New solid state oxygen and hydrogen conducting materials.
Towards their applications as high temperature electrochemical devices and gas separation membranes

Thesis submitted by
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To apply for the Degree of Doctor

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Valencia, July 2013
A mis padres y a mi hermana

A Sergio
"But I don’t want to go among mad people," Alice remarked. "Oh, you can’t help that," said the Cat: "we’re all mad here. I’m mad. You’re mad."

"How do you know I’m mad?" said Alice. "You must be," said the Cat, or you wouldn’t have come here."

Lewis Carroll, Alice in Wonderland
Outline

Context and Scope of the thesis ............................ 11

Chapter 1. Introduction ........................................ 15

1.1. Introduction to mixed ionic-electronic conductors (MIEC) ...... 17
1.2. Defect Theory .............................................. 19
1.3. Applications .................................................. 33
    Oxygen separation membranes .................................. 33
    Solid oxide fuel cells ........................................... 37
1.4. References .................................................... 50

Chapter 2. Methodology ........................................ 59

2.1. Material synthesis and sample preparation ...................... 61
    Material synthesis ............................................. 61
    Sample preparation ............................................. 63
2.2. Structural characterization .................................. 66
2.3. Thermal Analysis ........................................... 75
2.4. Electrochemical characterization ................................ 78
2.5. References .................................................... 85

Chapter 3. Structural-transport properties relationships on
Ce$_{1-x}$Ln$_x$O$_{2-δ}$ system (Ln = Gd, La, Tb, Pr, Eu, Er, Yb, Nd) and effect of
cobalt addition ..................................................... 89

3.1. Introduction ................................................... 91
3.2. Results and discussion ...................................... 95
3.3. Conclusions .................................................. 114
3.4. References .................................................... 115

Chapter 4. Microstructure optimization of Ce$_{0.9}$Pr$_{0.1}$O$_{2-δ}$ +Co 2mol% 121

4.1. Introduction ................................................... 123
4.2. Results and discussion

4.3. Conclusions

4.4. References

Chapter 5. Study of the bulk transport properties and oxygen surface exchange of the mixed ionic electronic conductor Ce$_{1-x}$Tb$_x$O$_{2.8}$ + Co (x=0.1, 0.2, 0.3, 0.5)

5.1. Introduction

5.2. Results and discussion

5.3. Conclusions

5.4. References

Chapter 6. LSM-ceria composite cathodes for SOFC

6.1. Introduction

6.2. Results and discussion

6.3. Conclusions

6.4. References

Chapter 7. Ionic transport improvement on CaTiO$_3$ based perovskites

7.1. Introduction

7.2. Results and discussion

7.3. Conclusions

7.4. References

Chapter 8. Ceria-based stable dual-phase membranes for fast oxygen separation

8.1. Introduction

8.2. Results and discussion

8.3. Conclusions

8.4. References
Chapter 9. Mixed protonic-electronic conductors based on lanthanum chromites

9.1. Introduction

9.2. Results and discussion

9.3. Conclusions

9.4. References

Chapter 10. Conclusions and remarks

10.1. Final remarks

Scientific Contribution

List of Figures and Tables

Appendix

Appendix 1

Appendix 2

Appendix 3

Appendix 4

Resumen / Resum / Summary

Acknowledgments/Agradecimientos
Context and Scope of the Thesis

Increasing energy demand in a world where economy and sustainability are claimed to come together has given a boost to renewable sources of energy that avoid the emission of greenhouse gases. Nevertheless, under a global vision the majority of constructed power plants are still working with fossil fuels, especially coal-fired as it is cheaper and more abundant than oil and natural gas. The concentration of atmospheric CO$_2$ has recently reached 400 ppm, far higher than the safety limit of 350 ppm and with the last precedent 3 million years ago. Therefore, decreasing the emission of greenhouse gases and especially CO$_2$ is a main issue that has gained great focus.

Up to date, a variety of mixed electronic and ionic (oxygen ions or protons) conducting materials are being investigated to separate oxygen or hydrogen from reforming streams at high temperature. The separation occurs through dense ceramic oxides providing selectivities close to 100%. These solid oxide materials may be used in the production of electricity from fossil fuels (coal or natural gas), taking part of the separation and storage of CO$_2$ system. Dense oxygen transport membranes (OTM) can be used in oxyfuel combustion plants or in catalytic membrane reactors (CMR), while hydrogen transport membranes (HTM) would be applied in precombustion plants. Furthermore, these materials may also be used in components for energy systems, as advanced electrodes or electrolytes for solid oxide fuel cells (SOFC) and proton conducting solid oxide fuel cells (PC-SOFC) working at high and moderate temperature.
However, the most widely studied materials for these purposes present a series of drawbacks that make them impractical for application to gas separation. The harsh working conditions established by the targeted processes include high temperatures and low O$_2$ partial pressures ($pO_2$), probably combined with CO$_2$ and SO$_2$ containing gases. Thus, the need to discover new stable inorganic materials providing high electronic and ionic conductivity is still present.

This thesis proposes a systematic search for new mixed ionic-electronic conductors with different crystalline structure and/or composition of the crystal lattice, varying the nature of the elements and the stoichiometry of the crystal. Hence, the main purpose of this thesis is to find breakthrough solid materials stable at high temperature (600-1000 °C) which are sufficient high mixed electronic and ionic conductors, either oxygen ion or proton conducting. This main objective may break-down in several parts:

- to obtain stable crystalline phases with possible ion conduction paths through the crystal lattice. The considered crystalline phases include mixed oxides of the family of the perovskites, fluorites and spinels. Besides, physical mixtures of crystalline phases are taken into account (fluorite-spinel composites)
- to develop a technique for evaluating transport characteristics of a set of materials simultaneous and automatically
- to find and optimize new mixed ionic-electronic materials with high ionic conductivity ($\sigma>0.05$ S/cm) and high chemical stability at high temperature in the presence of CO$_2$ and SO$_2$ and at low oxygen partial pressures
- to find and optimize new mixed protonic-electronic conductors ($\sigma>0.005$ S/cm) with high temperature and chemical stability in the presence of CO$_2$ and trace sulfur.
The best candidates are studied in depth to understand their properties and their possible applications in gas separation, catalytic membrane reactors and SOFC components.

The obtained results are structured as follows. Chapter 1 of the thesis briefly describes the fundamental principles of solid oxides chemistry and the state of the art on the applicability of the mixed ionic-electronic conducting materials. Chapter 2 collects the procedures developed for the material synthesis and specimen fabrication. The techniques employed to characterize the samples and to determine their transport properties are outlined as well as the different setups used for the electrochemical characterization.

From this point, the following chapters may be divided in two main blocks. The first block comprises from Chapter 3 to Chapter 8 and deals with oxygen ion conducting materials. The second block, Chapter 9, is devoted to proton conducting perovskites.

In Chapter 3, a large series of lanthanide-doped ceria is characterized in order to have an overall understanding of the structural, oxygen vacancy concentration and transport properties. Besides, the addition of cobalt oxide on this system is similarly studied. The chosen lanthanides include a large range of ionic radii and different metals exhibiting variable oxidation states under the typical operating conditions encountered for these materials. Transport properties are correlated with structural features induced by the different ionic radii and the variable oxidation state of the dopants. Chapter 4 completes the study of the transport properties dependence on the microstructure. The sintering temperature of Ce0.9Pr0.1O2-δ +Co 2mol% is systematically modified in order to observe the influence in particle size, CoOx location and total conductivity. In Chapter 5, the structural and bulk ionic and electronic transport properties of Tb-doped ceria are evaluated as a function of Tb concentration.
Surface properties are also reviewed. The potential use of these materials as high-temperature oxygen transport membrane is assessed. Finally, Chapter 6 includes additional metal dopants on the Ce_{0.9-x}M_{x}Gd_{0.1}O_{2.4} structure with the aim of modifying the transport properties. The performance of these materials as a component of La_{1-x}Sr_{x}MnO_{3}-based composite cathode is evaluated.

The second family of materials comprises CaTiO_{3}-based perovskites. Chapter 7 reports the search of dopants either in A or B site that provide stable perovskites with high mixed ionic electronic conductivity. The screening drives to iron and iron-magnesium doped CaTiO_{3} as best candidates, which are further structural and electrochemically studied.

Chapter 8 describes the development of composite materials composed of Fe_{2}NiO_{4} spinel material, combined with several lanthanide-doped cerias. The optimization of the synthesis method and the composition is reported as well as the electrochemical characterization and stability of the material.

Chapter 9 focuses on proton conducting perovskites based on La_{0.85}Sr_{0.15}CrO_{3}. Doping strategies that aim to improve the ionic or protonic conductivity are evaluated by DC-conductivity in reducing conditions. The applicability of these materials is evaluated and the best candidates are tested as anode material for proton conducting SOFC.

Finally, general conclusions are extracted taking into account the objectives above described.
Chapter 1. Introduction

Abstract

The present chapter shows a global vision about the defect chemistry of mixed ionic electronic conductors (MIEC). The main defect formation reactions are described. Secondly, the fundamentals of membrane and oxygen ion or proton conducting solid oxide fuel cells are explained, as well as their main requirements. Besides, a brief literature overview on the state of the art of the materials used for these applications is given.
1.1. Introduction to mixed ionic-electronic conductors (MIEC)

Mixed ionic–electronic conductors (MIECs) are materials that conduct both ions and electronic charge carriers (electrons and/or holes). Normally, MIECs are ionic semiconductors, though the definition also holds for metals, glasses and polymers. [1] Besides, mixtures of two phases, one conducting ions the other conducting electrons/holes are considered as heterogeneous MIECs, named composites. The ionic species in solids may be oxygen anions, protons or metal cations, from which the materials presented in this work are oxygen ionic and protonic conducting oxides. The use of the term MIEC is usually reserved to those materials in which the ionic and electronic conductivities ($\sigma_i$, $\sigma_e$) neither differ by more than 2 orders of magnitude nor release too low values ($\sigma_i, \sigma_e > 10^{-5} \Omega^{-1}\cdot\text{cm}^{-1}$). [2] In fact, the conductivity of a material is proportional to the mobility and concentration of the carrier species (ions, vacancies, electrons or holes), which are examined below.

Ionic conduction in solids occurs by the movement of oxide ions through a crystal lattice, where ionic species of comparable size and opposite charge to the ions exist. In the crystal matrix, an empty site in the way of the ion is needed, which must have enough energy to overwhelm the barrier impeding the diffusion. Besides, these conditions have to be fulfilled continuously all along the sample. Thus, the movement of the ions from one to the other side of the material takes place via interstitial sites or by hopping into a vacant site or a complex combination of both interstitial and vacant sites. [3] There are three different ways to create ionic defects in a material (i) intrinsically, by thermal
excitation, and extrinsically (ii) by changing the stoichiometry via modifications of the oxygen sublattice, or (iii) by doping with aliovalent cations.

Thermal excitation causes the smaller ion (usually the cation) to be displaced from the lattice position to an interstitial site, which create a Frenkel pair or Schottky disorder as illustrated in Figure 1.1. Frenkel pairs consist of a vacancy defect at its original site and an interstitial defect at its new location. Schottky disorder involves an equal number of cations and anions moving from the lattice to the surface. On the other hand, the change of stoichiometry modifies the composition by interaction with the environment, as introduces oxygen vacancies or interstitial oxygen in the structure, e.g. CeO$_{2-\delta}$ and Sr$_4$Fe$_4$Co$_2$O$_{13+\delta}$. Finally, the strategy of doping may introduce mobile interstitial donors or mobile vacancies on the anion sublattice, which are mobile at elevated temperatures.

![Figure 1.1. a) Frenkel defect: cation in interstitial hole, and b) Schottky disorder: missing pair of ions.](image)

Electronic (electron/hole) conductivity occurs via delocalized states in the conduction/valence band or via localized states by a thermally assisted hopping mechanism. Similarly to the ionic conductivity, there are three ways of creating these charge carriers, (a) thermal excitation, which generates an electron-hole pair across the band gap, (b) deviation from stoichiometry that
introduces native defects that may act as donors (oxygen vacancies) or acceptors (interstitial oxygen), and (c) doping, which also introduce donors or acceptors. [4] In some cases, unless donor level lies above the bottom of the conduction band, the electronic conduction by doping may require also thermal excitation.

It is evident that the defect concentration in an ionic or electronic conducting material is subjected to external factors such as temperature, oxygen partial pressure \((pO_2)\) and other thermodynamic parameters. In fact, some ionic conducting materials become \(p\) or \(n\)-type electronic conductors in certain environments. As an example, ionic conductor \(\delta-Bi_2O_3\) becomes an \(n\)-type electronic conductor at low \(pO_2\). [5, 6] In the majority of applications of MIEC, they appear in the form of polycrystals; consequently, the grain boundaries (GB) are critical for the microstructure and the conductivity. The GB misorientation angles result in inter-grain blocking effect that may come from two natures (1) intrinsic charge barriers in grain-to-grain contacts, which is effectively explained by the space-charge effect where oxygen vacancies are highly depleted and (2) extrinsic segregation of impurities that block the transport of ions. [7, 8] Increasing the temperature and doping concentration minimize the ‘intrinsic’ blocking effect, while ‘extrinsic’ blocking effect may be still present up to 1000 °C. [7, 9] Thus, all these factors should be taken into account. An accurate defect model could allow the systematic optimization of the electrical and transport properties of a material through cation substitution (doping), as well as the prediction of the behaviour under operating conditions. In the next section, an explanation of the defect generation is detailed.

1.2. Defect Theory

Key defect reactions containing the dominant electronic and ionic defects in an oxygen deficient oxide include oxidation/reduction, intrinsic lattice
defect generation, intrinsic electron-hole generation, and ionization reactions as well as overall charge and mass balance relations. The reaction equations and their respective mass action laws are listed below. [10]

The intrinsic disorder is integrated in the equilibrium state of crystals. It is not related to impurity content or any other extrinsic origin such as deviation from stoichiometry. To achieve the equilibrium concentration of defects, the temperature needs to be sufficiently high to reach the equilibrium in reasonable times. All the movements inside a crystal that cause the lattice disorder must satisfy several conservation rules, i.e., conservation of mass, charge and structure (lattice site ratios). There are different types of intrinsic lattice disorder consistent with the conservation rules, although one type uses to be favoured compared with all others in a real case. For the compounds studied in this thesis, mainly ionic binary oxide materials, with high co-ordination number and small difference in sizes of cations and anions, the major types of intrinsic ionic disorder (schematized in Figure 1.1) are anion Frenkel (also called Anti-Frenkel) and Schottky disorder, whose defect reaction equations and their respective mass action laws are written below in Kröger-Vink notation (Eqs. 1.1-1.4). [11]

\[ O_0^\chi \rightarrow V_0^- + O_i'' \]  
\[ K_F = [V_0^-][O_i''] = k_F \exp \left( \frac{-E_F}{kT} \right) \]  
\[ \text{null} \rightarrow 3V_0^- + V_{M1}'' + V_{M2}'' \]  
\[ K_S = [V_0^-]^3 [V_{M1}''][V_{M2}''] = k_S \exp \left( \frac{-E_S}{kT} \right) \]

where \( O_0^\chi \) is an occupied oxygen site, \( V_0^- \) is an oxygen vacancy, \( O_i'' \) is an oxygen interstitial, \( V_{M1}'' \) and \( V_{M2}'' \) are the vacancies in the metals M1 and M2. Notation in brackets denotes the concentration of corresponding species and
$K_F(T)$ and $K_S(T)$ are the equilibrium constants defining the anion Frenkel and Schottky reactions, respectively.

