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Oxide Diffusion in Innovative SOFC cathode materials

Y. Hu\textsuperscript{a}, V. Thoréton\textsuperscript{b}, C. Pirovano\textsuperscript{b}, E. Capoen\textsuperscript{b}, C. Bogicevic\textsuperscript{a}, N. Nuns\textsuperscript{b}, A.-S. Mamede\textsuperscript{b}, G. Dezanneau\textsuperscript{a} and R. N. Vannier\textsuperscript{a}\textsuperscript{*}

Oxide diffusion was studied in two innovative SOFC cathode materials, Ba\textsubscript{0.75}Co\textsubscript{0.25}O\textsubscript{4} and Ca\textsubscript{0.6}Co\textsubscript{0.4}O\textsubscript{3−δ} derivatives. Although oxygen diffusion was confirmed in the promising Ba\textsubscript{0.75}Co\textsubscript{0.25}O\textsubscript{4}, it was not possible to derive accurate transport parameters because of an oxidation process at the sample surface which has still to be clarified. In contrast, the oxygen diffusion in the well-known Ca\textsubscript{0.6}Co\textsubscript{0.4}O\textsubscript{3−δ} thermoelectric materials was improved when calcium was partly substituted with strontium, likely due to an increase of the volume of the rock salt layers in which the conduction process takes place. Although the diffusion coefficient remains low, interestingly, fast kinetics towards the oxygen molecule dissociation reaction were shown with surface exchange coefficients higher than those reported for the best cathode materials in the field. They increased with the strontium content, the Sr atoms potentially playing a key role in the mechanism of oxygen molecule dissociation at the solid surface.

