cell, \( V_{uc} \) is the unit-cell volume, and \( \beta \) is a constant. The results of these estimations (Fig. 12) indicate, first of all, that the hole mobility is thermally activated. This fact and the relatively low mobility level both provide evidence in favor of the small-polaron mechanism, in agreement with the conductivity activation energies (Table 1). Second, the \( \mu_p \) values tend to decrease
with increasing x, suggesting that the iron cations are essentially excluded from the hole migration processes.

Fig. 13 illustrates another important trend revealed in the course of the electrical measurements, namely, strong hysteresis phenomena on redox cycling in moderately reducing atmospheres. The corresponding p(O₂) range extends down to substantially lower oxygen chemical potentials than those known for the experimental problems, associated with stagnated diffusion and equilibration processes involving gaseous oxygen-containing species (e.g., [42]). At 973–1273 K the sluggish equilibration kinetics and resultant experimental uncertainties are usually observed at the oxygen partial pressures from 10⁻⁵–10⁻⁴ atm down to 10⁻¹¹–10⁻⁸ atm, depending on temperature. In the present case, the low-p(O₂) limit of the nonequilibrium plateau-like behavior indicative of “frozen” oxygen stoichiometry on reduction (Fig. 13) varied from 10⁻¹⁴ down to 10⁻²⁰ atm at 1023–1223 K, although the time used for equilibrating was up to 80 h. Notice that a significantly faster equilibration and a good reproducibility of the electrical properties were observed beyond the plateaus. Consequently, only the data collected in the oxidation regime (Fig. 8) were considered as equilibrium, except for 973–1073 K when the oxygen content seems to be essentially frozen in the entire p(O₂) range. The hysteresis phenomena may originate either from the perovskite phase transition or from large differences in the oxygen exchange kinetics on reduction and on oxidation. Whatever the microscopic mechanisms, the slow reduction kinetics makes it possible to increase electronic conductivity of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋₄ electrodes under intermediate-temperature SOFC anodic conditions by periodic current pulses.

3.7. Anodic performance

Typical anodic polarization curves of porous (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋₄ electrodes with CGO interlayers applied onto LSGM solid electrolyte are presented in Fig. 14. Table 5 compares their electrochemical performance with other anode compositions, including LSCM [12] and (La,Sr)(Ti,Cr,Mn)O₃−δ perovskites [14]. All the data were collected in the electrochemical cells with similar electrode thicknesses and geometry described elsewhere [12], using identical Pt gauze current collectors, in flowing 10%H₂–90%N₂ gas mixture humidified at room temperature. The electrochemical activity of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋₄ tends to moderately decrease with increasing x, in correlation with the electronic conductivity variations. The opposite trends observed for the oxygen permeability and ionic conductivity (Fig. 10 and Table 4) confirm previous conclusion [12] that the impact of ionic transport is much smaller with respect to electronic, probably due to the presence of ceria interlayers and/or overall low level of the ion diffusivity. These factors make it possible to increase the electrode performance via infiltration of CeO₂−δ, which is a mixed ionic–electronic conductor under anodic conditions. On the other hand, the electrode resistivity variations with composition (Table 5) cannot be simply ascribed to the only limiting effect of electron transfer, and indicate also non-negligible contributions

Table 4
Partial oxygen–ionic conductivity and ion transference numbers of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋₄ at T = 1223 K and p(O₂) = 10⁻¹⁷ atm, calculated from the oxygen permeability and total conductivity data.

<table>
<thead>
<tr>
<th>x</th>
<th>Total conductivity σ, S cm⁻¹</th>
<th>Ionic conductivity σ₀, S cm⁻¹</th>
<th>Transference numbers t₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.3</td>
<td>1.41</td>
<td>0.056</td>
<td>0.040</td>
</tr>
<tr>
<td>0.4</td>
<td>0.54</td>
<td>0.079</td>
<td>0.15</td>
</tr>
</tbody>
</table>

![Fig. 12](image-url) Oxygen partial pressure dependencies of the normalized hole mobility (A) and p(δN) ratio (B) calculated from the conductivity and Seebeck coefficient data at 1023–1223 K, and temperature dependencies of these parameters at p(O₂) = 10⁻² atm (C). Solid lines are a guide for the eye.

![Fig. 13](image-url) One example of the hysteresis behavior of the electrical properties of (La₀.₇₅Sr₀.₂₅)₀.₉₅Cr₁₋ₓFeₓO₃₋₄ on redox cycling.
of other factors, such as metal–oxygen bonding energy affecting oxygen exchange kinetics [7]. For instance, the polarization resistance of (La0.75Sr0.25)0.95Cr0.3Mn0.5Ti0.2O3−δ is lower than that of (La0.75Sr0.25)0.95Cr0.35Mn0.35O3−δ, although the electronic conductivity of LSCM is higher.

4. Conclusions

The Mössbauer spectroscopy and thermogravimetric studies combined with electrical measurements show that the incorporation of moderate amounts of iron into acceptor-doped LaCrO3 enable to stabilize Fe3+ states in a wide p(O2) range necessary for the intermediate-temperature electrochemical applications. Perovskite-type (La,Sr)[Cr,Fe]O3−δ exhibit moderate TECs compatible with common solid electrolytes materials, relatively low oxygen-stoichiometry variations, and substantially high oxygen permeability and ionic conduction in reducing atmospheres. These perovskites possess also a higher electrochemical activity under anodic conditions and lower costs compared to LSCM. At the same time, a number of functional properties of (La,Sr)[Cr,Fe]O3−δ, primarily electronic conductivity at low p(O2) and chemical expansivity, are worse with respect to their Mn-containing analogs. When iron concentration increases, these disadvantages become more pronounced. Namely, the electronic conductivity decreases with increasing x in (La0.75Sr0.25)0.95Cr1−xFe0.5O3−δ, while the polarization resistance, reducibility and chemical expansion tend to increase. The developments of new electrode compositions require, therefore, an optimization of Cr:Fe concentration ratio in the range 0.1 ≤ x ≤ 0.3, and infiltration of catalytically active additives with significant mixed conductivity, such as doped ceria or CeVO3:δ.

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