The intrinsic electronic disorder involves thermal excitation of electrons from chemical bonds to higher energy states. In a stoichiometric solid oxide, the ground state is a nominally filled valence band and a nominally empty conduction band. The excitation of an electron across the bandgap not only allows it to move freely through the empty states of the conduction band, but also leaves an empty electronic state at the top of the valence band. This hole can also transport charge as the electrons can move laterally into its empty state, in the opposite direction of electrons moving in the conduction band. Hence, the equilibrium reaction for the electronic excitation process that leaves behind an empty state must complete the conservation rules, and will be written as Eq. 1.5.

$$null \rightarrow h^\prime + e'$$

(Eq. 1.5)

where $e'$ is an electron in the conduction band and $h'$ represents a hole in the valence band. So the mass action equation for this process is written as follows:

$$K_E = np = k_E \exp \left( \frac{-E_g}{kT} \right)$$

(Eq. 1.6)

where $n = [e']$ and $p = [h']$ are the widely used shorter notation representing the concentration of negative and positive carriers, respectively. $E_g$ is the energy of the bandgap that needs to be overwhelmed to create the disorder.
In the case of solid oxides where the motion of electrons is more restricted, the electronic transport occurs by discrete jumps between localized electrons via the so called hopping. Electronic defects in an ionic crystal induce a distortion in the lattice, which becomes polarized and traps the charge carrier inside the electrostatic field (*self-trapping*) as depicted in Figure 1.2a. The electron or hole trapped in a particular atom, ion or vacancy is considered as a particle named ‘**polaron**’, which can propagate throughout the crystal by hopping from one atom to another. Two kind of polarons can be distinguished depending on their binding energy. If the interaction with the lattice is strong and the polaron is localized, it is called ‘**small polaron**’ and the lattice distortion reaches distances close to the cell parameter. The small polaron hopping occurs between sites of optic phonon frequency, causing the low mobility (less than 0.1-1 cm² V⁻¹ s⁻¹). By increasing temperature, the random hopping is activated so the small polaron mobilities rise. On the other hand, a weak interaction with the lattice produces longer distance distortions, which is known as ‘**large polaron**’, typical of transition metal oxides where metals may be in different oxidation states, *e.g.* in olivine [(Mg₁₋ₓFeₓ)₂SiO₄], charge carriers are electrons transferred from Fe²⁺ to Fe³⁺ ions. [13, 14] Large polaron mobilities are high at low temperatures (1.75x10⁵ cm² V⁻¹ s⁻¹), which decrease with rising temperature.
Introduction

through phonon scattering vibrations. [15] Focusing on carrier mobilities, the distinction between small and large polarons is shown in Figure 1.3.

The temperature dependence of the mobility is often of the form

\[ \mu = \mu_0 T^{-3/2} e^{-E_A/kT} \]  
(Eq. 1.7)

The localized centers hosting the electrons or holes may be lattice cations or anions or impurity centers if they are close enough to each other.

![Diagram](image)

Figure 1.3. Electron (e), and hole (h), mobilities in oxides. The lower limit for large polarons corresponds to a mean-free path equal to the interatomic distance. The upper limit for small polarons corresponds to one hop every lattice vibration period. Note the opposite trends with temperature: for small polarons and ions, thermal activation is needed, so higher temperatures raise the rate; for large polarons, phonon scattering reduces the mobility at higher temperatures. [13]

Extrinsic ionic disorder involves the ionic defects that result from the presence of impurity ions dissolved in the lattice, forming a substitutional solid solution. If an isovalent ion replaces the host element, there is not need of creating other compensating defect and the properties of the solid solution are similar to the origin. Aliovalent ions carry a charge with respect to the perfect
crystal, so that other oppositely charged defect must be present to keep the neutrality, *i.e.*, an ionic defect. There are two possible compensating ionic defects, named vacancies or interstitials, which are different depending on the type of impurity. Donor impurities (D, positively charged) promote either cation vacancies or anion interstitials, while acceptor impurities (A, negatively charged) forms either cation interstitials or anion vacancies. Only one of these ionic defects results predominant from the impurity content. Features as electrostatic environment, space availability or formal charge on the defect will determine the most probable compensating defect.

In a material that exhibits different sort of defects, oppositely charged, the possibility of electrostatic attraction gives rise to the defect complexes and associates formation. These defects break under increasing temperature, so that their enthalpy of formation is negative. The defect complexes bind an impurity center with an oppositely charged ionic defect (sited on nearest-neighbour cation sites) turning it into a neutral inmobile complex. The subsequent drop of mobile charges produces and accelerated decrease in ionic conductivity with decreasing temperature.

In a more real scenario, it is necessary to consider that the crystal is not isolated and the thermodynamic equilibrium with the surrounding needs to be reached. This is achieved by means of electronic compensation, so that extrinsic electronic disorder appears. The interaction of doped crystals with the gaseous ambient leads to gain or loss non-metallic constituent until the ionic defects introduced by an impurity or a dopant have been eliminated and replaced by electrons or holes of equivalent charge. As a consequence, the material becomes nonstoichiometric. When doping with donor impurities the compensation occurs via the electrons left after the loss of the oxygen surplus relative to the host lattice. Acceptor dopants posses less oxygen per cation than host oxide, so that
neutral external oxygen will enter in the lattice by taking two electrons from the valence band and leaving two compensating holes as overall result. [10]

Summarizing the effects of these mentioned cases in which an impurity or doping element is added, an ionization equation must be taken into account, different whether the dopant is an acceptor or donor:

**Acceptor dopant**

\[ A \rightarrow A^+_c + h' \]

\[ K_{Ac} = [A^+_c]p = k_{Ac}e^{\frac{-E_A}{kT}} \]

**Donor dopant**

\[ D \rightarrow D^- + e' \]

\[ K_{Dn} = [D^-]n = k_{Dn}e^{\frac{-E_D}{kT}} \]

Each system will show those defects corresponding to the most favourable energetic situation. These charge-balancing defects may be used to tailor the transport properties of the material.

On the other hand, the reaction of a semiconductor compound with the surrounding gas phase components gives rise to intrinsic nonstoichiometry. The nonstoichiometry denotes a variable composition of the same crystal structure that must be adjusted with the environment by a combination of ionic and electronic defects.

**Reduction reaction**

In reducing atmospheres, the oxide material reacts with the atmosphere by loosing oxygen.
\[ O_0^{\overline{X}} \rightarrow V_0^- + 2e^- + \frac{1}{2} O_2 \]  
(Eq. 1.12)

The atomic defects generated in an occupied oxygen site, \( O_0^{\overline{X}} \), are oxygen vacancies that retain two electrons from the doubly ionized oxygen, \( V_0^- \). The mass action law is written in Eq. 1.6

\[ K_R = [V_0^-] n^2 pO_2^{\frac{1}{2}} = k_R \exp \left( \frac{-E_v}{kT} \right) \]  
(Eq. 1.13)

where \([V_0^-]\) and \(n\) are the oxygen vacancy and electron concentration, \(pO_2\) is the oxygen partial pressure and \(K_r(T)\) is the equilibrium constant defining the reaction.

**Oxidation reaction**

In oxidizing atmospheres, the oxide material reacts with the atmosphere by incorporating oxygen in the lattice.

\[ \frac{1}{2} O_2 + V_0^- \rightarrow O_0^{\overline{X}} + 2h^- \]  
(Eq. 1.14)

\[ K_D = [V_0^-]^{-1} n^2 pO_2^{-\frac{1}{2}} = k_D \exp \left( \frac{-E_d}{kT} \right) \]  
(Eq. 1.15)

Finally, the electroneutrality condition inside the crystal has to be maintained, which is represented by a general equation incorporating all the charged defect species mentioned above. It can be written as Eq. 1.16 and Eq. 1.17 for Frenkel and Schottky dominion:

\[ n + 2[O^{\prime \prime}] + [A'] \rightarrow p + 2[V_0^-] + [D'] \]  
(Eq. 1.16)

\[ n + [V_A^+] + [A'] \rightarrow p + 2[V_0^-] + [D'] \]  
(Eq. 1.17)
If the aliovalent impurity or dopant content exceeds the concentration of defects resulting from intrinsic disorder, it gives rise to extrinsic non-stoichiometry. Combination of the equations written above describes a way to evaluate the predominant type of carriers, i.e., the concentration of electrical species present at a given temperature, $pO_2$ and dopant level. As not all the defects occur simultaneously it is possible to simplify the equations including only one defect species on either side of the equality (Eqs. 1.18-1.21). Hence, restricted ranges of $pO_2$ and temperature usually define four defect regions dependent on the external conditions for acceptor and donor doped materials, i.e., reduction, ionic compensation, electronic compensation and oxidation regions, summarized in Table 1.1.

<table>
<thead>
<tr>
<th>Region</th>
<th>Acceptor</th>
<th>Donor</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Reduction</td>
<td>$n \approx 2[V_0^+]$</td>
</tr>
<tr>
<td>II</td>
<td>Ionic compensation</td>
<td>$2[V_0^+] \approx [A']$</td>
</tr>
<tr>
<td>III</td>
<td>Electronic compensation</td>
<td>$p \approx [A']$</td>
</tr>
<tr>
<td>IV</td>
<td>Oxidation</td>
<td>$p \approx 2[O_{2}^{2-}]$</td>
</tr>
</tbody>
</table>

Table 1.1. Summary of electroneutrality equations in different regions.

Substituting these approaches solves the mentioned equation system, so that the temperature and $pO_2$ dependence of the electrical species may be obtained. These dependencies are used in the analysis of the defect chemistry from the raw electrical data of the different systems studied in this thesis.

For an acceptor doped material, the approximation for the reduction reaction dominant region (I) indicates that the electron generation is compensated by doubly ionized oxygen vacancies. When it is substituted in Eq. 1.13 gives the following relation:

$$n = \left(2K_R\right)^{\frac{1}{2}} pO_2^{-\frac{1}{2}}$$  (Eq. 1.24)
Hence, the electrical conductivity at the reduction dominant region will have a \( pO_2^{-1/6} \) dependence due to electron conduction (i.e., electronic conductivity). By substituting Eq. 1.24 on the electron-hole formation equation, Eq. 1.6, it is possible to obtain \( p \) carriers dependence on \( pO_2 \):

\[
\begin{align*}
  p &= 2^{-\frac{1}{3}} K_R^{-\frac{1}{3}} K_e \ pO_2^{\frac{1}{6}} \\
  &\quad \text{(Eq. 1.25)}
\end{align*}
\]

Similarly, the equation for oxygen vacancies is obtained by substituting Eq 1.18 into Eq. 1.13:

\[
\begin{align*}
  [V_0^-] &= 2^{\frac{2}{3}} K_R^{-\frac{1}{3}} pO_2^{\frac{1}{6}} \\
  &\quad \text{(Eq. 1.26)}
\end{align*}
\]

Finally, the oxygen partial pressure dependence for oxygen interstitials comes from the substitution of Eq. 1.26 in Eq. 1.2:

\[
\begin{align*}
  [O_0^\prime] &= 2^{\frac{2}{3}} K_R^{-\frac{1}{3}} K_F pO_2^{\frac{1}{6}} \\
  &\quad \text{(Eq. 1.27)}
\end{align*}
\]

In region II, where less reduction conditions appear, the electrons left from the production of \( O_2 \) are invested in cation reduction. Thus, by substituting Eq. 1.19 in Eq. 1.13 the electronic density will be determined by:

\[
\begin{align*}
  n &= (2 K_R)^{\frac{1}{2}} \left[ A_e'\right]^{-\frac{1}{2}} pO_2^{-\frac{1}{4}} \\
  &\quad \text{(Eq. 1.28)}
\end{align*}
\]

Analogous analyses can be done for the rest of \( pO_2 \) dependence of the electrical species in other regions, and the results are summarized in Table 1.2 for acceptor and Table 1.3 for donor doped materials.
However, pure electronic conduction is expected since the mobility of electrons is usually orders of magnitude higher than that of oxygen vacancies.

I. Ho observed that electrons and oxygen vacancies are the dominant defects in Region I. Concentration merely represents the ionized acceptor concentration. It can be material Kröger in the activity of a component of the system, usually representation of the variation in defect concentrations as a function of changes material.

<table>
<thead>
<tr>
<th>Defect</th>
<th>Region I</th>
<th>Region II</th>
<th>Region III</th>
<th>Region IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>( n )</td>
<td>((2K_R)^{\frac{1}{2}}pO_2)</td>
<td>((2K_R)^{\frac{1}{2}}[A'_c]^\frac{1}{2}pO_2)</td>
<td>([A'_c]^{-1}K_E)</td>
<td>((2K_FK_D)^{\frac{1}{2}}K_EpO_2)</td>
</tr>
<tr>
<td>( p )</td>
<td>((2K_R)^{\frac{1}{2}}K_EpO_2)</td>
<td>((2K_R)^{\frac{1}{2}}[A'_c]^\frac{1}{2}pO_2)</td>
<td>([A'_c])</td>
<td>((2K_FK_D)^{\frac{1}{2}}pO_2)</td>
</tr>
<tr>
<td>([V'_O])</td>
<td>(2^{-\frac{2}{3}}K_R^{\frac{1}{2}}pO_2)</td>
<td>(\frac{1}{2}[A'_c])</td>
<td>((2K_R)^{\frac{1}{2}}[A'_c]^{-\frac{1}{2}}pO_2)</td>
<td>((2K_F)^{\frac{1}{2}}[A'_c]^{\frac{1}{2}}pO_2)</td>
</tr>
<tr>
<td>([O''_O])</td>
<td>(2^{\frac{2}{3}}K_R^{\frac{1}{2}}K_EpO_2)</td>
<td>(2[A'_c]^{-1}K_F)</td>
<td>([A'_c]^{-2}K_OK_EpO_2)</td>
<td>(2^{-\frac{2}{3}}(K_OK_F)^{\frac{1}{2}}pO_2)</td>
</tr>
</tbody>
</table>

Table 1.2. Region dependent equation for defect species concentration in acceptor doped material.

<table>
<thead>
<tr>
<th>Defect</th>
<th>Region I</th>
<th>Region II</th>
<th>Region III</th>
<th>Region IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>( n )</td>
<td>((2K_R)^{\frac{1}{2}}pO_2)</td>
<td>([D_c])</td>
<td>((2K_RK_F)^{\frac{1}{2}}[D_c]^{-\frac{1}{2}}pO_2)</td>
<td>(2^{\frac{2}{3}}K_O^{-\frac{1}{2}}K_EpO_2)</td>
</tr>
<tr>
<td>( p )</td>
<td>((2K_R)^{-\frac{1}{2}}K_EpO_2)</td>
<td>([D_c]^{-1}K_E)</td>
<td>((2K_RK_F)^{-\frac{1}{2}}K_E[D_c]^{\frac{1}{2}}pO_2)</td>
<td>(2^{\frac{2}{3}}K_O^{-\frac{1}{2}}pO_2)</td>
</tr>
<tr>
<td>([V'_O])</td>
<td>(2^{-\frac{2}{3}}K_R^{\frac{1}{2}}pO_2)</td>
<td>([D_c]^{-2}K_RpO_2)</td>
<td>(2[D_c]^{-1}K_F)</td>
<td>((2K_F)^{\frac{1}{2}}K_O^{-\frac{1}{2}}pO_2)</td>
</tr>
<tr>
<td>([O''_O])</td>
<td>(2^{\frac{2}{3}}K_R^{\frac{1}{2}}K_EpO_2)</td>
<td>([D_c]^2K_R^{-1}K_EpO_2)</td>
<td>(\frac{1}{2}[D_c])</td>
<td>(2^{-\frac{2}{3}}(K_OK_F)^{\frac{1}{2}}pO_2)</td>
</tr>
</tbody>
</table>

Table 1.3. Region dependent equation for defect species concentration in donor doped material.

The Kröger-Vink diagram (also called Brouwer diagram) is a useful representation of the variation in defect concentrations as a function of changes in the activity of a component of the system, usually \(pO_2\). For an acceptor doped material Kröger-Vink diagram is plotted in Figure 1.4, where the acceptor concentration merely represents the ionized acceptor concentration. It can be observed that electrons and oxygen vacancies are the dominant defects in Region I. However, pure electronic conduction is expected since the mobility of electrons is usually orders of magnitude higher than that of oxygen vacancies.
Otherwise, in Region II, oxygen vacancies are maintained while the electron density decreases with increasing $pO_2$, thus becoming dominating defects. In Region III, holes are the dominant charge carriers. In Region IV, both holes and oxygen interstitials dominate, although conduction is likely by holes since their mobility is generally much higher than that of interstitials.

A similar Kröger-Vink diagram is depicted in Figure 1.5 for donor doped material (simply the ionized donor concentration is to be considered). In Region I, the electron is expected to be the dominant electrical carrier with a $pO_2^{-1/6}$ dependence. In Region II the concentration of ionized donors sets the electron density, so that it would be $pO_2$ independent. In Region III, where higher $pO_2$ is found, the electrical conductivity may switch to ionic towards the middle of this region if the intrinsic electron-hole product becomes low enough. Progressing to sufficiently high $pO_2$, the higher amount of holes will again shift the balance to electronic conduction. Therefore, in Region IV, the material is likely to be dominated by hole conduction, $pO_2^{1/6}$ dependent.
Figure 1.4. Kroger-Vink diagram for acceptor doped material. The acceptor concentration represents the ionized acceptor concentration.