Introduction

With the possibility of cogeneration of electricity and heat, Solid Oxide Fuel cells are promising. First units were commercialised in Japan in 2009 and demonstrators are running in Europe. Their principle is based on a dense ceramic electrolyte which separates two compartments, one containing hydrogen, and the other containing air. Because of a difference of oxygen chemical potential between the two compartments, oxygen molecules are reduced at the cathode into oxide ions, which migrate through the electrolyte to the anode to react with hydrogen molecule and produce water. Hydrogen is usually produced by methane reforming, however internal reforming of methane is also envisaged. Typical temperatures of use range are in between 700 and 1000°C. The main electrolyte materials which are currently studied are Yttria Stabilised Zirconia (YSZ), Gadolinia Doped Ceria (GDC) and lanthanum gallates (LSGM). At the anode, a Ni-YSZ cermet containing 40% in volume of nickel is usually developed, but it is prone to carbon deposition when used with hydrocarbon fuels. It is also sensitive to sulphur poisoning and suffer from nickel agglomeration and redox stability under prolonged usage. Alternative materials such as the chromo-manganites (La\textsubscript{0.75}Sr\textsubscript{0.25})Ca\textsubscript{0.5}Mn\textsubscript{0.5}O\textsubscript{3} or titanates such as lanthanum doped SrTiO\textsubscript{3} have been proposed. At the cathode, the strontium substituted lanthanum manganite (La\textsubscript{1−x}Sr\textsubscript{x}MnO\textsubscript{3}) is conventionally used. However, it displays low oxide ion conductivity and composites with YSZ were developed to allow its use at temperature lower than 800°C. To operate at lower temperature, Mixed Ionic Electronic Conductors (MIEC) are usually preferred to allow the oxygen reduction reaction to be sprayed on the whole electrode volume and not only at the electrode/electrolyte interface. As an alternative to LSM, the La\textsubscript{0.6}Sr\textsubscript{0.4}Co\textsubscript{0.2}Fe\textsubscript{0.8}O\textsubscript{3−δ} (LSCF) perovskite has been widely studied\textsuperscript{1}. When used with a GDC electrolyte and after optimisation of its microstructure, an area specific resistance (ASR) of only 0.13 Ω.cm\textsuperscript{2} was obtained at 603°C for this compound\textsuperscript{2}, the target for the cathode being 0.15 Ω.cm\textsuperscript{2} at 700°C. These high performances are likely due to the presence of Co(III) in the structure which is known to be particularly active for the reduction of oxygen molecules into oxide ions\textsuperscript{3}. Indeed, cobaltites are known to display high oxide ion transport parameters. Among these, the La\textsubscript{0.6}Sr\textsubscript{0.4}CoO\textsubscript{3−δ} composition is among the best catalyst for the oxygen molecule reduction reaction with a surface exchange coefficient (k\textsuperscript{*}, cm/s) of 1.1×10\textsuperscript{−6} cm/s associated to an oxygen self-diffusion tracer (D\textsuperscript{O*}, cm\textsuperscript{2}s) coefficient of 2.0×10\textsuperscript{−9} cm\textsuperscript{2}s at 860°C under 230 mbar\textsuperscript{4}. One drawback of these materials is their high thermal expansion coefficient with a value of 23×10\textsuperscript{−6} K\textsuperscript{−1} for this later composition which is twice the thermal expansion of YSZ (10×10\textsuperscript{−6} K\textsuperscript{−1})\textsuperscript{5} and GDC (12×10\textsuperscript{−6}K\textsuperscript{−1})\textsuperscript{6}. However, partial substitution of cobalt with iron in LSCF led to a good compromise with a TEC in the range of 14-15×10\textsuperscript{−6} K\textsuperscript{−1} from 500 to 700°C\textsuperscript{7}. More recently, promising performances as SOFC cathodes were shown for two innovative cobaltites, Ba\textsubscript{2}Co\textsubscript{4}O\textsubscript{14} and Ca\textsubscript{3}Co\textsubscript{4}O\textsubscript{11.5δ}. An ASR of 0.5 Ω.cm\textsuperscript{2} was first measured on a symmetric cell made of a 70 wt% Ba\textsubscript{2}Co\textsubscript{4}O\textsubscript{14} - 30 wt% GDC composite electrode, deposited on a GDC
electrolyte. The good electrochemical performances of this compound were later confirmed by Li et al. who obtained ASR of only 0.133 Ω cm² and 0.068 Ω cm² at 750 and 850 °C and powder density of 450 mW/cm² and 770 mW/cm² on a cell made of a LSGM electrolyte with a Ba₂CoO₄₋₈ – samarium doped ceria composite cathode at 800 and 850°C, respectively. This compound exhibits a three dimension structure based on a [CoO₉₃] blocks sandwiched in between [Ba₂CoO₃] layers⁸,¹⁰ (Fig 1a). With cobalt atoms at difference valences, it displays high electronic conductivity with a value of 240 S cm⁻¹ at 650°C but as other cobaltites, its thermal expansion coefficient is high (22×10⁻⁶ K⁻¹) which supposes the use of a composite at least for mechanical compatibility between the electrode and the electrolyte. After optimisation of its composition and thickness, an ASR of only 0.08 Ω cm² at 700°C was recently measured on symmetric cell made of a 50 wt% Ba₂CoO₄₋₈ – 50 wt% GDC composite electrode with a GDC electrolyte¹¹.

In contrast, a thermal expansion coefficient of only 10×10⁻⁶ K⁻¹ was measured for the well-known Ca₃Co₂O₇-δ thermoelectric compound¹² whose structure is a misfit composed of CdI₂-type [CoO₂] layers sandwiched in between [Ca₃Co₂O₇] rock salt slabs¹³,¹⁴ (Fig 1b). With an electronic conductivity of 100 S cm⁻¹ at 700°C and oxygen vacancies in the rock salt layers, Mixed Ionic Electronic Conductivity was expected for this compound. When tested in symmetrical cell with a GDC electrolyte, ASR of 4.94 Ω cm² for the pure compound and 1 Ω cm² for a 70 wt% Ca₃Co₂O₇-δ - 30 wt% GDC composite electrode were first measured¹⁵,¹⁶. After optimisation of composition and thickness it was decreased to 0.5 Ω cm² at 700°C for a 50 wt% Ca₃Co₂O₇-δ - 50 wt% GDC composite electrode with a thickness of 21 µm, and further decrease to 0.35 Ω cm² when calcium was partly substituted with 10% of strontium¹⁷. These good performances were confirmed by Samson et al.¹⁸, who reported a polarization resistance of 0.64 Ω cm² at 600°C for a 50 vol% Ca₃Co₂O₇-δ - 50 vol% GDC composite electrode. Using a samarium doped ceria electrolyte, Zou et al.¹⁹ measured a maximum power density of 430 mW cm⁻² at 700°C for a 70 wt% Ca₂₊₂Ba₁CoO₉₋₈ - 30 wt% Ce₁₋₃SmₓO₂₋₉₅ | Ce₁₋₉Smₓ₀₂₋₉₅ | Ce₁₋₉Smₓ₀₂₋₉₅-Ni button cell. More recently, oxygen diffusion in this material was confirmed by combing °¹⁸O/O isoexchange, electrical conductivity relaxation and pulse isotopic exchange technique²⁰. Despite low diffusion with coefficient of 2.7×10⁻¹⁰ cm².s⁻¹ at 700°C, this study showed high kinetics toward oxygen dissociation with a surface exchange coefficient as high as 1.6×10⁻⁷ cm².s⁻¹.