Figure 1.5. Kröger-Vink diagram for donor doped material. The donor concentration represents the ionized donor concentration.
From this point, the conductivity obtained from these general systems will be the sum of the contribution of all charge carriers: \( t \), total; \( i \), ionic; \( n \), electronic and \( p \), holes (Eq. 1.29). For each defect, \( x \), that contribution is proportional to (i) the charge carriers, \( c \) (carriers/cm\(^3\)), (ii) to the charge that they carry, \( z_x e \) (C/carryer), being \( z_x \) the number of unit charges on the carrier and \( e \) the unit electron charge (1.6 x 10\(^{-19}\) C) and (iii) to the mobility \( \mu_x \), defined as the ability to move in an electric field (Eq. 1.30). Therefore, having into account Eq. 7, a general conductivity expression in an intermediate \( pO_2 \) range is given in Eq. 1.31.

\[
\sigma_T = \sigma_i + \sigma_n + \sigma_p \\
(\text{Eq. 1.29})
\]

\[
\sigma_T = 2e[V_o']\mu_{v_o'} + c_ne\mu_n + c_pe\mu_p \\
(\text{Eq. 1.30})
\]

\[
\sigma_T = \sigma_i^0 e^{(-E_i/kT)} + \sigma_n^0 (pO_2)^{-1/4} e^{(-\Delta H_{red}/kT)} + \sigma_p^0 (pO_2)^{+1/4} e^{(-\Delta H_{ox}/kT)} \\
(\text{Eq. 1.31})
\]

In a MIEC material, the conductivity data analysis permits to distinguish electronic an ionic components by their \( pO_2 \) dependency. As mentioned, the mobility of ions is far lower than the mobility of electronic carriers, so that ionic conductivity will be only observable if electron and hole concentration is minimum, that is, the \( pO_2 \) is about providing \( p = n \). If ionic conductivity in this range is higher than electronic, the experimental observation would be a conductivity independent on \( pO_2 \). Otherwise, \( n \) or \( p \) electronic conductivities would be evident by \( \sigma \propto pO_2^{-1/4} \) and \( pO_2^{+1/4} \) dependencies, respectively.

In conclusion, solid oxide materials are able to behave differently depending on their surrounding. This opens the possibility of tailoring both the structure and atmosphere to achieve desired properties for a number of
applications, *i.e.*, solid oxide fuel cells (SOFC), solid oxide electrolyser cells (SOEC) oxygen or hydrogen separation membranes, catalyst and gas sensors. The work presented in this thesis was focused in obtaining MIEC materials suitable for applications in SOFC components and oxygen separation membranes.

### 1.3. Applications

**Oxygen separation membranes**

Oxygen production at high temperature through ceramic membranes represents a current challenge that would enable the introduction of more efficient and environmental-friendly power generation and chemical production processes. In fact, reliable mass-scale $O_2$ separation in membrane modules would allow the cost-effective operation of *Oxyfuel* power plants, and integrated gasification combined cycle (IGCC) plants, which can integrate CO$_2$ capture and sequestration strategies in order to minimize SO$_2$ particulates and CO$_2$ emissions. [16, 17] Oxyfuel combustion principle is the removal of nitrogen from the oxidizer to carry out the combustion process in an oxygen-enriched gas stream, thus reducing gas volume and recycling the flue gas to lower the flame temperature. Hence, an air separation unit (ASU) has to be added to the power plant. The current state of technology development for ASU applies high-energy consuming cryogenic distillation units, while alternative oxygen separation technologies such as ion transport membranes would enable thermal integration and energy savings. [18] Furthermore, the development of catalytic membrane reactors (CMR) allows increasing the reaction conversion by shifting the equilibrium of the reaction. Several chemical processes as partial oxidation of
methane (POM) to synthesis gas, oxidative coupling of methane (OCM), water splitting, selective ammonia oxidation or NO\textsubscript{x} to N\textsubscript{2} conversion would benefit from the development of highly-permeable and CO\textsubscript{2} stable ceramic membranes. [19-25] The thermal integration of the membrane in the process permits its intensification, which involves several advantages with respect to conventional processes, \textit{i.e.}, the improvement of product selectivity and conversion, energy conservation, safer operation, reduced reactor/combustor volume and higher process performance.

For these reasons, many researches are devoted to find membrane materials that can work in the processes mentioned above. Membranes based on MIEC materials enable \textit{in situ} separation of oxygen in a membrane reactor fed by air. [26, 27] In MIEC ceramic membranes only oxygen ions are mobile within the ceramic lattice and all other gases, such as nitrogen, carbon monoxide, argon or methane are excluded to pass through the membrane to the permeate side. Thus, MIEC oxygen transport membranes are able to separate oxygen from air at high temperature in a process 100\% selective to oxygen being competitive with conventional cryogenic separation methods. In the opposite direction to the oxygen a continuous flux of electrons keeps the electroneutrality without any external circuit. The oxygen partial pressure difference across the membrane is the driving force for the membrane to work.
The mechanism of permeation in a ceramic membrane is depicted in Figure 1.6. In the feed side, oxygen molecules in air are dissociated and reduced to oxygen ions at the membrane reducing surface; the oxygen ions are then transported across the ceramic membrane to the oxidizing surface in the permeate side, where is combined back to oxygen gas. These three processes can be seen as a sum of series resistance (\(R = R_{\text{interface, feed}} + R_{\text{bulk}} + R_{\text{interface, permeate}}\)). When the kinetics of the membrane is dominated by the bulk diffusion \(R_{\text{bulk}}\) the oxygen transport across the membrane is given by the modified Wagner’s equation (Eq. 32). It predicts that the oxygen flux is proportional to (i) the temperature, (ii) the oxygen potential gradient, which is related to the \(pO_2\) difference between feed and permeate sides of the membrane, (iii) the ionic and electronic conductivities, \(\sigma_i\) and \(\sigma_{el} (\sigma_h + \sigma_e)\), respectively, and inversely proportional to membrane thickness, \(L\).

\[
j_{O_2} = \frac{RT}{16F^2L} \int_{p_{O_2,\text{feed}}}^{p_{O_2,\text{perm}}} \frac{\sigma_i \sigma_{el}}{\sigma_i + \sigma_{el}} d\ln pO_2
\]  

(Eq. 1.32)

Thus, decreasing the membrane thickness can increase the oxygen flux through the membrane. However, there is a certain thickness in which the diffusion process is sufficiently fast and the kinetics of the membrane becomes governed by the oxygen surface exchange rate at the gas/solid interfaces.
(R_{\text{interface}}). This value is known as the characteristic thickness $L_c$, which is defined by the ratio of oxygen diffusion constant $D \, (m^2 \, s^{-1})$ and oxygen surface exchange coefficient $k \, (m \, s^{-1})$, distinctive for each membrane material, as expressed in Eq 1.33.

$$L_c = \frac{D}{k} \quad \text{(Eq. 1.33)}$$

In order to decrease the resistances in the interfaces, oxygen reduction and oxidation catalyst may be attached in, respectively, feed and permeate sides of the membrane, as depicted in Figure 1.6.

Up to date, the most common type of material that exhibits both ion and electron conducting properties are perovskites e.g. $\text{Ba}_{1-x}\text{Sr}_x\text{Co}_{0.8}\text{Fe}_{0.2}\text{O}_3$ (BSCF) and $\text{La}_{1-x}\text{Sr}_x\text{Co}_{0.8}\text{Fe}_{0.2}\text{O}_{3.5}$ (LSCF), which enable oxygen fluxes up to $12.2 \, \text{ml} \cdot \text{min}^{-1} \cdot \text{cm}^{-2}$ (BSFC) and $0.5 \, \text{ml} \cdot \text{min}^{-1} \cdot \text{cm}^{-2}$ (LSFC) in 1 mm thick membranes measured at 900 ºC using argon as sweep gas. [28-33] However, the main disadvantage of these materials is the limited chemical stability under a large oxygen concentration gradient, with one side of the membrane exposed to air oxidizing atmosphere and the other side to reducing or CO$_2$ containing atmosphere. [34] Even at 300 ppm [35], CO$_2$ presence is a major issue for perovskite-based membranes (e.g. BSCF) due to the carbonation reaction of the earth alkali metals included in their structure. Improved stability against oxygen partial pressure gradients and carbonation is expected when avoiding Co alkaline-earth elements and doping by only transition metals and lanthanides instead.

On the other hand, fluorite materials such as stabilized bismuth oxide (SBO) or $\text{Ce}_{1-x}\text{Ln}_x\text{O}_{2.5}$ ($\text{Ln}=$lanthanide) have been attempted as oxygen separation membranes, as they are more stable in operational gases and against large oxygen concentration gradients. However, they are mainly ionic
conductors and the scarcity of electronic conductivity limits the oxygen permeation performance. By doping with multivalent cations, as terbium and praseodymium, mixed conduction in ceria may be further extended to higher oxygen partial pressure range (10^{-2}-10^{-1} atm), which enables its application as oxygen permeable membranes. As an example, Fagg et al. developed cobalt-doped Ce_{0.8}Pr_{0.2}O_{2-δ} membranes that offer low but competitive levels of oxygen permeation flux at temperatures below 850 °C when compared with perovskite materials, since they are thermally and CO₂ stable. [36]

As it is difficult to meet all the requirements in one single-phase material, dual phase composite membranes made from an oxygen-ion conducting oxide and a primarily electronic material (MIEC or metal) have been investigated, e.g. Ce₀.₈Gd₀.₂O₂₋δ (CGO) combined with La₀.₇Sr₀.₃MnO₃₋δ (LSM) or La₀.₈Sr₀.₂Fe₀.₈Co₀.₂O₃₋δ (LSFC), [37] or Er₀.₂₅Bi₀.₇₅O₃₋δ mixed with Ag, Pt or Au. Nevertheless, the interaction of properties was found to be detrimental for the total conductivity and oxygen permeability.

**Solid oxide fuel cells**

Fuel cells are the most efficient devices for power generation as they convert the chemical energy of fuel directly to electrical energy. Since the combustion process is avoided, fuel cells produce power with minimal pollutant exhausts. Fuel cells are mainly composed of a pure ion conducting electrolyte and two electrodes: an anode and a cathode. There exist different types of fuel cells, whose classification depends on the constituent electrolyte, i.e., polymer electrolyte membrane fuel cell (PEMFC), alkaline fuel cell (AFC), phosphoric acid fuel cell (PAFC), molten carbonate fuel cell (MCFC) and solid oxide fuel cell (SOFC).
In SOFC the electrolyte is a solid, nonporous metal oxide, which is a pure ionic (O$^{2-}$ or H$^+$) conductor at high temperature. It allows the fast conduction of ions between the two electrodes while the conductivity of electrons must be negligible. [38-40] The electrolyte is enclosed between two porous electrodes, the cathode, and the anode. Figure 1.7 shows the working scheme of a SOFC made by an oxygen ion conductor electrolyte that is fed by H$_2$ in the anode side and air or O$_2$ in the cathode. Oxygen atoms are reduced on the porous cathode surface by electrons, so that oxide ions flow through the ceramic electrolyte. In the porous anode, the oxide ions react catalytically with fuel (hydrogen) and release electrons, giving off water and heat. An external circuit conducts the electrons from the anode to the cathode and provides the electrical energy. [41]  

![Figure 1.7. Schematic of an individual SOFC based on oxygen-ion conductors fed by H$_2$](image-url)
The overall reaction for the process when H₂ is used as fuel can be written as

\[ \text{H}_2 + \frac{1}{2} \text{O}_2 \rightarrow \text{H}_2\text{O} \]  
(Eq. 1.34)

The chemical energy from this reaction is directly converted into electrical energy, so that the maximum electrical work \( W_{el} \) obtainable in a fuel cell operating at constant temperature and pressure is given by the change in Gibbs free energy \( \Delta G \) of the electrochemical reaction:

\[ W_{el} = \Delta G = -zFE \]  
(Eq. 1.35)

where \( z \) is the number of electrons participating in the reaction, \( F \) is Faraday’s constant \((96,487 \text{ C mol}^{-1})\), and \( E \) is the ideal potential of the cell.

The Gibbs free energy change is also given by the state function of the enthalpy change, \( \Delta H \) and the entropy change, \( \Delta S \):

\[ \Delta G = \Delta H - T\Delta S \]  
(Eq. 1.36)

The total thermal energy available is \( \Delta H \). The available free energy is equal to the enthalpy change less the quantity \( T\Delta S \), which is the energy consumed in the entropy change within the system. Reactions in fuel cells that have negative entropy change generate heat (such as hydrogen oxidation), while those with positive entropy change (such as direct solid carbon oxidation) may need an extra heat contribution if the irreversible generation of heat is smaller than the reversible absorption of heat. Thus, a fuel cell operating reversibly, i.e., the cell is in equilibrium without external current, will produce a heat equal to \( T\Delta S \). At this point, where all the Gibbs free energy is converted to electrical energy, it is possible to obtain the ideal equilibrium potential or open circuit voltage (OCV) of the fuel cell, \( (E) \).
By substituting Eq 1.35 in Eq 1.36 under conditions different than standard, it is possible to obtain Nernst equation, which relates the ideal standard potential \( E^o \) for the cell reaction and the ideal equilibrium potential at other partial pressures of reactants and products.

\[
E = E^0 + \frac{RT}{2F} \ln \frac{\Pi(\text{reactant fugacity})}{\Pi(\text{product fugacity})}
\]  
(Eq. 1.37)

In the particular system of Eq. 1.34, \( E \) will be rewritten by taking into account the partial pressures of the gases involved.

\[
E = E^0 + \frac{RT}{2F} \ln \frac{pO_2^{1/2}pH_2}{pH_2O}
\]  
(Eq. 1.38)

For the overall cell reaction, the cell potential increases with an increase in the partial pressure (concentration) of reactants and a decrease in the partial pressure of products. For example, for the hydrogen reaction, operating at higher reactant pressures can increase the ideal cell potential at a given temperature. [41]

When no current is being drawn from the fuel cell, the cell voltage is at a maximum, the open circuit voltage \( E \). As current is extracted from the cell, additional irreversible losses result in a decrease in the cell voltage \( E_{\text{cell}} \), according to

\[
E_{\text{cell}} = E - iR - \eta_c - \eta_A
\]  
(Eq. 1.39)

where \( iR \) involves the ohmic losses within electrodes, interconnects, and current take-offs due to the finite resistance of the materials used, \( \eta_A \) refers to the overpotential at the anode, reflecting losses due to both electrode kinetics and mass transport limitations, and \( \eta_c \) refers to the equivalent overpotential at
the cathode. Thus, materials selection and microstructure are fundamental in minimizing these losses and, therefore, in maximizing fuel cell performance.

High temperature (ca. 1000 °C) SOFC technology (HT-SOFCs) is based on the oxide-conducting yttria-stabilised zirconia (YSZ) electrolyte. This working regime provides highly efficient conversion to power, internal reforming, rapid electrocatalysis with non-precious metals and high quality heat by-product, which can be used for co-generation, or for use in combined cycle applications. SOFC systems have demonstrated one of the highest efficiencies (ranging from 40 to 70 %) of any power generation system, combined with very low environmental impact (virtually no acid gas or solid emissions). [42-44] The modular and solid state construction of SOFC devices allows flexibility to build stationary power generation plants in the 2 kW to hundreds of MW capacity range. [42] SOFCs are able to work not only with H₂ fuel, but also several hydrocarbons as CH₄, alcohols and CO, giving rise to CO₂ and H₂O as residue.

However, SOFCs present various drawbacks, usually derived from the high operation temperature. Thermal expansion mismatches between component materials may lead to failure of solid electrolyte and sealing difficulties between cells in the flat plate configurations. The aggressive operation conditions also cause corrosion of metal components (such as the interconnects) and the modification of the materials, which may develop insulating phases. All these issues need to be taken into account for thermal cycling and stack life. [45] Hence, the material selection is very restricted and results in difficult and high cost fabrication processes.