In the present paper, oxygen diffusion in these two innovative cathodes was investigated with an emphasise put on the impact of calcium partial substitution with strontium on the Ca₃Co₂O₇₋δ oxygen transport properties.

**Experimental**

Ba₂CoO₄₋₈ and (Ca₁₋ₓSrₓ)₂Co₉₋₈ox (x=0.1, 0.2) powders were prepared by solid-state reaction from the corresponding carbonates and oxides BaCO₃ (Prolabo – 99.5%, dried at 100°C), CaCO₃ (Prolabo – 99.5%, dried at 100°C), SrCO₃ (Aldrich – 99.8%, dried at 100°C) and Co₉O₄ (Alfa Aesar, 99.7%). Reactants were carefully ground in an agate mortar and then calcined repeatedly at 900°C with intermediate grinding. Purity of samples was checked by X-ray diffraction (D8 advance Bruker AXS diffractometer, Cu Kα radiation, 1D LynxEye PSD detector). The Fullprof software [v2.05, July 2011] was used for refinement of the unit cell parameters²¹. An Accupyc II 1340 Micromeritic pycnometer was used to measure the powder density. The absolute oxygen stoichiometry of the sample was determined by iodometric titration²². Thermal analyses were conducted under air, using a TG/DTA Setaram Setys apparatus. 40 mg of the sample powder was introduced into a Pt crucible, and the experiment was carried out at a heating rate of 5°C/min under flowing air (0.4 L/h) from room temperature to 950°C. A blank experiment with an empty crucible was performed to correct the data for effects of buoyancy. Thermogravimetry was also carried out for evaluation of the oxygen stoichiometry of the sample as a function of pO₂. In this case, a Setaram 92-1750 apparatus was used. The experiment was carried out with 50 mg of powder in a Pt crucible under flowing O₂/N₂ mix (5 L/h), with three successive oxygen partial pressures (100, 200 and 400 mbar), and temperature ranging from 600 to 750°C (50°C step) for each pressure. For this set-up, the buoyancy was negligible and the data were directly used to derive the thermodynamic factor at each temperature.

Dense ceramics, 3 mm thick, were prepared by Spark Plasma Sintering at the IdF Platform in Thiais (France). A 20 mm (diameter) carbon die was used. The samples were sintered within 15 minutes with a 2 minutes dwell at 850°C, under a 50 Mpa (8.8 kN) pressure. To remove carbon residues the pellets were annealed for 12 h at 800°C. A Hitachi S-3400N was used to characterise the samples’ surface.

For Isotopic Exchange and Electrochemical Conductivity Relaxation, samples surfaces were roughly polished in a first step before polishing down to 1/40 of micron with successive SiC papers and diamond pastes (Ra~10-20nm) in a second step. In case of calcium co-baltites, to avoid evolution of the surface roughness, samples were annealed for 72 h at 900°C before the second polishing in order to release residual strains in the material.
The $^{18}\text{O}/^{16}\text{O}$ Isotope Exchange Depth Profiling technique was applied to all samples to derive the oxygen self-diffusion tracer ($D^s$, cm²/s) and surface exchange coefficients ($k^s$, cm/s). The diffusion time was chosen to get diffusion lengths significantly smaller than the sample dimensions to consider a 1D diffusion model perpendicular to the surface of the dense ceramic into a semi-infinite medium. In these conditions, assuming first order surface exchange reaction and constant $^{18}\text{O}$ concentration during the exchanges, the solution for a semi-infinite solid to the Fick's 2nd law of diffusion is given by the following equation:

$$C'(x,t) = \frac{C(x,t) - C_{bg}}{C_p - C_{bg}} = \text{erfc} \left( \frac{x}{2 \sqrt{D^s \times t}} \right) - \left[ \exp \left( \frac{k^s x^2}{D^s} + \frac{k^s t x^2}{D^s} \right) \times \text{erfc} \left( \frac{x}{2 \sqrt{D^s \times t}} + k^s \frac{t}{D^s} \right) \right]$$

where $C_{bg}$ is the natural $^{18}\text{O}$ ratio, $C_p$ the $^{18}\text{O}$ ratio in the enriched exchange atmosphere, $x$ the distance from the surface of the specimen and $t$ the time of the isotope exchange.

Exchange conditions are reported in Table 1. Experiments were carried out in between 650 and 750°C under a pressure close to 210 mbar with a pre-annealing under normal oxygen (search grade oxygen 99.995% ALPHAGAZ) to reach thermodynamic equilibrium prior to the exchange under labelled oxygen. For both annealings, under the $^{18}\text{O}$ enriched atmosphere or pre-annealing, the samples were annealed for the required time and quenched. To determine the $^{18}\text{O}$ content in the oxygen gas used for the exchange, a silicon wafer was oxidised overnight at 1000°C under 200 mbar.

Table 1. Conditions of $^{18}\text{O}/^{16}\text{O}$ isotope exchange.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Pressure (mbar)</th>
<th>Pre-annealing time (hours)</th>
<th>Annealing time (min)</th>
<th>$^{18}\text{O}$ ratio in gas phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>648</td>
<td>210</td>
<td>240</td>
<td>2860</td>
<td>0.79</td>
</tr>
<tr>
<td>700</td>
<td>210</td>
<td>96</td>
<td>2765</td>
<td>0.80</td>
</tr>
<tr>
<td>750</td>
<td>218</td>
<td>96</td>
<td>2790</td>
<td>0.80</td>
</tr>
<tr>
<td>(Ca$_{1-x}$Sr$_x$)$_2$Co$<em>3$O$</em>{6.6}$</td>
<td>x=0.10, 0.20</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>604</td>
<td>207</td>
<td>160</td>
<td>725</td>
<td>0.88</td>
</tr>
<tr>
<td>649</td>
<td>211</td>
<td>100</td>
<td>567</td>
<td>0.67</td>
</tr>
<tr>
<td>694</td>
<td>219</td>
<td>61</td>
<td>392</td>
<td>0.80</td>
</tr>
<tr>
<td>741</td>
<td>210</td>
<td>20</td>
<td>55</td>
<td>0.9</td>
</tr>
</tbody>
</table>

A ToF-SIMS$^3$ machine (ION-TOF GmbH, 25 keV Bi$^+$ LMIG analysis gun, DSCS sputter gun) was used to measure the $^{18}\text{O}$ diffusion profiles using the same experimental protocol as described in ref 20. Both depth analysis from the sample surface (depth profile mode)$^3$ and edge analysis of the cut sample (line scan mode)$^2$ were performed and profiles merged. Depth of craters, after SIMS depth profiling, were measured with a KLA Tencor Alpha-step IQ profilometer. The same apparatus was used to measure the samples’ roughness. Electrical Conductivity Relaxation was also carried out on strontium doped Ca$_3$Co$_4$O$_9$.$\delta$. For this purpose, plane-parallel samples (17 mm in length and 1 to 2.4 mm in width) were cut from the strontium substituted dense ceramics using a diamond saw. Electrical conductivity was measured using the 4 points configuration while changing the oxygen concentration of the surrounding atmosphere as shown in ref 20. A gold paste was used to paint gold contacts which were sintered at 750 °C for 1h at a rate of 100 °C/h before experiment. The distance between the two internal electrodes was about 14 mm.

Once the sample was at equilibrium at given temperature, the $p$O$_2$ was step-wise changed and the conductivity was monitored as a function of time. The variation of $p$O$_2$ was performed by an abrupt switch among air and mixtures of O$_2$/Ar with oxygen partial pressures of 0.05 atm and 0.01 atm, respectively, at a 5L/h flow rate. In addition, the temperature-dependent electrical conductivity was measured under air with 5L/h flow rate at a very slow speed of 25 °C/h.