Intermediate temperatures SOFCs (IT-SOFCs) working at ca. 500-700°C have been developed to face the mentioned disadvantages and to improve the durability. Combined with the ability of SOFC to use fossil fuels, the new operation conditions reduce the cost of the fuel cell since less-expensive materials of construction may be used.
More recently, new proton conducting SOFCs (PC-SOFCs) have been developed, whose electrochemical reactions are based on the transport of protons or protonated species. The main difference regarding an oxygen ion electrolyte is the side in the fuel cell in which the water is produced. In the case of pure proton conductivity, hydrogen is oxidized in the anode to form protons, which are incorporated into the electrolyte. These protons are consumed at the cathode in the reduction of oxygen to produce water, which can easily be removed by the oxygen (air) flow. PC-SOFCs allow lower operating temperatures and achieve very high fuel utilization efficiencies, since water is formed at the air electrode and the fuel is not diluted. [46] However, there is still a need to further develop protonic electrolytes and electrode materials to achieve an optimum combination of proton transport properties, compatibility with other cell components, and thermochemical stability.

**General requirements for SOFC**

Target power densities of fuel cells are typically of the order of 1 kW kg\(^{-1}\), which means that the area specific resistivity (ASR) of the cell (electrolyte plus both electrodes) should not exceed 0.5 Ω cm\(^2\). This involves general requirements for the principal components, as low individual resistivity, chemical compatibility between materials and similar thermal expansion, high temperature stability of all the components and high strength and toughness. Traditionally, the electrolyte is made from a ceramic such as YSZ, the anode is Co-ZrO\(_2\) Ni-ZrO\(_2\) or Ni-YSZ cerments, and the cathode is Sr-doped LaMnO\(_3\) (LSM) or LSM-YSZ cerers. New strategies and materials for each cell component have been studied and developed in an attempt to reduce the ASR of the cell, which are described below.
**Electrolyte**

Solid electrolytes consist of a dense non-porous microstructure that avoids the permeation of neutral gases. The electrolyte material must possess high oxygen ion or proton conductivity (> 0.05 S·cm⁻¹) at the operating temperature while electronic conductivity should be negligible both in oxidizing and reducing atmospheres to prevent losses from leakage currents. The most common oxygen conductor electrolytes are doped zirconia-based materials, especially the mentioned YSZ and scandia-doped zirconia (ScSZ). Other stabilizing dopants have been used, as CaO, MgO and certain rare earth oxides such as Nd₂O₃, Sm₂O₃, Yb₂O₃. Also Sm, Gd, Y or Ca doped ceria (SDC, CGO and YDC, respectively), which are ionic conductors, have been tested as electrolyte materials. The main problem with the cerias is that they react with the electrodes during the sintering at high temperatures and develop electronic conductivity both at temperature above 600 ºC and in the reducing atmospheres of the anode side due to the Ce⁴⁺ reaction. Other works focused on perovskite-structured electrolytes. Ishihara *et al.* developed Sr-Mg doped lanthanum gallate ceramics (LSGM), which demonstrates higher ionic conductivities than ZrO₂ and CeO₂ based oxides at intermediate operating temperatures, while keeping pure ionic behaviour in a wide pO₂ range (1-10⁻²⁰ atm). [47, 48] However, in an overall electrochemical cell LSGM reacts with almost every electrode materials and the performance is quickly degraded due to the phase change and formation of minor phases. [49]

Bismuth oxide is the suggested electrolyte material showing the highest ionic conductivity. However, it is not stable in reducing atmospheres and undergoes a phase transition from the monoclinic to the high temperature cubic phase, at approximately 730 ºC where develops electronic conductivity. Many metal oxides (MO, M₂O₃, M₂O₅) dopants have been tried in order to stabilize bismuth oxide, although lanthanides (especially Er₂O₃) or Y₂O₃ have been
studied in more detail. Y$_2$O$_3$-Bi$_2$O$_3$ structure was stable over a wide range of temperatures and shows an ionic conductivity about 0.2 S cm$^{-1}$ at 700 ºC, but it still develops $n$-type electronic conductivity at low $pO_2$. [50] Er$_2$O$_3$-Bi$_2$O$_3$ was stable to oxygen partial pressures lower than 10$^{-20}$ atm while being one of the highest ion conductors in air within this system. Finally, BIMEVOX and Aurivillius phases are families based on Bi$_4$V$_2$O$_{11}$ in which V is replaced by a metal ion (as Cu or Ni). They are even lower temperature ionic conductors that allow high oxygen diffusion rates, but reveal surprisingly slow surface exchange kinetics. [51]

To overcome the material hitches, electrolyte performance enhancement is addressed on the cell configuration optimization. The construction of thin electrolyte films supported on porous electrode substrates allows decreasing the operation temperature. Also double layered electrolytes have been proposed, where a zirconia-coated side is exposed to the fuel gas to block the electrons, and the material that may be reduced is in contact with the cathode. Some examples are the thin coat of ZrO$_2$/stabilised bismuth oxide or YSZ deposited by sputtering over CGO; However, possible interaction and interdiffusion between phases still appear as a consequence of the electrolyte sintering temperature and influences on the performance of the composite electrolyte. This brings about the concept of protecting layers to avoid the rapid degradation of the fuel cells, as cathode-interlayer-electrolyte or sandwiching YSZ between two CGO layers. [52-55]

Proton conducting solid electrolytes might eventually overcome some of the limitations of oxygen ion conductors cells, since the lower activation energy of proton mobility regarding oxygen vacancies enables operation at lower temperatures. Nevertheless, they are still in an early stage of development. The first compounds with high proton conductivities were based on SrCeO$_3$ and
Introduction

BaCeO$_3$, which reached power densities up to 0.2 W cm$^{-1}$ at 800 °C. However, these materials react to carbonates even with low levels of CO$_2$ to form alkaline earth hydroxides at high water activities, which excludes any hydrocarbon fuel. [56] Among these materials, BaZrO$_3$ is an exception in stability, but requires high sintering temperatures that lead to high grain boundary resistances. A trade-off between stability and lower grain boundary resistance may be obtained by partial substitution of Zr by Ce in the perovskite structure. Others like yttrium-doped barium zirconate (BaZr$_{1-x-y}$Y$_x$M$_y$O$_{3-δ}$) have been investigated, using M=Pr, Fe and Mn as dopants. Among these, the highest total and protonic conductivity at elevated temperatures under reducing atmospheres was found for BaZr$_{0.8}$Y$_{0.15}$Mn$_{0.05}$O$_{3-δ}$. [57] More recently, new classes of proton conducting oxides are derived from acceptor-doped rare earth ortho-nobates, LnNbO$_4$ (Ln$_{1-x}$M$_x$NbO$_{4-x/2}$, where Ln = Y, La, Nd, Gd, Tb, Er; M = Ca, Sr, Ba and x ≤ 0.1). They showed protonic conduction and a general stability against CO$_2$. LaNbO$_4$ based materials exhibit the highest proton conductivity, which is close to 0.001 S cm$^{-1}$ for La$_{0.99}$Ca$_{0.01}$NbO$_4$ under 2.5% H$_2$O at ~950 °C. [58] La$_{6-x}$WO$_y$ with 0.4<x<0.7 (LWO) is a mixed protonic-electronic conductor (mainly protonic below 800 °C) highly stable in CO$_2$ containing atmospheres. The applicability of LWO as a protonic electrolyte has been proved by recording the OCV in a Pt/LWO/Pt cell as a function of the temperature. [59-62] Proton conductivities of 0.005 S cm$^{-1}$ have been achieved for La$_3.8$WO$_{11.7}$ at 800 °C in wet air. [63] Latest works have shown that compounds of Ln$_6$WO$_{12}$ (Ln = La, Nd, Gd, Er, Eu) series exhibit mixed proton-electron conductivity at high temperature in humid reducing or oxidizing atmospheres. [61, 62]

Cathodes

The electrochemical reduction of oxygen takes place in the cathode side. The reaction described in Eq. 1.40 occurs when the oxygen ion conductor, the
electronic conductor and the gas phase come in contact, sites that are known as triple-phase boundaries (TPBs).

\[ \frac{1}{2} O_2 \text{(gas)} + 2e^- \text{(cathode)} \rightarrow O^{2-} \text{(electrolyte)} \]  
(Eq. 1.40)

Therefore, high electronic conductivity (> 100 S cm\(^{-1}\)) for the electron to reach the oxygen and high catalytic activity for the oxygen reduction reaction are required. Ionic conductivity is also desired in order to increase the TPBs area all along the cathode. Besides, the cathode material should not react neither with the electrolyte nor the interconnectors (or the protecting layers used to protect them, typically spinel-structured materials) to avoid changes of phase or blocking impurities. This compatibility must be extended to the thermal expansion coefficient (TEC), since great differences with the materials in contact may lead to cracks and failures in the components. The porosity of the electrode must be adequate to enhance the diffusion of oxygen gas to the cathode/electrolyte interface. [64, 65]

Typical HT-SOFC cathode material is a composite made of LSM mixed with YSZ, to provide electronic and ionic conductivity, respectively, and to improve TEC compatibility with the YSZ electrolyte. In this situation, the electrochemically active reaction sites (TPB) increase orders of magnitude over that of porous cathodes exhibiting only electronic conductivity.

However, the new trends of reducing the operating temperature also involve the degradation of the electrode kinetics and results in large interfacial polarization resistances. At the reduced temperatures of IT-SOFC, the standard cathode material LSM cannot be used because of its low activity for the oxygen reduction reaction (due to its low oxygen diffusivity). The approach in this case is to use single-phase MIEC cathodes permitting both oxide ion and electron mobility within the cathode material, e.g., La\(_{1-x}\)Sr\(_x\)CoO\(_{3-\delta}\) (LSC),
La$_{x}$Sr$_{1-x}$Co$_{1-y}$Fe$_{y}$O$_{3-\delta}$ (LSCF), Pr$_{1-x}$Sr$_{x}$FeO$_{3-\delta}$ (PSF), Pr$_{1-x}$Sr$_{x}$Co$_{1-y}$Fe$_{y}$O$_{3-\delta}$ (PSCF), Pr$_{1-x}$Sr$_{x}$Co$_{1-y}$Mn$_{y}$O$_{3-\delta}$ (PSCM) or Ba$_{0.5}$Sr$_{0.5}$Co$_{0.8}$Fe$_{0.2}$O$_{3-\delta}$ (BSCF), but they still lack in ionic conductivity or phase stability. [66-68] Cobalt free cathodes, both single phase or forming a composite with a ceria based fluorite have been also tested with the aim of improving TEC compatibility, but a drop in electrocatalytic properties is experienced. [69-71]

Recently, Co containing materials based on YBaCo$_4$O$_7$ swedenborgites, with CGO and YSZ compatible TEC have been developed, which demonstrated low polarization resistances compared to LSCF reference material. [72,73] Layered perovskite oxides LnBaCo$_2$O$_5$ (LBCO) (Ln=Lanthanide) with A-site Ln$^{3+}$ and Ba$^{2+}$ cation-ordered structures are new potential cathode materials. Particularly, the Pr based family performs the best electrochemical results due to the fast oxygen transport kinetics. [74] On the other hand, single phase fluorites, as Ce$_{1-x}$Pr$_x$O$_{2.6}$ (CPO) have been proposed as cathode since they present high levels of oxygen non-stoichiometry and significant electronic contribution from electron hopping between adjacent Pr ions, even at high $pO_2$. [75] By using these strategies, the cathode reaction occurs over the full electrode surface area, rather than being limited to the TPB. Thus, the overall cathode performance depends essentially on the rate of defect transport through the solid MIECs, gas transport through the pores to promote surface coverage in the MIECs, and the catalytic activity of interfaces. [76]

On the other hand, the use of PC-SOFC requires the development of tailor-made suitable electrodes for these particular electrolytes, since the main issue is the chemical compatibility between the cathode and the electrolyte materials. For barium-zirconate based electrolytes cobalt-free composites as Sm$_{0.5}$Sr$_{0.5}$Fe$_{0.8}$Cu$_{0.2}$O$_{3-\delta}$-Ce$_{0.8}$Sm$_{0.2}$O$_{2-\delta}$ (SSFCu–SDC) or Sm$_{0.5}$Sr$_{0.5}$FeO$_{3-\delta}$-BaZr$_{0.1}$Ce$_{0.7}$Y$_{0.2}$O$_{3-\delta}$ (SSF-BZCY) have been reported as suitable composite cathodes. [77, 78] La$_0.99$Ca$_{0.005}$NbO$_4$ (LCNO) proton conductor electrolyte is
chemically and mechanically compatible with typical cathode material La$_{0.75}$Sr$_{0.25}$Cr$_{0.5}$Mn$_{0.5}$O$_3$ (LSCM), while reacts with BSCF cathode and with LSM (La$_{0.7}$Sr$_{0.3}$MnO$_3$) causing La and Nb electrolyte cations to diffuse in the LSM phase. [38,79] The incorporation of the proton conducting material in a composite cathode as LCNO-LSM cer-cer increase in the TPB length providing reaction sites along the whole thickness of the electrode. [59] On the same line, specific cathodes for LWO-based cells have been developed performing a composite of LSM (La$_{0.8}$Sr$_{0.2}$MnO$_3$) and LWO. [80] Analogously to oxygen ion SOFC, the integration of the electrolyte material in the cathode structure has some advantages as the improvement of chemical and mechanical stability and decrease in processing cost, so that many researches are addressed in this trend. [81-83]

Anodes

The anode side hosts the fuel oxidation reaction written in Eq. 1.41. Similarly to cathode side, the reactions take place as long as an oxygen ion conductor, electronic conductor, and the gas phase converge.

$$H_2 (fuel) + O^{2-} (electrolyte) \rightarrow H_2O + 2e^- (anode) \quad \text{(Eq. 1.41)}$$

Thus, the anode requires high ionic conductivity to ensure electrical contact with the electrolyte, and high electronic conductivity that provides good electrical contact with the interconnector. As for cathodes, the use of a MIEC material instead of a pure electronic conductor allows extending the active sites from the TPBs to the whole anode surface area. An anode material with high catalytic activity is necessary to promote chemisorption and dissociation of fuel at the surface and the product reaction with the oxygen ions provided by the electrolyte. Similar TEC for the anode and electrolyte prevents from cracks and problems during sintering and sealing. Also chemical stability with the materials
in contact is needed to avoid ion or electron blocking phases. Finally, the porosity must be adequate and physically stable to allow gas transport to and from the electrolyte.

Two-phase ceramic-metal composites (cermets) may combine conduction and catalytic properties. Actually, the most commonly used anode materials are Ni/YSZ cermets, which display excellent catalytic properties for fuel oxidation and good current collection, but do exhibit disadvantages, such as low tolerance to sulphur and carbon deposition when using hydrocarbon fuels, or the fatal mechanical failure upon re-oxidation or highly moistured environments. Nickel-free SOFC anodes try to overwhelm these problems. For example, ceria based materials, which become electronic conductors at the anode reducing atmosphere have been used as anodes, being limited by their low total conductivity. Similar problem is observed for titanate based oxides, combined with fast anode degradation. [84] Perovskite structured chromites are stable in reducing conditions. LaCrO$_3$ has been used as interconnect materials for SOFC, but their stability in both reducing and oxidizing atmospheres at high temperatures make them anode suitable materials. Substituting alkali earth and transition metal elements on the La and Cr sites of LaCrO$_3$ may modify their electronic and catalytic properties. Typically, Sr is introduced in La site, while other transition elements into the Cr-site of La$_{1-x}$Sr$_x$Cr$_{1-y}$M$_y$O$_3$ (M = Mg, Mn, Fe, Co, Ni) may improve the catalytic properties for methane fed SOFC. [85-87] The use of chromite based perovskites has been also proposed as anode material for LWO electrolyte based PC-SOFC, since the typical anode materials for the barium zirconates proton conductors, NiO or nickel-based cermet combinations (NiO or Ni /Ba(Ce$_{0.8-x}$Zr$_x$)$_{0.1}$O$_{3-d}$), react with the LWO electrolyte. [88, 89] La$_{0.85}$Sr$_{0.15}$CrO$_{3-δ}$ (LSC) based anodes have been tested as compatible and promising anode materials for LWO electrolytes, although surface reaction processes limit their operation.
Interconnectors

The interconnectors must conduct the electron flow from the anode to the cathode by an external path. Thus, the materials require high electronic conductivity with negligible ionic conductivity. Metals are commonly used, which selection is determined by the operation temperature and TEC of the other components.

The work developed in this thesis aims to find new solid oxide materials that mean an improvement as any ionic, electronic or MIEC material. Regarding to the transport behaviour ascertained and stability, they will be classified and tested for their most suitable applications.

1.4. References


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Introduction


Chapter 2. Methodology

Abstract

This chapter describes the procedures developed for the material synthesis and specimen fabrication. The techniques employed to characterize the samples and determine their transport properties are also explained. The different assemblies used for the electrochemical characterization are outlined as well.
2.1. Material synthesis and sample preparation

Material synthesis

Different methods have been applied to prepare the materials depending on the desirable characteristics for the powders and on the metal precursor nature. Ceria doped compounds have been fabricated by co-precipitation method. CaTiO$_3$ perovskites have been formed and doped by solid state reaction, commonly known as traditional ceramic method. Doped (LaSr)CrO$_3$ perovskites have been synthesized by Pechini sol-gel route, [1] and the same method has been employed for dual-phase nanocomposites. The details of the fabrication are described below.

Co-precipitation

The co-precipitation method allows the synthesis of nanometric size powders. In this work this method has been employed on the fabrication of lanthanide doped ceria materials, as it is shown in the scheme of Figure 2.1. A clear solution of commercial lanthanide nitrates mixture is prepared in deionized water at 50 °C. A (NH$_4$)$_2$CO$_3$ solution is dropped into the previous solution to achieve total precipitation and the final NO$_3^−$/CO$_3^{2−}$ molar ratio is 0.75. The resulting precursor powders are dried at 100-150 °C after filtration and rinsing with water. In some cases, the cerias have been codoped with cobalt. The cobalt addition is done over the dried precursor powder by incipient wetness impregnation, i.e., calculated 2% molar of Co(NO$_3$)$_2$·6H$_2$O is dissolved in deionized water (volume according with the pore volume) and mixed with
the intermediate dried powder. Finally, each powder is calcined during 5 hours in air atmosphere at 800 ºC to decompose the residual nitrates and carbonates and to favour the formation of the fluorite phase.

![Diagram of the fabrication process](image)

Figure 2.1. Scheme of the fabrication of the lanthanide doped ceria. The addition of Co by impregnation is indicated. The calcination temperature ensures the decomposition of the nitrates and carbonates from the precursors.

**Solid state reaction**

The solid state reaction is the most common synthesis process to produce conventional ceramic powders. The method consists of mixing the precursor oxides, carbonates, hydroxides or salts dispersed in acetone or other solvent in a ball mill, which reduces and homogenizes the particle size. The ball mill uses 3 mm diameter zirconia balls and turn at 50 rpm. After the milling, the powder is separated from the zirconia balls and dried. Then, the powder is fired at temperatures of at least two thirds of the melting point for periods up to 10 h. [2] This method presents several drawbacks. One the one hand, it delivers low homogeneity and purity powders with broad particle size distribution. [3] On the other hand, the high-energy consumption to reach the required high
temperatures and the issues concerning to the toxicity and evaporation of some elements have to be taken into account.

*Pechini or citrate complexation route.*

The Pechini route presents some advantages such as obtaining carbonate-free, chemically homogeneous final oxide compounds with a high relative density and small size. It is a sol–gel technique that involves the production of an amorphous-like gel containing a metal complex. Citric acid, ethylenediaminetetraacetic acid (EDTA) or glycine can be used as chelating agent that prevents partial segregation of metal components, which could occur in the case of different stabilities with the metal ions in solution. [4] This complexation of metal ions in citric acid is followed by dehydration at low temperature. In some cases, as in this work, a polyol (*e.g.* ethylene glycol) is added to the reaction batch to promote the polymerization with chelating agent and produce an organometallic polymer. Finally, thermal decomposition of the complex to form the structural phase, *e.g.* fluorite, perovskite or spinel, is carried out at 600 °C or 750 °C.

**Sample preparation**

*Rectangular probes*

This sample configuration is used in electrochemical experiments as DC-conductivity. The as prepared powder is mixed and dispersed in acetone and ball milled with zirconia balls during 12-15 h. After rinsing and drying, rectangular probes (4 x 0.4 x 0.2 cm³) of the initially fired nanopowder were uniaxially pressed at 125 MPa during 1 minute and subsequently sintered 5 hours at high temperature (1000-1450 °C) in air atmosphere. Final dimensions
will depend on the shrinkage undergone during the sintering process, as observed in the examples on Figure 2.2.

Figure 2.2. Main steps for rectangular probes fabrication. From left to right: initial powder, ball milling step, rinsing of the powder, pressing, high temperature calcination and final configuration of sintered samples.

**Membranes and electrolytes**

Membranes and electrolytes are obtained from the sintered powders by the same process, which is schematized in Figure 2.3. In this work, membranes consist of flat dense discs. The homogeneity is very important for the sintering process, which is achieved by ball milling the powder following the same process explained for solid state reaction precursors. The typical particle size of the milled powder is selected and homogenized by separating them through steel sieves of a particular mesh size (200-400 µm). A 26 mm diameter steel dye is used to uniaxially press the material at 125 MPa during 3 minutes. Afterwards, the green disk is exposed to a high temperature sintering process in order to densify the specimen. The temperature and time of the process depend on the material. Permeation measurements are performed on 15 mm diameter and 500-900 µm thick disks, which are ground by sandpaper to reach that dimensions (Figure 2.3). In some cases both disk sides were screen printed by 20-30 µm of a catalytic layer (metals, MIECs or composite materials) in order to improve the surface catalytic behaviour.
Methodology

Figure 2.3. Main steps for membrane fabrication. From left to right: initial powder, ball milling step, rinsing and drying of the powder, uniaxial pressing, high temperature calcination and final configuration of sintered samples. Afterwards, a catalytic layer may be screen printed on the membrane surface.

Electrodes

Thick film SOFC anode and cathode electrodes have been attached to the electrolyte by screen-printing technique. This is the most economical, simple and widely used technique to produce films with a thickness in the range of 10–100 µm. [5] It consists of applying highly viscous ink made of the ceramic powder, an organic binder (ethyl cellulose) and a plasticizer (terpineol). A three-roll mill refines the powder and the other ingredients to form an uniform slurry. This slurry is forced to pass through a 9 mm diameter mesh to develop the electrode. The layer deposition is then dried and sintered at high temperature. After the drying step, more than one layer may be deposited by this method, depending on the targeted thickness. The thickness for the porous layers is ~15 µm and the area of the electrode is ~0.6 cm², as depicted in Figure
2.4. In addition a gold mesh is screen printed on the electrode surface to contact the current collector.

Figure 2.4. Main steps for electrode fabrication. From left to right: initial powder, ball milling step, rinsing and drying of the powder, ceramic ink fabrication, screen printing of the electrode on the electrolyte surface and final electrode configuration after high temperature calcination. In addition, the electrode on the right is screen printed with a gold mesh for current collection.

In this thesis, the electrodes have been tested as symmetrical cells. They consist of two porous electrodes of the same material attached to both sides of the dense electrolyte. The same atmosphere is present in both sides of the symmetrical cell for the analysis.

2.2. Structural characterization

Scanning Electron Microscopy

Scanning electron microscopy (SEM) allows for high-resolution imaging of surfaces by using high-energy electrons (1.5-20 KeV) generated by
a heated tungsten filament. The SEM has a magnification up to $10^6 \times$ and great depth of field. For that purpose, an incident beam of monochromatic electrons causes a secondary emission across the sample surface, which is collected to form an image of the surface. The quality and contrast obtained in the image depends mainly on the conductivity and surface topography of the sample. By detecting different emissions of the sample, it is possible to obtain diverse contrast images, *i.e.*, backscattered electrons (BSE), Auger electrons (AES) and Energy dispersive x-ray spectroscopy (EDS). BSE causes different contrast subjected to the atomic number, $Z$, of the elements, so changes in composition can be distinguished; EDS is a qualitative and quantitative chemical microanalysis technique to characterize the elemental composition of the volume analysed. It is performed in conjunction with a SEM, but uses the X-rays that are emitted from the sample due to the electron beam. The combination of SEM with EDS allows the analysis of the sample morphology and the composition of the different phases that are present. The minimum detection limits are about 0.1 weight percent, depending on the element and matrix.

Conducting materials that allow the transport of the incident beam electrons do not need any handling of the sample. However, for poor conductors or insulators a conducting layer that does not modify the topography needs to be added. This is achieved by coating the sample in vacuum with Au or graphite (for EDS) using a sputter coater.

Two SEM devices were used on the characterizations, namely the Jeol JSM 6300 and Jeol JSM 5410, both with an acceleration voltage of 20 kV. The EDS microanalyses were carried out with a detector from Oxford Instruments. The Software INCAEnergy/Wave served for the interpretation of the X-ray patterns obtained. More recently, a ZEISS Ultra55 field emission SEM (FE-SEM) has been also used.
Transmission Electron Microscopy

Transmission Electron Microscopy (TEM) utilizes energetic electrons to provide morphologic, compositional and crystallographic information on samples. The beam of electrons is transmitted through an ultra-thin specimen (<200 nm), interacting with it as the beam passes through. The electrons that are not absorbed by the sample are detected forming a two dimensions image, which is magnified and focused onto an imaging device.

The TEM device employed for the experiments is a Philips CM10, which has allowed obtaining particle sizes and morphologies of the nanoparticles. In addition, a transmission electron microscope FEI Tecnai F20 (with acceleration voltage of 200 kV) has been used. TEM lamellas were prepared by means of a FIB (Focused-Ion Beam, FEI Helios Nanolab 400s) with the subsequent thinning performed in a BAL-TEC RES-120.

X-ray diffraction

X-ray diffraction (XRD) is a non-destructive technique that allows the identification of the crystalline phases and orientation of crystalline materials, their lattice parameters (10-4 Å), strain, crystallite size and thermal expansion, which define the fingerprint of the substance. In a crystalline material, the atoms are arranged in a regular pattern. The unit cell is the smallest volume element that by repetition in three dimensions describes the crystal. Three axes describe the dimensions of the unit cell: a, b, c and so the angles between them α, β, γ. Different combinations of these parameters release fourteen kinds of cells covering all possible point lattices or all crystals, which were classified by August Bravais. Since some of the Bravais lattices can be expressed by other simpler lattices, they can be represented by seven primitive lattices, summarized in Table 2.1. [6]
Table 2.1. Summary of the seven crystal systems and Bravais lattices. [6] The axial length and angles, and lattice symbols are also indicated.

XRD measurements are based on the physical principle that a monochromatic X-ray beam with a wavelength $\lambda$ comes into contact with the crystalline material at a particular angle. The diffraction is produced only when the distance traveled by the rays reflected from successive planes differs by a complete number of wavelengths. As shown in Figure 2.5, the diffracted angle from the family of crystallographic planes $(h k l)$ has a value of $2\theta$, which is employed to determine the distance between layers of atoms in a sample using the Bragg's law:

$$n\lambda = 2d_{hkl} \sin \theta$$  \hspace{1cm} (Eq. 2.1)
where \( d \) is the \( h k l \) interplanar spacing (nm), \( \theta \) is the angle of incidence of X-rays (rad), \( \lambda \) is the wavelength of X-rays (m); and \( n \) is an integer representing the reflection order.

\[
\frac{\sin^2 \theta}{h^2+k^2+l^2} = \frac{\lambda^2}{4a^2} \quad \text{(Eq. 2.2)}
\]

where \( a \) is the cell parameter and \( h, k, \) and \( l \), the Miller indices defining the plane. The plane spacing for solving other systems is provided in table of the Appendix 2.

On the other hand, XRD peak profile is also employed to estimate the particle size of the crystallites in a powder sample. It is worthy here to emphasize the difference between the grain size of a crystalline powder sample and the size of crystallites. The term ‘particle size’ or ‘grain size’ is used when the size of individual crystals is less than about 100 nm, because a particle of a real crystalline powder sample generally consists of many fine units called
Methodology

‘crystallites’, which can be considered as single crystals, as drawn in Figure 2.6. Although the size of crystallites may be the same as that of grain size in some cases, they are essentially different physical quantities. The size of crystallites is related with the peak width, broader as smaller the crystallite. Crystallites in a crystalline powder smaller than 5 nm will show a diffraction peak profile clearly different from that of the same sample with the standard size of 0.5-10 µm in diameter, which will present sharper peaks. [6]

![Figure 2.6. Schematic diagram for grain size and crystallite of a crystalline powder sample. A particle or grain of a real crystalline powder generally consists of many crystallites, which can be considered as single crystals.](image)

Calculations of crystallite size commonly employ the Scherrer’s equation (Eq. 2.3) for estimating the size of crystallites from a measured diffraction peak profile.

\[
t = \frac{0.9 \cdot \lambda}{\beta \cdot \cos \theta}
\]  
(Eq. 2.3)

where \( t \) is diameter of crystallites and \( \beta \) represents the full width at half the maximum peak intensity. As order of magnitude crystallites down to about 50 nm, may show peaks of ca. 0.2 degrees broad. [6]

The measurements developed during the present thesis were carried out by a Cubix fast diffractometer, using CuK\(\alpha_{1,2}\) radiation (\(\lambda_{1,2}=1.5406 \ \text{Å} \) and an
X’Celerator detector in Bragg-Brentano geometry. XRD patterns were recorded in the $2\theta$ range from 2º to 90º and analyzed using X’Pert Highscore Plus software. Some studies required obtaining the XRD patterns at high temperature (HT-XRD), which were performed by an Anton Paar XRK-900 reaction chamber attached to the diffractometer, either in dry air atmosphere or in N$_2$ up to 900 ºC.

Density measurements

Density features of the materials were calculated from mass and volume values, obtained by a helium pycnometer, and comparing them with the theoretical values. This method measures the volume of gas (He) displaced by a known mass of powder, and gives the real density of the material by (Eq. 2.4). The sample must be completely dried. The density measurements were carried out at least three times to average out the variation in values measured.

$$\rho_e = \frac{\text{Sample mass}}{\text{Displaced volume}}$$

(Eq. 2.4)

The theoretical density ($\rho_T$) is determined from the molecular weight and the cell parameters obtained by XRD. As an example, for a cubic fluorite structure, which has 4 lanthanide and 8 oxygen atoms, $\rho_T$ is calculated as follows:

$$\rho_T = \frac{\sum n_x n [4 \cdot AW_n + 8 \cdot AW_O]}{[CV] [N_A]}$$

(Eq. 2.5)

Where $n$ represents the number of different elements forming the material; $x$ is the molar fraction of the element in the material; $AW$ is the atomic weight of the n-lanthanide and the oxygen; $CV$ is the volume of the unit cell and $N_A$ is the Avogadro’s number.


\[ E_K = h\nu + E_b + E_w \]  (Eq. 2.6)

Being \( E_k \) the kinetic energy of the electrons, \( h\nu \) the energy of the photons, \( E_b \) is the binding energy characteristic of each element, and \( E_w \) is the work function of the spectrometer. The measurement yields a spectrum with a series of photoelectron peaks whose peak areas determine the composition of the materials surface. The shape of each peak and the binding energy can be slightly altered by the chemical state of the emitting atom. Thus, these abnormalities can be used to extract chemical bonding information. As a limitation, XPS is not sensitive to hydrogen or helium, and must be carried out in ultra high vacuum conditions. [8]

Sintered pellets of the materials under study were quenched from high temperature in the desired atmosphere to air saturated liquid N\(_2\) to imitate the surface in working experimental conditions. Measurements were carried out on a SPECS spectrometer with a MCD-9 detector and using a non-monochromatic Al K X-ray source. Spectra were recorded using analyzer pass energy of 30 V, an X-ray power of 100 W and under an operating pressure of \( 10^{-9} \) mbar. CASA software has been used to perform the spectra treatment. Binding energies (BE)
were referenced to the C 1s peak at 284.5 eV. The energy regions of O 1s and C 1s transitions and the specific ones of the materials were recorded.

Raman spectroscopy

Raman spectroscopy is a useful tool for obtaining information about the structure and properties of molecules from their vibrational transitions. Raman vibrational bands are characterized by their frequency (energy), intensity, and band shape (environment of bonds). The frequencies of these molecular vibrations depend on the masses of the atoms, their geometric arrangement, and the strength of their chemical bonds. [9, 10] Thus, it is possible to detect structural and compositional changes by the anomalies in position, intensity and shape of the bands.