To characterise the nature of the atoms at the uppermost surface of the sample, Low-Energy Ion Scattering (LEIS) was also applied to a dense ceramic (Ca$_{1-x}$Sr$_x$)$_2$Co$_3$O$_{6.6}$ with x=0.1 composition which had been heated at 800°C in air during 12 hours prior to the analysis. The analysis was performed using a Qtacl$^{100}$ (ION-TOF GmbH) apparatus. Before analysis, the sample was in situ submitted to an atomic oxygen treatment (2-3 $\times$ 10$^{-4}$ mbar) to remove possible organic surface contamination. The study was carried out with a ‘He’ (3keV) beam on a 1mm$^2$ analysed area.

**Results and discussion**

**Characterization of powder and ceramics.** Synthesised powders and dense ceramics purity was confirmed by X-ray diffraction. As shown in Fig. 2, a good pattern matching was obtained for both type of ceramics.

Fig. 2. XRD calculated profile compared to experimental data collected on dense pellets of Ba$_3$Co$_4$O$_{14.5}$ (a), and (Ca$_{1-x}$Sr$_x$)$_2$Co$_3$O$_{6.6}$ with x=0.10 (b).
Corresponding unit-cell parameters are given in Table 2. For Ba$_2$Co$_9$O$_{14}$, they are in good agreement with the parameters reported by Ehora et al. (5.6963(8) Å, 28.924(6) Å) and Sun et al. (5.6958(4) Å, 28.909(4) Å), the small difference observed between the powders and the ceramics, on one hand, and these authors, on the other hand, being likely due to small difference in the oxygen content due to different thermal histories. For the refinement of (Ca$_{9-x}$Sr$_x$)$_3$Co$_9$O$_{27}$, the phases were considered as a mixture of two phases, one corresponding to the [CoO$_2$] layers and the second one corresponding to the [Ca$_3$CoO$_{14}$] rock salt layers, with common $a$, $c$ and $\beta$ cell parameters, constrained to be the same, and different $b_{ij}$ and $b_{RS}$, refined independently. The evolution of the unit cell parameters with the strontium content is in good agreement with the results reported by Wang et al., who showed almost no variation of the $b_{ij}$ unit cell characterising the [CoO$_2$] layers periodicity and an increase of the $b_{RS}$ unit cell with strontium content in good agreement with the higher radius of Sr$^{2+}$ (1.18 Å) compared to Ca$^{2+}$ (1.00 Å).

As shown by Ehora et al., Ba$_2$Co$_9$O$_{14}$ decomposed into CoO and a cubic BaCoO$_2$ form at 1030°C. To check the stability of the calcium cobalites, TGA and DTA analyses were carried out on the two strontium doped compositions. Their thermograms are compared to the signals measured for the non-doped compound in Fig. 3. They exhibit similar behavior. The first transition observed at 130°C for the pure compound disappears when strontium is introduced in the structure, the second transition at 550°C, corresponding to the beginning of a mass loss is maintained at the same temperature but the intensity of the associated endothermic peak on heating decreases with the increase of strontium content. All compounds decomposed before 950°C, the temperature of decomposition being slightly lowered when strontium content increases. This decomposition process is reversible but it is worth noting that the recombination happens at a lower temperature when substitution rate increases.

Table 2. Ba$_2$Co$_9$O$_{14}$ and (Ca$_{9-x}$Sr$_x$)$_3$Co$_9$O$_{27}$ $x=0.1$, 0.2 unit cell parameters from XRD on powder and dense ceramics.

<table>
<thead>
<tr>
<th></th>
<th>Powder</th>
<th>Dense ceramics</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Ba$_2$Co$<em>9$O$</em>{14}$ (R 3m)</td>
</tr>
<tr>
<td>$a$ (Å)</td>
<td>5.6945(3)</td>
<td>5.700(1)</td>
</tr>
<tr>
<td>$b$ (Å)</td>
<td>28.905(2)</td>
<td>28.926(6)</td>
</tr>
<tr>
<td>(Ca$_{9-x}$Sr$_x$)$_3$Co$<em>9$O$</em>{27}$ $x=0.10$</td>
<td>4.8429(3)</td>
<td>4.8433(9)</td>
</tr>
<tr>
<td>$b_{ij}$ (Å)</td>
<td>2.8253(5)</td>
<td>2.825(4)</td>
</tr>
<tr>
<td>$b_{RS}$ (Å)</td>
<td>4.5968(6)</td>
<td>4.563(2)</td>
</tr>
<tr>
<td>$c$ (Å)</td>
<td>10.8730(4)</td>
<td>10.881(2)</td>
</tr>
<tr>
<td>$\beta$ (°)</td>
<td>98.12(2)</td>
<td>98.14(3)</td>
</tr>
<tr>
<td>(Ca$_{9-x}$Sr$_x$)$_3$Co$<em>9$O$</em>{27}$ $x=0.20$</td>
<td>4.8518(3)</td>
<td>4.8547(6)</td>
</tr>
<tr>
<td>$b_{ij}$ (Å)</td>
<td>2.8225(5)</td>
<td>2.821(1)</td>
</tr>
<tr>
<td>$b_{RS}$ (Å)</td>
<td>4.6304(8)</td>
<td>4.6362(8)</td>
</tr>
<tr>
<td>$c$ (Å)</td>
<td>10.8995(7)</td>
<td>10.904(2)</td>
</tr>
<tr>
<td>$\beta$ (°)</td>
<td>98.123(7)</td>
<td>98.12(2)</td>
</tr>
</tbody>
</table>

Because of lack of stability at high temperature, it was not possible to sinter these compounds by conventional sintering but dense ceramics were obtained by spark plasma sintering with relative densities higher than 96% as confirmed by SEM (see Fig. 4).

Fig. 3. Comparison of TGA and DTA corresponding to the Ca$_2$Co$_9$O$_{14}$ parent compound (CCO, black solid line), (Ca$_{9-x}$Sr$_x$)$_3$Co$_9$O$_{27}$ with $x=0.10$ (CSCO10, red line) and (Ca$_{9-x}$Sr$_x$)$_3$Co$_9$O$_{27}$ with $x=0.20$ (CSCO20 black doted line).

Fig. 4. SEM images of Ba$_2$Co$_9$O$_{14}$ (a), and (Ca$_{9-x}$Sr$_x$)$_3$Co$_9$O$_{27}$ with $x=0.10$ (b) and 0.20 (c) showing dense ceramics.

$^{18}$O/$^{16}$O isotope exchange in Ba$_2$Co$_9$O$_{14}$

The normalised $^{18}$O/$^{16}$O isotope concentration in function of penetration depth are given in Fig. 5 for the three studied temperatures. Both data recorded in depth-profile and line-scan mode are reported. A good junction between the two sets of data was
observed. However, whatever the temperature of exchange, a high oxygen content was measured at the surface of the samples in the first microns depth, the thickness of this high oxygen content layer increasing with the temperature. It was therefore not possible to fit these data to the diffusion equation [1].

The oxygen transport properties in the two strontium doped Ca$_3$Co$_4$O$_{9+}$ were characterised by both Electrical Conductivity Relaxation and the $^{18}$O/$^{16}$O Isotope Depth Profile Technique. The electrical conductivity at thermodynamic equilibrium corresponding to the x=0.10 composition under different oxygen pressures is given in Fig. 6a. An increase of the electrical conductivity when the oxygen partial pressure increased was observed, confirming the p-type behaviour of these materials. As shown in Fig. 6b, the electrical conductivity slightly decreased when strontium was introduced in the structure. The evolution of electrical conductivity corresponding to the (Ca$_{1-x}$Sr$_x$)$_3$Co$_4$O$_{9+}$ (x=0.1) composition at 750°C with surrounding atmosphere changes is shown Fig. 7. It is worth noting that oxidation and reduction are reversible processes.

Surprisingly, despite very good electrochemical performances, rather low oxygen diffusion and kinetics toward oxygen molecule dissociation were shown. However, one must be careful with these results. The increase of $^{18}$O at the sample surface is not clearly understood. It is likely due to an oxidation process when cooling the sample but this hypothesis still has to be confirmed.