Raman spectra were measured with a Renishaw inVia Raman spectrometer equipped with a Leica DMLM microscope and a 514-nm Ar+ ion laser as an excitation source. A ×50 objective of 8-mm optical length was used to focus the depolarized laser beam on a spot of about 3 µm in diameter. The Raman scattering was collected with a charged coupled device (CCD) array detector.

UV-vis spectrophotometry

The electronic properties of a semiconductor may be determined from the analysis of its energy band diagram. The separation between the energy of the lowest conduction band and that of the highest valence band is called the band-gap ($E_g$). UV–vis diffuse reflectance spectroscopy is a commonly employed optical method to evaluate the $E_g$. It describes the electronic behavior present in the structure of the solid giving information about the electronic transitions of the different orbitals of a solid.
The UV–vis technique measures the increase in the absorbance at a given wavelength (band gap energy) due to the optical excitation of the electrons from the valence band to the conduction band. Graphical Kubelka-Munk \((K-M)\) transformation (Eq. 2.7) is commonly employed to obtain the optical \(E_g\).

\[
F(R) = \frac{(1-R)}{2R}^2
\]  
(Eq. 2.7)

where \(R\) is the reflectance; \(F(R)\) is proportional to the extinction coefficient \(\alpha\). Any electronic transition has been considered here, so the plot directly represents \(F(R)\) versus the energy in eV. The optical gap has been inferred by linear extrapolation of the absorbance from the high slope region obtained from the spectra. [11]

By doping the materials under study, it is possible to introduce electronic levels that will make the band gap narrower. These changes are followed by UV-vis spectra of the compounds recorded on a Varian 5000 UV-Vi-NIR spectrophotometer. The wavelength range is 200–800 nm (with a lamp change at 350 nm) using BaSO\(_4\) as reference material.

### 2.3. Thermal Analysis

Thermal analysis techniques detect the transformations occurring in a sample under a temperature programme in a controlled atmosphere.

**Thermogravimetry/Differential Thermal Analysis (TG/DTA)**

Chemical and thermal stability of materials can be studied by TG technique, which is commonly used to investigate decomposition, dehydration,
oxidation/reduction reactions, reaction pathways, kinetics of reaction and the quantity of a specific component that occurs accompanied by a mass change. The TG device registers the mass changes undergone by the sample as a function of temperature, time and atmosphere. The TG is combined with DTA, which records exothermic or endothermic processes that enclose transformations in the material when compared with an inert reference material. DTA is mainly used for studying transitions, chemical reactions, adsorption, crystallisation, melting and sublimation. In addition it is possible to measure the heat of reaction and to determine kinetic parameters. [12]

In the current study, the thermobalance was used to monitor changes in the oxygen non-stoichiometry, to ensure the complete decomposition of precursors (carbonates, nitrates and polymers) and to test mass changes of the material in the reaction against CO₂. The TG/DTA analysis was performed on Mettler-Toledo StarE equipment, which has different gases available, such as synthetic Air (21 vol% O₂ and 79 vol% N₂), N₂ and 5% CO₂/air. The powders were analysed in these atmospheres in the temperature range from room temperature to 1000 °C, with heating and cooling rates of 10 °C/min.

Temperature programmed desorption

Temperature programmed desorption (TPD) techniques are important for the determination of the kinetic and thermodynamic parameters of desorption processes or decomposition reactions. The method consists of heating a sample and simultaneously detecting certain absorbed species that have enough energy to escape, by means of a mass analyser. The mass spectrometer measures the mass of gas phase ions produced from the ionisation of the molecules and atoms of a specimen and allows qualitative and quantitative determination of its elemental composition. These ions are separated according to their mass to charge ratio (m/z) and then detected. [13]
TPD measurements were performed in order to observe the oxygen release with temperature. Around 100 mg of powdery material sintered at high temperature are placed in a quartz reactor and heated up and cooled down in air up to it 1000 °C. Then the sample is heated up at 10/min up to 1000 °C in the desired atmosphere, *i.e.* He, N₂, Ar or CO₂. The ions and molecules delivered are monitored and recorded with a mass spectrometer Omnistar (Balzers), by following the corresponding ionic masses: *m/e* = 32 and 16 amu for oxygen, or *m/e* = 44 amu for CO₂.

**Temperature programmed reduction**

Temperature programmed reduction (TPR) method is a widely used tool for the characterization of metal oxides and mixed metal oxides. Quantitative information of the reducibility of the oxide's surface, as well as the heterogeneity of the reducible surface can be obtained by TPR. A reducing gas mixture (typically 3% to 17% hydrogen diluted in argon or nitrogen) flows over the sample, while a thermal conductivity detector (TCD) measures the changes in the thermal conductivity of the gas stream. The TCD signal is then converted to concentration of active gas using a level calibration. Integrating the area under the concentration vs. time (or temperature) yields the total gas consumed.

In the present work, a 2910 Micromeritics system was used to carry out the measurements. Thus, 100 mg of sample was degassed under Ar flow for 1 h and then was subjected to reduction under H₂/Ar (1/9) flow and heating rate of 10 °C/min till 1000 °C. The H₂ consumption was measured by a TCD.
2.4. Electrochemical characterization

**DC total electrical conductivity**

Electrical conductivity measurements are conducted by standard four-point DC technique. Compared to two-point measurements, this configuration eliminates the influence of the contact resistance and wiring.

![Diagram](image)

Figure 2.7. Four point configuration for DC conductivity measurements. The contacts are made with silver paste and wires.

Samples consisting of sintered rectangular bars are contacted using silver wire and paste, as shown in Figure 2.7. The measurements are carried out in a temperature range from 400 to 800 °C by cooling down (1 °C/min) in constant different O\(_2\) contained atmospheres (Linde calibrated gas mixtures checked by an YSZ oxygen sensor). Exceptionally, the temperature range may be extended to 400-1000 °C. A new combinatorial setup for measuring simultaneously the DC conductivity of ten samples has been developed (Figure 2.8a and b). It consists of a constant \(pO_2\) chamber that is inserted in a tubular oven. The samples and the YSZ sensor are radially attached around the gas inlet. A programmable current source (Keithley 2601) supplies the constant current and a sixteen channel multimeter (Keithley 3706) detects the voltage drop (Figure 2.8c). The I-V slope provides the resistance imposed by the
sample, from which the electrical conductivity ($\sigma_{\text{total}}$) is extracted by applying the Ohm’s law:

$$\sigma_{\text{total}} = \frac{1}{a\cdot b} \cdot \frac{1}{R} \quad \text{(Eq. 2.8)}$$

where $l$ is the distance between contacts, $a$ is the width and $b$ is the thickness of the sample and $R$, the resistance of the material. The conductivity measurements are thermally activated and were analysed on the basis of Arrhenius behaviour

$$\sigma(T) = \frac{A}{T} \exp\left(-\frac{E_a}{kT}\right) \quad \text{(Eq. 2.9)}$$

The activation energy ($E_a$) is extracted from the slope of the graphs. Once the highest temperature (800 °C) was reached, the samples were stabilized for two hours in order to warrantee the high-temperature reduction state corresponding to the specific $pO_2$. 
Figure 2.8. a) Combinatorial DC conductivity setup, b) detail of the sample chamber containing the ten samples and c) from top to bottom, furnace controller, programmable current source (Keithley 2601) that and sixteen channel multimeter (Keithley 3706).

Pulse Isotopic Exchange (PIE)

The fast surface exchange of neutral oxygen and reduction/dissociation to $O^{2-}$ is a basic requirement for conduction in dense materials. It is characterized by the surface exchange coefficient $k$. Pulse isotopic exchange (PIE) [16] technique allows working in a surface exchange dominated regime since it uses powders of a size smaller than the characteristic thickness ($L_c=D/k$). Therefore, it provides a more reliable method to measure $k$ than other techniques as the electrical conductivity relaxation (ECR). [17]

For PIE experiments, the powder was uniaxially pressed at 400 MPa. The disk was sintered at 1450 ºC for 10 h in air, and crushed into a coarse powder. The fraction passing through 125 µm stainless steel sieve was pre-
annealed at 1200 °C for 10 h in air, and sieved again to remove agglomerates. For the measurement, approximately 80-100 mg of sample powder were loaded between two quartz wool plugs in a quartz tubular microreactor with inner diameter of 2 mm. The length of the packed-bed was about 10 mm. A 16O2/Ar gas mixture was used as carrier gas with a flow rate 50 ml/min (STP). The response to 18O2/N2 pulse (500 µl), with the same pO2 than the carrier gas was analyzed by online mass spectrometry at the exit of the reactor. A six-port valve with sample loop was used for injection of the 16O2/N2 pulse into the 16O2/Ar carrier gas flow stream. Oxygen isotope gas was greater than 97 atom% 18O2 (Cambridge Isotope Laboratory). Nitrogen used as diluent for the 18O2 gas was also used for internal calibration of the mass spectrometer. Prior to measurements, the powder packed-bed was pre-treated at 800 °C for 0.5 h under flowing synthetic air in order to remove possibly adsorbed water and CO2, and subsequently cooled to room temperature at a rate of 5 °C/min. The pulse-response isotope exchange measurements were performed in the temperature and pO2 ranges 450–900 °C and 0.21 atm, respectively. The reactor was equilibrated 0.5 h at fixed temperature and pO2 before data collection. The packed-bed microreactor was designed to approximate ideal plug flow behaviour. [18] The isotopic exchange measurements details are explained by Yoo et al. [19]

**Permeation tests**

Permeation measurements were performed on a cylindrical two chambers quartz reactor schematized in Figure 2.9. The inner chamber is 16 mm diameter with a flat holed bottom that supports the membrane. Sealing was done using gold rings heated at temperature >1000 °C and pressed by a spring sited in the lid. Permeation measurements were performed on the 15 mm diameter disks mentioned in section 2.1.2. Both disk sides may be coated by micrometric layer of catalyst e.g. Pt ink (Mateck, Germany).
Oxygen was separated from feed synthetic air using argon as sweep gas. The flow rate of all gas feed streams was controlled using mass flow controllers (MFCs). The standard flow rates used were 100 mL min\(^{-1}\) for feed and 150 mL min\(^{-1}\) for sweep, although they are modified in the different experiments in order to determine their influences on the oxygen separation. In fact, the oxygen permeation has been followed under different parameters, i.e., temperature (500-1000 °C), sweep gas flow or fuel gas.

From the oxygen content measured in the argon side (permeate side) and the argon flow rate, the total oxygen permeation rate was calculated, assuming the ideal gas law. The oxygen content in the permeate side was analyzed using micro-GC Varian CP-4900 equipped with Molsieve5A, PoraPlot-Q glass capillary and CP-Sil modules. The Molsieve5A detects O\(_2\), N\(_2\),
He and H₂. Qualitative analysis of water concentration in the permeate is done in the PoraPlot-Q channel and the CP-Sil measures heavier compounds. Appropriate sealing was confirmed by measuring a tracer concentration in the permeate stream. An acceptable sealing was achieved when the N₂ concentration was lower than 5% of the permeated O₂. Data reported are achieved at steady state after 30 min of stabilization and each test was repeated three times to minimize analysis relative error, which was found to have a standard deviation of $10^{-4}$.

Oxygen concentration is calculated from the O₂ signal area and its response factor, previously calibrated. Total permeation is calculated as the product of O₂ concentration and Ar flux. Permeation flux ($\text{mL} \cdot \text{min}^{-1} \cdot \text{cm}^{-2}$) is extracted from the oxygen flux and the effective area ($A_{\text{effective}}$) of the membrane by the following equation:

$$J(O_2) = \frac{\%O_2_{\text{permeate}}}{100 \%O_2_{\text{permeate}}} \cdot \frac{Q_{\text{Ar}}}{A_{\text{effective}}} \text{(mL} \cdot \text{min}^{-1} \cdot \text{cm}^{-2})} \quad \text{(Eq. 2.10)}$$

**Electrochemical impedance spectroscopy (EIS) and Alternating Current (AC) electrode testing**

EIS is based on the response of an electrochemical cell material to the AC passing through the cell. The different processes occurring inside the fuel cell, *i.e.*, gas transfer to the reaction sites, the electrochemical reaction at those sites, the transfer of ions and electrons in the bulk and in the grain boundary and their combination at the cathode, have different rates. They can be disclosed by measuring in a range of frequencies, whose impedance response is in the complex plane. Thus, it covers circuit elements that exhibit much more complex behavior than that observed by simplified DC conductivity, which is considered
as an ideal resistor. [20] Experimentally, it is carried out by passing a small sinusoidal current (or voltage) through the interface and measuring the phase shift and amplitude, or real and imaginary parts, of the resulting voltage/current signal:

\[ v(t) = V_m \sin \omega t \]  
\[ i(t) = I_m \sin(\omega t + \theta) \]  
\[ Z(\omega) = \frac{v(t)}{i(t)} = \text{Re}(Z) + i\text{Im}(Z) \]

where \( Z \) is the system impedance, \( \omega \) the frequency and \( \theta \) the phase shift between voltage and current intensity.

![Equation 2.11](image1.png)

\[ \text{Eq. 2.11} \]

\[ \text{Eq. 2.12} \]

\[ \text{Eq. 2.13} \]

Figure 2.10. Equivalent circuits a) simple parallel resistance-capacitor (RC) circuit and its corresponding perfect semicircle response and b) double RC circuit, represented as a double arc.

EIS data analysis is typically made by graphical representation and modelling to an electrical equivalent circuit that allows the physical interpretation of the single characteristics. Nyquist-plot represents real (\( Z' \)) and imaginary (\( Z'' \)) part of impedance values, whose spectra have a semicircle...
shape. The high frequency intercept (for $\omega \rightarrow \infty$) with the real axis represents the purely ohmic resistance, $R_0$ of the cell, whereas the intercept at the lower frequency (for $\omega \rightarrow 0$) represents the differential resistance, which can be obtained from the corresponding I-U characteristic at the given operating point. The difference between the low and high frequency intercept is the so-called polarisation resistance ($R_p$) of the cell that is the sum of each single polarisation resistance caused by the loss mechanisms inside the cell. Figure 2.10 shows two examples of typical equivalent circuits and the corresponding representation in the complex plane. A simple parallel resistance-capacitor (RC) circuit will deliver a perfect semicircle response as shown in Figure 2.10a, whose extremes are in the origin and in $R_0$, and the maximum is found at $\omega_M = 1/(RC)$. Figure 2.10b is a double RC circuit, represented as a double arc if $R_3C_3 << R_2C_2$.

Electrodes in symmetrical cells were tested by EIS in two-point configuration. Input signal was 0 V DC – 20 mV AC amplitude signal in the $0.01 – 3 \cdot 10^5$ Hz frequency range. This signal was generated by a Solartron 1470E and a 1455A FRA module equipment. For anodes, EIS measurements were performed in the 650-900 ºC range, under moistened atmospheres (2.5% vol. H$_2$O) at different pH$_2$ (100%, 50% and 5% H$_2$). On the other hand, cathodes in symmetrical cells were measured under air in the 700-900 ºC temperature ranges. In all the cases, the total flow remained constant (100 mL·min$^{-1}$). Zview™ 2 fitting programme was employed to analyze the impedance spectra.

2.5. References


Chapter 3. Structural-transport properties relationships on $\text{Ce}_{1-x}\text{Ln}_x\text{O}_{2-\delta}$ system ($\text{Ln} = \text{Gd, La, Tb, Pr, Eu, Er, Yb, Nd}$) and effect of cobalt addition

Abstract

A large series of doped ceria has been prepared by co-precipitation method and completely characterized in order to have an overall understanding of the structural, oxygen vacancy concentration and transport properties. Several lanthanides have been dissolved in the $\text{CeO}_2$ fluorite structure and the effect of the final sintering temperature and the addition of cobalt oxide on the structural properties have been studied. The chosen lanthanides (Gd, La, Tb, Pr, Eu, Er, Yb and Nd) include a large range of ionic radii and different metals exhibiting variable oxidation states. The materials have been characterized by powder XRD, HT XRD, $\mu$-Raman spectroscopy, helium pycnometry and DC conductivity. Transport properties have been correlated with the structural features induced by the different ionic radii and variable oxidation state of the dopants. The highest ionic conductivity has been obtained for the less distorted cells (Gd and Nd doped ceria), which represent the optimum balance between Coulomb interactions, steric effects and vacancy distribution. The lowest $E_a$ value has been found for materials with long cell parameters.
3.1. Introduction

Pure CeO$_2$ is a fluorite structured oxide in which the cations have eightfold coordination. At low oxygen partial pressures, Ce$^{4+}$ cations are easily reduced to the Ce$^{3+}$ valence state in order to charge compensate the oxygen vacancy formation while the fluorite structure is retained. This material exhibits $n$-type electronic conductivity in reducing environments due to the formation of small polarons, which migrate by a thermally activated hopping. [1] The ceria fluorite phase is very tolerant to such reduction and to the dissolution of other oxides that introduce oxide vacancies and thus extend the electrolytic region (area of predominant ionic conductivity) over that of ceria. Nevertheless, both acceptor doping and reduced cation formation result in a strain termed chemical expansion, which produces stresses or constrictions in the structure. [1] Such strains affect the conductivity and might result in mechanical failure of the solid state device. [2, 3] Actually, the highest ionic conductivities are reported for cations causing the lowest size mismatch with respect to Ce$^{4+}$, *i.e.*, gadolinium and samarium. [4, 5]
Figure 3.1. Fluorite cubic structure of CeO$_2$. Doping occurs by the Ln substitution of a Ce cation.