**Oxygen transport in strontium doped Ca$_3$Co$_4$O$_9$**

Data obtained for both compositions in the 650-750°C range were fitted to the theoretical model of diffusion. In these conditions, only the diffusion coefficient could be derived with precision, reflecting a $D_{chem}/k_{chem}$ characteristic length much smaller than the size of the sample and suggesting much higher surface exchange kinetics for these materials than chemical diffusion$^{28}$. Corresponding $D_{chem}$ values are given in Table 4 shown in Fig. 8. $D_{chem}$ values obtained from the air↔0.05 atm and air↔0.01 atm are close and similarly to what was obtained for the pure material$^{20}$, moreover the chemical diffusion coefficients are independent of whether the relaxation is a reduction or an oxidation process.

As shown in Fig. 8, as expected the diffusion process is thermally activated and activation energies of 1.03±0.03eV and 0.98±0.04eV were measured for the x=0.1 and x=0.2 compositions, respectively, slightly lower than the activation energy displayed by the pure compound (1.2±0.02eV) indicating easier diffusion for the strontium doped compound which also exhibit higher oxygen diffusion coefficients.
This increase of oxygen diffusion coefficient with substitution ratio was confirmed by $^{18}$O/$^{18}$O isotope exchange. The normalized $^{18}$O diffusion profile corresponding to the ($\text{Ca}_{x} \text{Sr}_{1-x})\text{Co}_{O_{3+\delta}}$ with $x=0.1$ composition, exchanged at 649°C during 567 minutes under 211 mbar dry oxygen is shown in Fig. 9. Again a good junction was obtained between data collected in depth profile mode and line scan mode.

However in this case, as shown for the parent compound$^{20}$, a small decrease of the $^{18}$O content was noticed at the sample surface, attributed to a possible carbonation or hydroxilation when back to ambient atmosphere. In insert, the same plot in decimal logarithm scale has been drawn to compare experimental and theoretical profiles far in depth. As previously noticed for the pure compound, a "tail" can be seen, suggesting most likely an anisotropic diffusion$^{20}$. Nevertheless, as shown in Fig. 9, a good fit between the calculated and experimental data was obtained. The derived $k^*$ and $D^*$ parameters are given in Table 5. Their evolution as a function of temperature is given on an Arrhenius plot in Fig. 10.

The tracer diffusion can be correlated to chemical diffusion via the thermodynamic factor $\gamma$ according to the following equation,

$$D_{\text{chem}} = \gamma \cdot D^*$$

where $\Delta \ln p_{\text{O}_2}$ and $\Delta \ln c_{\text{O}}$ are the corresponding variations in the oxygen partial pressure and oxygen content, and $\sigma_e$ and $\sigma_i$ are the electronic and ionic conductivity, respectively.

For sake of comparison, the evolution of $D_{\text{Eyr}}^*=D_{\text{chem}}/\gamma$ is also reported in Fig. 10. Values are in good agreement. The slightly higher values observed for $D_{\text{Eyr}}^*$ compared to the tracer diffusion coefficient values can be explained by a lack of accuracy in the determination of the thermodynamic coefficient because of small oxygen stoichiometry variations (see value of the thermodynamic coefficient in Table 6).

Table 6. ($\text{Ca}_{x} \text{Sr}_{1-x})\text{Co}_{O_{3+\delta}}$) thermodynamic coefficient derived from TGA analysis as a function of temperature and oxygen partial pressure.

<table>
<thead>
<tr>
<th>Temp. (°C)</th>
<th>$x=0.1$</th>
<th>$x=0.2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>650</td>
<td>211</td>
<td>313</td>
</tr>
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<td>750</td>
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An increase of the diffusion coefficient with the strontium content is confirmed and may be explained by the increase of the rock salt layers volume which may facilitate the oxygen
mobility within these layers. However, with a value in the range of $10^{-10}$ cm$^2$.s$^{-1}$ it is still low which justify the addition of gadolinia doped ceria to improve electrochemical performances.$^{15-19}$

With surface exchange coefficient of $3.0 \times 10^{-6}$ cm.s$^{-1}$ and $4.0 \times 10^{-6}$ cm.s$^{-1}$ at 700°C for both $x=0.1$ and $x=0.2$ compositions, these materials exhibit very fast kinetics for the oxygen reduction reaction compared to the best cathode materials in the field since values of only $1.3 \times 10^{-7}$, $1.1 \times 10^{-6}$ and $1.3 \times 10^{-7}$ were reported for La$_2$NiO$_{4+\delta}$ at 700°C$^{29}$, La$_{0.6}$Sr$_{0.4}$CoO$_{2+\delta}$ at 680°C$^{4}$ and GdBa$_2$O$_{5+\delta}$ at 686°C$^{30}$, respectively.