By the mentioned modifications, doped cerias are able to combine high oxygen-ion mobility, redox catalytic properties, chemical compatibility with water and carbon dioxide at high temperatures and sufficient resistance to reduction under relatively low oxygen partial pressures. Therefore, depending on the main characteristics introduced by the dopant, these materials have been suggested and studied for a number of applications, e.g. electrolyte [6, 7] and anode [8, 9] in solid oxide fuel cells (SOFCs), mixed-conductive membranes for oxygen separation [10, 11] and partial oxidation of hydrocarbons, [12, 13] oxygen sensors, [14] oxygen storage materials and catalysts for industrial relevant processes. [15-18]

Special attention has been paid to doping with lanthanide (Ln) oxides as depicted in Figure 3.1, which are soluble in ceria up to values around 40%. [19,20] The addition of trivalent lanthanides promotes the ionic conductivity since an oxygen vacancy, $V_0^-$, is formed for every two substituting ions as indicated by the equation
where following the Kröger-Vink notation employed in the Chapter 1, $\text{Ln}_{\text{Ce}}$ means an acceptor dopant substitution in a Ce site. The oxygen vacancies are mostly free at low dopant content, however, as the dopant level increases defect associations due to Coulombic attractions might appear [4]. It has been widely reported that Sm provides high ionic conductivity combined with high resistance to reduction, followed by Gd. [5] That is why these two dopants are the most commonly used in ceria-based electrolytes for SOFC. [2, 4, 11] Doping with La increases the lattice constant of the ceria phase favouring the mobility of the ions, but the material becomes quite reducible instead. [5, 15, 21] Dy has been proposed on ceria systems as yttrium co-dopant and Er has been used as dopant in the catalytic ceria-zirconia system. [22, 23] The addition of elements with two valence states has been less studied. They increase the ionic conductivity, but some of these materials develop mixed ionic electronic conductivity, i.e., ceria doped with Tb and Pr. The rise in electronic conductivity and the lattice expansion undergone at low partial pressures, persuade the use of these materials as SOFC electrolyte. However, at high oxygen partial pressures ionic and $p$-type electronic conductivity are available in some materials and make them suitable for $\text{O}_2$ permeable membranes and SOFC cathodes. [24, 25] Specifically, Pr has been employed as dopant for gravimetric gas sensors because it increases the level of nonstoichiometry in ceria under oxidizing conditions. [26] However, it has a doping limit close to 30% due to the high and non linear thermal expansion behaviour and large chemical expansion upon reduction, which may lead to the device failure. [27] Thus, initial dopant content of 10% has been chosen. Trivalent lanthanides as Gd, La, Nd and Er have been added. Pr, Tb, Eu and Yb are elements with two possible valence states.
On the other hand, doping with lanthanide elements does not generally affect the sintering properties, with the exception of Yb, Nd and La. Their pellets or probes remain fragile and crumble on the edges despite they may give densities greater than 85-90% with regard to theoretical density. [28] Many metal oxides have been proposed to improve the density such as MnO$_2$, Mn$_2$O$_3$, Ga$_2$O$_3$, Co$_3$O$_4$, CoO and Fe$_2$O$_3$. [29, 30] They act as sintering aid because of their melting point, which is lower than that of ceria-lanthanide and thus promotes the atomic diffusion throughout the particle surface under high temperatures causing the sintering at lower temperatures. Cobalt oxide has been demonstrated to be an attractive sintering aid since it is able to reduce the sintering temperature more effectively than other metal oxides. [31] Cobalt oxide has been added to the same series of doped ceria in order to check whether it increases the mixed conductivity of these materials as previously reported for Gd and Pr doped ceria. [32, 33]

Previous reported works about series of Ln doped ceria are based on theoretical studies or on the collection of literature conductivity values, which are usually either quite scattered or does not take multivalent dopants into account. [34-36] The screening presented in this chapter deals with a systematic and exhaustive study of both structural and transport properties of several Ln substituted ceria compounds and the effect of Co as a sintering aid. First of all, the materials have been systematically prepared and characterized by powder XRD, μ-Raman spectroscopy, helium pycnometry and DC conductivity. Specifically, Raman spectroscopy is a widely used technique for structural analysis of ceria and lanthanide doped ceria due to its sensitivity to any structural changes (cell parameter and particle size) as well as to the oxygen vacancy concentration. [37, 38]
3.2. Results and discussion

Lanthanide doped ceria compounds have been prepared by the co-precipitation method explained in Chapter 2.1, in order to synthesize powders of nanometric size. Thermogravimetric analysis of Ce$_{0.9}$Tb$_{0.1}$O$_{2-\delta}$ and Ce$_{0.9}$Er$_{0.1}$O$_{2-\delta}$ precipitate powder dried at 150 ºC are shown in Figure 3.2. The decomposition of the precursor proceeds through two stages. The first weight loss may be attributed to the evaporation of the absorbed moisture and the release of ammonia and water. The second stage may correspond to the decomposition of the nitrates and (hydroxy)-carbonates. That proves that the mass is completely stabilized at 600 ºC and calcination at 800 ºC ensures that there will not remain any phase related to the precursor salts. Afterwards, the samples are sintered at 1300 ºC to perform the electrochemical experiments where dense specimens are required, so that characterization at this temperature was also carried out.

![Figure 3.2 Thermogravimetry analysis of Ce$_{0.9}$Tb$_{0.1}$O$_{2-\delta}$ and Ce$_{0.9}$Er$_{0.1}$O$_{2-\delta}$, in air. The TG was performed up to 800 ºC, the calcination temperature of the as prepared materials.](image-url)
XRD patterns of Ce$_{0.9}$Ln$_{0.1}$O$_{2-\delta}$ and Ce$_{0.9}$Ln$_{0.1}$O$_{2-\delta}$ +Co 2mol% series (Ln=La, Gd, Eu, Pr, Tb, Yb, Er) are depicted in Figure 3.3. After calcination at 800 ºC all the compounds show the expected cubic fluorite structure for CeO$_2$ (ICDD 00-043-1002). Diffraction peaks corresponding to any other lanthanide oxide or precursor are not observed, which means that the dopants are fully incorporated to the lattice of ceria. Likewise, the addition of Co does not cause the formation of any impurity phase. The stability is maintained at 1300 ºC. In this case, XRD patterns are sharper due to the crystallite size growing induced by the increasing temperature. XRD peak positions of doped specimens are shifted regarding to the pure CeO$_2$, which indicates deviations from its cell dimensions. The shift to shorter angle implies the enlargement of the unit cell, which happens for all dopants with ionic radius ($r_d$) bigger that of Ce$^{4+}$ ($r_c = 1.024$ Å), while deviations to higher $2\theta$ denote the shrinkage of the unit cell (dopants with $r_d < r_c$). The unit cell of the present cubic system is characterized by the lattice or cell parameter ($a$). It was extracted by fitting (4 2 2) reflection peak with a Gaussian function. [39] The lattice parameters for the $(h k l)$ plane were calculated using the equation 2.2 (Chapter 2) for a cubic system.
Figure 3.3. XRD pattern of Ce$_{0.9}$Ln$_{0.1}$O$_2$-$\delta$ and Ce$_{0.9}$Ln$_{0.1}$O$_2$-$\delta$ + Co 2mol% calcined at 800 ºC.

Figure 3.4 depicts the cell parameter of all the materials as a function of the trivalent ionic radii in 8-fold coordination given by Shannon for the two different annealing temperatures, 800 and 1300 ºC. [40] The cell parameter follows a linear correlation except for Pr and Tb. Despite the two possible oxidation states (+2/+3) of Eu and Yb, they remain trivalent in the fluorite structure. On the other hand, multivalent Pr and Tb (+3/+4) would need an apparent ionic radii between those corresponding to +3 and +4 oxidation states in order to preserve the general trend. By applying the lever rule, it is possible
to estimate, at room temperature, the amount of trivalent and tetravalent cations, which gives rise to 40% and 62% of trivalent cations for Pr and Tb respectively.

Figure 3.4. Lattice parameter of doped ceria as a function of the trivalent ionic radius of the dopant cation (and tetravalent for Pr and Tb) in eight-fold coordination.

Within the range of accuracy of the XRD device, Co addition does not cause any common cell parameter change. Several studies about Co addition in cerias attempted to elucidate the location of Co atoms. Chen et al. [41] studied the solubility of Co in CeO₂ and found a value of 3 mol% after sintering at 1580 °C. Also Co has been reported to be soluble less than 0.5 mol% in CGO, at 900 °C and diffuse into the grain boundary to form a film of roughly 0.5 nm thickness together with Gd or clustering particles. [42] For Pr doped ceria, the addition of Co at 2 mol% results in the apparent dissolution of Co while the addition of 5 mol% shows traces of Co oxides, observed by XRD, which are segregated along the grain boundaries. [33] On the contrary, Co causes shrinkage in the cell parameter of Tb doped ceria indicating that it might be soluble in the structure to some extent. Regardless the location, Co enhances the
sinterability and the grain size growth, as inferred from the sharpening of the XRD peaks of samples calcined at the same temperature with respect to the Co-free samples (Figure 3.3). [27]

SEM images for Tb, Pr, Eu and La doped ceria and analogous pictures of these materials with Co are shown in Figure 3.5. All the specimens were calcined at the same temperature (1300 °C). The mentioned differences in grain size when Co is added are observed, i.e., an average size of 0.3±0.1 µm is observed for samples without Co, while the average size increases up to 1.3±0.3 µm for Co containing compounds. This suggests that Co doping accelerates the densification rate and the grain growth remarkably. Works of Zhang et al. on the sintering behavior of several dopants included Co oxide. They reported that Co doping reduces the activation energy for grain growth of CeO₂, which would explain why Co doping leads to a rapid grain enlargement. [43, 44] The SEM pictures do not show difference of phases that could appear from CoOₓ or from its reaction with the doped ceria, although this is not concluding due to the limitations of the device and will be studied in more detail in Chapter 4. On the other hand, all samples show porosity, which should be taken into account depending on the application of the material, i.e., dense membranes or SOFC electrolyte, or porous electrodes.
Figure 3.5. SEM images of Ce$_{0.9}$Ln$_{0.1}$O$_{2-δ}$ and Ce$_{0.9}$Ln$_{0.1}$O$_{2-δ}$ + Co 2mol% calcined at 1300 °C (Ln = Tb, Pr, Eu and La).

The experimental density values have been obtained from mass and volume values acquired by a helium pycnometer and compared with the theoretical values calculated from the unit cell volume obtained by XRD. The results summarized in Table 3.1 indicate that almost all the materials achieve > 97% of the estimated density. Pr-doped ceria shows the highest deviation with a 92% of the expected value, which is ascribed principally to closed porosity. However we should take into account the experimental error obtained in these measurements (≈ 3%), which also explains densities over 100% for Gd doped ceria.
The real operating conditions in which the object materials of this thesis should work involve high temperatures and gradients of $pO_2$. In this context, different redox processes dependent on temperature and $pO_2$ may occur. Therefore, the evolution of cell parameters associated to thermal expansion and oxygen vacancy formation is studied by means of high temperature XRD (HT-XRD). The measurements are done by heating the specimens in air up to 900 °C and cooling them down in nitrogen. The fluorite structure is maintained in the whole temperature range and corresponding cell parameters are plotted in Figure 3.6 for some selected compositions. Ln= La, Gd and Eu show a linear dependency of cell parameter with temperature, which relates to thermal expansion coefficients (TEC) in air and in He respectively. On the other hand, Tb- and Pr-doped cerias show an increase in the slope at ca. 480 and 550 °C, respectively, when they are heated up in air. Accordingly, an additional chemical expansion coefficient (CEC) is calculated, which is assigned to the reduction of Ln$^{3+}$ to Ln$^{2+}$ and the associated release of molecular oxygen. CEC is not observed for Eu doped ceria, which confirms that there is not reduction of the cation either with the temperature or with lower $pO_2$ (N$_2$). When Pr and Tb

<table>
<thead>
<tr>
<th>Materials</th>
<th>$\rho_{\text{theor}}$ (±0.001) g/cm$^3$</th>
<th>$\rho_{\text{exp}}$ (±0.25) g/cm$^3$</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>CeO$_{2-d}$</td>
<td>7.209</td>
<td>7.11</td>
<td>98.6</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Pr$</em>{0.1}$O$_{2-d}$</td>
<td>7.197</td>
<td>6.63</td>
<td>92.1</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Yb$</em>{0.1}$O$_{2-d}$</td>
<td>7.345</td>
<td>7.29</td>
<td>99.3</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Eu$</em>{0.1}$O$_{2-d}$</td>
<td>7.190</td>
<td>6.76</td>
<td>94.0</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Tb$</em>{0.1}$O$_{2-d}$</td>
<td>7.304</td>
<td>7.25</td>
<td>99.3</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Gd$</em>{0.1}$O$_{2-d}$</td>
<td>7.245</td>
<td>7.45</td>
<td>102.8</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$La$</em>{0.1}$O$_{2-d}$</td>
<td>7.061</td>
<td>7.05</td>
<td>99.8</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Er$</em>{0.1}$O$_{2-d}$</td>
<td>7.314</td>
<td>7.12</td>
<td>97.4</td>
</tr>
</tbody>
</table>

Table 3.1. Theoretical and experimental density of doped cerias.
Doped samples are cooled down in N₂; the material cannot be re-oxidized due to the lack of molecular oxygen necessary for re-oxidation and only thermal expansion can be observed for the material in the achieved reduced state. TEC values in both atmospheres and CEC values in air are summarized in Table 3.2.

![Figure 3.6. HT-XRD heating up in air (left side) and cooling down in N₂ (right side) of Ce₀.₉Ln₀.₁O₂₋δ. In the insets, the calculated cell parameters (a 700 ºC) at 700 ºC in air (left side) and in N₂ (right side) are plotted as a function of the trivalent ionic radii (and tetravalent for Pr and Tb).](image)

Based on the HT-XRD cell parameters and the ionic radii for the different oxidation states, it is possible to estimate the amount of trivalent cation for Pr and Tb by comparing with those theoretical cell parameters of totally reduced and oxidized samples. The insets of plot the cell parameters in air (left) and N₂ (right) at 700 ºC as a function of the ionic radii (trivalent and tetravalent...
when required). From room temperature (RT) to 700 °C in air the estimated amount of trivalent cation is increased from 40% to 53% for Pr and from 62% to 78% for Tb. After heating the materials up to 900 °C in air only 9.4% of Tb$^{4+}$ remains and 16% of Pr$^{4+}$ is still present. By changing the atmosphere to N$_2$ both materials are completely reduced and the concentration of trivalent cations remains fixed during the cooling down in N$_2$ to room temperature. The relative amount of Pr and Tb cations in each of their oxidation states proved by HT-XRD is related to changes in the oxygen vacancy concentration whose importance will be discussed latter in the transport measurements.