![Fig. 10. Arrhenius plot of $k^*$ and $D^*$ compared to $D_{\text{chem/\gamma}}$ for composition (Ca$_{1-x}$Sr$_x$)$_3$CoO$_{4+\delta}$ with $x=0.1$ (a) and $x=0.2$ (b).](image1)

To go further in the understanding of oxygen transport in these materials, the $x=0.1$ composition was characterised by LEIS. The initial spectrum corresponding to an estimated removal of 0.2 monolayer is compared to the one measured after an estimated removal of 3 monolayers in Fig. 11. Despite a bulk contamination with sodium which was also observed at the surface of powders and ceramics of the parent compound$^{20}$, mainly strontium and calcium were initially observed at the uppermost sample surface. No cobalt was initially observed, just a sub-surface signal at low energy side from the surface peak$^{31}$ in the background indicating cobalt atoms are beneath the first measured atomic layer. After ionic sputtering, cobalt was evidenced at the outermost surface and an increase of calcium was also noticed. Although, one must be careful with this technique since strontium may also segregate at the sample surface, the same behaviour was observed for other compounds. For SmCo$_3$, Fullarton et al. also showed that samarium was preferentially at the surface with only 5% of cobalt at the outmost surface and a segregation of strontium at the surface was evidenced for strontium doped compounds$^{32}$. Similarly, Viitanen et al. showed the absence of cobalt and iron in the outermost atomic layer of a La$_{0.6}$Sr$_{0.4}$Co$_{0.8}$Fe$_{0.2}$O$_{3}$ membrane$^{33}$. We also showed calcium oxide at the uppermost surface of the Ca$_3$Co$_2$O$_7$ parent compound. Here, strontium seems to be mainly at the sample surface and its presence may partly explain the increase of kinetics towards the oxygen dissociation reaction when strontium content increases.

![Fig. 11. $^4$He$^+$ (3keV) LEIS spectra recorded on a dense ceramics of (Ca$_{1-x}$Sr$_x$)$_3$CoO$_{4+\delta}$ with $x=0.1$ after annealing at 800°C for 12 hours prior to the analysis. The first spectrum (in black) corresponds to an estimated removal of 0.2 monolayer and the last spectrum (in red) to an expected removal of around 3 monolayers.](image2)

Conclusions

Although oxygen diffusion was confirmed in the promising Ba$_2$Co$_2$O$_7$, it was not possible to derive accurate transport parameters because of an oxidation process at the sample surface which has still to be clarified. In contrast, for Ca$_3$Co$_2$O$_{4+\delta}$ derivatives, it was shown that oxygen diffusion was improved when calcium was partly substituted with strontium likely due to an increase of the volume of the rock salt layers of the structure which may help for oxygen mobility. Although diffusion coefficient remains low, interestingly fast kinetics towards oxygen dissociation reaction were shown with surface exchange coefficient higher than those reported for the best cathode materials in the field. The surface exchange coefficient is increased when the strontium content increases which may be correlated to the presence of a higher amount of strontium at the sample surface which may play a key role in the mechanism of the oxygen molecule dissociation at the surface of the solid.

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**Notes**

*a* Laboratoire Structures, Propriétés et Modélisation des Solides (SPMS), Ecole Centrale Paris, Grande Voie des Vignes, 92295 Châtenay-Malabry, France

*b* Univ Lille Nord de France, CNRS UMR 8181, Unité de Catalyse et de Chimie du Solide - UCCS, ENSCI, Université Lille 1, CS 90108, 59652 Villeneuve d’Ascq Cedex, France

*corresponding author, email: rose-noelle.vannier@ensc-lille.fr, Tel +33(0)320436583, Fax +33(0)320436814*

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