<table>
<thead>
<tr>
<th>Ln</th>
<th>TEC in Air (K$^{-1}$)</th>
<th>TEC in N$_2$ (K$^{-1}$)</th>
<th>CEC in Air (K$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>11.9x10$^{-6}$</td>
<td>12.2x10$^{-6}$</td>
<td></td>
</tr>
<tr>
<td>Eu</td>
<td>11.9x10$^{-6}$</td>
<td>12.2x10$^{-6}$</td>
<td></td>
</tr>
<tr>
<td>Gd</td>
<td>11.9x10$^{-6}$</td>
<td>11.4x10$^{-6}$</td>
<td></td>
</tr>
<tr>
<td>Pr</td>
<td>11.3x10$^{-6}$</td>
<td>12.3x10$^{-6}$</td>
<td>11.51x10$^{-6}$</td>
</tr>
<tr>
<td>Tb</td>
<td>12.1x10$^{-6}$</td>
<td>12.3x10$^{-6}$</td>
<td>5.77x10$^{-6}$</td>
</tr>
</tbody>
</table>

Table 3.2. Thermal expansion coefficients calculated in Air and in N$_2$ for different La, Eu Gd, Pr and Tb doped ceria.

The creation of oxygen vacancies by the dopants causes changes in the lattice bond that may be detected by Raman spectroscopy. CeO$_2$ has a cubic fluorite structure that belongs to the O$^5_6$ (Fm$^{3}$m) space group. The F$_2$Raman-active mode frequency (at 465 cm$^{-1}$) can be understood as a symmetric breathing mode of the O atoms around each cation (Ce-O8 unit). Therefore, this unit should be very sensitive to any disorder in the oxygen sublattice due to induced non-stoichiometry. [45] The normalized Raman spectra obtained for Ce$_{0.9}$Ln$_{0.1}$O$_{2-δ}$ series (Ln=Pr, Yb, Eu, Tb, Gd, La) are plotted in Figure 3.7a, where they are compared with pure ceria. There are also
reported additional peaks close to $F_{2g}$ mode, mainly a shoulder at $\sim 570 \text{ cm}^{-1}$ attributed to the formation of oxygen vacancies (see inset of Figure 3.7a). The higher relative intensity observed for Pr and Tb of the $\sim 570 \text{ cm}^{-1}$ shoulder is ascribed to the higher optical adsorption for these coloured compounds (dark red) and not only to the oxygen vacancy existence. [37] There is also a broad band at 600 $\text{ cm}^{-1}$ consequence of either an ordering effect in the oxygen vacancies to produce confinement of the phonons within an inhomogeneous distribution of domains or structural distortions of the O sublattice. [46]

Figure 3.7. a) Raman spectra of the $\text{Ce}_{0.9}\text{Ln}_{0.1}\text{O}_{2-\delta}$ (Ln=La, Gd, Eu, Pr, Tb, Yb, Er) (sintered at 1300 ºC) from 400-750 $\text{ cm}^{-1}$ and b) $F_{2g}$ Raman shift as a function of the cell parameter $\text{Ce}_{0.9}\text{Ln}_{0.1}\text{O}_{2-\delta}$ and $\text{Ce}_{0.9}\text{Ln}_{0.1}\text{O}_{2-\delta} + \text{Co}$ calcined at 1300 ºC and 800 ºC (top) and after subtracting Gr"uneisen shifts (bottom). The inset shows a magnification of the band related to the oxygen vacancy formation.

The frequency shift ($\Delta \omega$) produced by a change in the lattice parameter ($\Delta a$) can be written in terms of the Gr"uneisen parameter:

$$\Delta \omega = -3\gamma \omega_0 \Delta a/a_0 \quad (\text{Eq. 3.2})$$
where \( \omega_0 \) and \( a_0 \) are the Raman frequency and the lattice parameter for CeO\(_2\) respectively, and \( \gamma \equiv (B/a_0) \text{d}\omega/\text{d}P \) is the Grüneisen constant (1.24 for CeO\(_2\) [37]). Therefore, the influence of the Ln doping on the structure can be detected by Raman spectroscopy. Figure 3.7b (top) plots the Raman shift of the F\(_{2g}\) mode as a function of the cell parameter. Raman mode shifts down as the cell parameter increases and only Pr doped CeO\(_2\) does not follow the linear tendency. The general trend agrees with expected Raman shift due to cell volume changes.

Other contributions, apart from volume, can be analyzed if the Grüneisen shifts are taken into account. Figure 3.7b (bottom) shows corrected Raman frequencies after subtracting Grüneisen shifts, as a function of the cell parameter. Now, the Raman shift increases with the cell parameter except for Pr, again out of the general trend. This shift up in frequency in the F\(_{2g}\) mode can be directly assigned to the increasing disorder produced by the oxygen vacancies. [37] It can be observed that La-doped material present the highest shift, while the behaviour of the Pr-doped compound remains unexplained in this work.

![Figure 3.8](image-url)  
Figure 3.8. Correlation length obtained from Raman line-shape fitting as a function of the cell parameter.
The defect concentration may be estimated from Raman intensity by using the spatial correlation length, [47] which considers them as spheres with diameters equal to the distance between two next oxygen vacancies. According to the spatial correlation model, Raman intensity $I(\omega)$ at a frequency $\omega$ can be written as [47]:

$$I = \int_0^1 \exp \left( -\frac{q^2 L^2}{4} \right) \frac{d^3 q}{[\omega - \omega(q)]^2 + (\Gamma / 2)^2}$$  \hspace{1cm} (Eq. 3.3)

where the wave vector $q$ is expressed in $2\pi/a$ units ($a$ is the lattice parameter), $L$ is the correlation length, and $\Gamma$ is the linewidth of the Raman mode (FWHM). $\exp \left( -\frac{q^2 L^2}{4} \right)$ is the Gaussian spatial correlation function and $\omega(q)$ is the phonon dispersion for selected mode, represented by parabolic fits to the CeO$_2$ phonon dispersion curves determined by Weber et al. [45] Figure 3.8 shows the correlation length obtained from Raman line-shape fitting as a function of the cell parameter. Correlation length is 1.7 and 1.3 nm for CeO$_2$ (grain size 85 nm) and Co doped CeO$_2$ (grain size 87.7 nm) respectively, while for the rare earth doped samples (with and without Co) the correlation length decreases and it is shorter than the unit cell. It has been previously reported that when the oxygen vacancy concentration reaches $10^{21}$ cm$^{-3}$, practically each unit cell is influenced by defects and then the correlation length, which is related to the short-range order parameter, should be attributed to the average oxygen-oxygen distance.

When 10% of Ln dopant is introduced in ceria samples there is an induced oxygen defect concentration of around $4 \times 10^{21}$ cm$^{-3}$ so, in this case, the correlation length corresponds to the oxygen – oxygen averaged distances, which is in all cases close to 2.71 Å. [48, 49] For pure and Co added ceria the defect concentration is three orders of magnitude smaller and the correlation length can be assigned to the distance between two next oxygen vacancies. This
defect concentration can be seen as the volume between two next oxygen vacancies, and assuming they are spherical, it is possible to calculate the defect concentration \( N \) as a function of the correlation length by \( N = \frac{3}{4\pi l^3} \). Then \( 5 \times 10^{19} \) cm\(^{-3} \) is obtained for CeO\(_2\) and \( 1 \times 10^{20} \) cm\(^{-3} \) for Co doped CeO\(_2\), which is in agreement with other reported values. [48]

Raman spectroscopy results demonstrate that the introduction of Ln\(^{3+}\) dopants produces a disorder due to the oxygen vacancies that increases with the cell parameter. The generated disorder is reflected in transport properties, whose features are analysed in Figure 3.9 and Figure 3.10. For the sake of simplicity dopants are displayed in two different plots and compared with CeO\(_2\) as a reference.

Figure 3.9a shows a representative sample of each kind of dopant: CeO\(_2\) as a reference, Eu (+2/+3), Tb and Pr (+3/+4) and Er (+3). The Arrhenius plot for the conductivity in air of the selected materials shows that any dopant improves the total conductivity of pure ceria due to the generation of extrinsic vacancies. Trivalent dopants may enhance the ionic conductivity better than those in a mixed valence as Tb or Pr, as they produce a fixed number of oxygen vacancies at this \( pO_2 \). The oxygen release and consequent reduction of the dopant caused by the rising temperature results in a change of the Arrhenius slope (apparent activation energy) for Tb and Pr doped ceria in air. Actually, the temperature of change in the slope matches that observed by HT-XRD analysis. On the other hand, the conductivity results for the rest of samples available in Figure 3.10a (Yb, Gd, La and Nd) do not show such transition. Indeed, it is expected for fixed valence dopants and indicates that Yb and Eu remain in their +3 oxidation state.
Figure 3.9. a) Arrhenius plot of the total conductivity in air of CeO\(_2\) and Ce\(_{0.9}\)Ln\(_{0.1}\)O\(_{2-\delta}\) (Ln=Tb, Pr, Eu and Er), and b) log-log plot of total conductivity at 700 °C of the Co-free and Co-containing CeO\(_2\) and Ce\(_{0.9}\)Ln\(_{0.1}\)O\(_{2-\delta}\) (Ln=Tb, Pr, Eu and Er) as a function of \(pO_2\).

Figure 3.10. a) Arrhenius plot of the total conductivity in air of Co-free Ce\(_{0.9}\)Ln\(_{0.1}\)O\(_{2-\delta}\) Ce\(_{0.9}\)Ln\(_{0.1}\)O\(_{2-\delta}\) (Ln=Yb, Gd, La and Nd) and b) log-log plot of the total conductivity versus \(pO_2\) at 700 °C of the Co-free and Co-containing CeO\(_2\) and Ce\(_{0.9}\)Ln\(_{0.1}\)O\(_{2-\delta}\) Ce\(_{0.9}\)Ln\(_{0.1}\)O\(_{2-\delta}\) (Ln=Yb, Gd, La and Nd).
Figure 3.9b shows the conductivity dependence on the $pO_2$ for the selected materials with and without Co taken at 700 °C. The studied range of $pO_2$ includes from oxidant (0.21-1 atm) to moderately reducing atmospheres (10^-5 - 0.21 atm), where the Ce^{4+} does not reduce to Ce^{3+}, since it has been demonstrated that this reaction initiates at oxygen partial pressures lower than 10^{-10} atm. [27, 50-52] Except for Tb and Pr, all the materials show a mainly ionic conductivity behaviour, which is independent on $pO_2$. Tb and Pr doped ceria present a dependency on $pO_2^{-1/6}$ at the most oxidizing atmospheres attributed to the dopant reduction as $pO_2$ decreases, and which turns into a $pO_2$ independent plateau. [53] Taking into account the defect chemistry explained in the Chapter 1 (see Table 1.2), the conductivity is identified as mainly ionic in these oxidizing conditions and the multiple oxidation state of some dopants makes the ionic conductivity $pO_2$ dependent (region not observed in single-oxidation-state acceptor doped systems as Er and Eu). In fact, this behaviour is well in agreement with the defect chemistry of an acceptor doped fluorite working in the boundary between region III and region IV developed in Figure 1.4. It is worthy to mention here that the defect concentration is also determined by temperature, so that there will be a different diagram at each temperature for any given compound. That means the $pO_2$ region edges will shift as a function of temperature and related to the enthalpy of the diverse defect reactions.

In general the addition of cobalt to doped ceria improves the total conductivity being especially evident for Tb and Pr. The conductivity enhancement is ascribed to the creation of percolating electronic paths through the grain boundary, where a nanosized Co-rich phase has been reported to be preferentially located. [27] Moreover, Pr and Tb with cobalt addition show enhanced total conductivity due to (a) the reducibility of the material at lower temperature with respect to the Co-free samples; (b) the electronic conductivity promotion; and (c) the higher sample density and grain packing due to the high
sintering activity of Co-containing ceria observed by SEM (Figure 3.5). [27, 31, 42]

Relationship between structural and transport properties

From the conductivity features, it may be stated that the dopants in fixed trivalent state contribute only ionic charge carriers to the total conductivity. In that case, for the same dopant concentration, differences in conductivity should be related not only to the defects, but also to the mobility of ions over the oxygen vacancies formed in the fluorite lattice. In the following section the correlation between structural and transport properties of these series of lanthanide doped ceria is discussed. It has been widely reported that a dopant minimizing the internal strain will maximize the ionic conductivity as it has been traditionally observed for Gd or Sm doped ceria. [54] Some computational studies are devoted to the understanding of the factors that affect the ionic conductivity in doped ceria, as dopant ionic radius or clustering phenomena and how they are reflected in the mobility of ions. [35, 55, 56] They allow identifying and evaluating the energy barriers for diffusion inside the ceria lattice, which are related to the total activation energy ($E_a$).

Fundamental physics behind the ionic conductivity determine that $E_a$ is equal to the sum of the vacancy formation energy ($E_f$) and the migration barrier ($E_m$) for pure ceria. The introduction of aliovalent dopants produces extrinsic oxygen vacancies whose effective charges are opposite so that they may attract each other forming a $\text{Ln}_\text{Ce}^2_\text{-V}_\text{O}^2_\cdot\cdot\cdot\cdot$ (vacancy-dopant) associate. Vacancy-dopant association energy ($E_{\text{ass}}$) also influences the vacancy mobility. Hence, $E_a$ can be understood as the sum of $E_m$ and $E_{\text{ass}}$ for bulk diffusion in doped ceria. Quantum mechanical calculations on Sm-doped ceria have revealed $E_m$ values around 0.46 eV, when there are no trivalent ions around a vacancy, while $E_m$ values
increase up to ~0.5 eV when neighboring dopant cations are present. This is due to the oxygen atom movement through a point close to large trivalent ions is less favorable than to move to smaller Ce4+ ions. [56, 57] On the other hand, $E_{\text{ass}}$ depends on the distribution and concentration of the trivalent ions. In the present work the concentration of trivalent ions is the same (10%) for all dopants except for Tb and Pr that have a variable concentration depending on temperature and $pO_2$. Thus, the vacancy-dopant distribution will be essential in the $E_{\text{ass}}$ performance.

![Figure 3.11: $E_a$ (top) and total conductivity at 700 ºC in air (bottom) of the different doped materials versus cell parameter measured at 700 ºC. Cell parameters at 700 ºC in air have been calculated from HT-XRD or by using TEC values for similar dopants. $E_a$ solid lines have been fitted to Eq. 4 only taken into account samples with trivalent dopants while dashed fits include Pr and Tb.](image)

The experimental $E_a$ values derived from the conductivity measurements corresponding to the different samples are represented in Figure
3.11 (top) as a function of the cell parameter at 700 °C in air. $E_a$ has been calculated over the full temperature range for samples with trivalent dopants. For Tb and Pr doped ceria, the plotted $E_a$ was the one in the 800-650 °C range that includes the temperature at which the cell parameter was measured.

$E_a$ decreases with the increasing cell parameter of the materials following a trend proportional to $1/a^2$ and tend to reach a fix value of 0.73 eV. The observed tendency can be easily related to coulombic interactions and steric effects due to the strong dependency of the displacement distance upon dopant type and relative dopant-vacancy positions. [58] The lowest $E_a$, obtained for samples with the largest cell parameters, are related to materials with an optimal balance between elastic (lattice distortions) and electronic defect interactions (redistribution of the electronic structure). When the cell parameter is shorter than that of pure ceria (which is approximately equal to Gd doped material) the structure is shrunk and $E_a$ follows Coulombic tendency. For longer cell parameters, from the point where the structure is not changed (Gd doped) up to the largest cell parameter (La doped) the Coulombic limit is overpassed and $E_a$ reaches a steady value defined by dopant concentration.

Experimental data has been fitted following Eq. 3.4

$$E_a = E_0 + \frac{A}{a^2-B^2}$$  \hspace{1cm} (Eq. 3.4)

where $a$ is the calculated cell parameter at 700 °C; $E_0$ is the limit where Coulomb interactions do not affect the total energy for sufficient large cell parameters; $A$ is a constant that principally includes charges and electric constants; and $B$ is the minimum cell parameter that allows oxygen diffusivity. [58]
Solid lines have been fitted to Eq. 3.4 including only samples with trivalent dopants and dashed fits also include Pr and Tb. The deviation from general trend of these dopants, especially Pr, is due to the remaining and variable tetravalent cations. This fact confirms the importance of the interatomic distances that govern $E_a$ behavior, as they present a cell parameter according to their mixed valence state (as discussed in Figure 3.4 and 3.6) and almost fall in the same trend as all the other trivalent dopants. In fact $E_a$ decreases for Pr and Tb doped samples at lower $pO_2$ (the more trivalent dopant the higher vacancy density, and thus longer cell parameter) while trivalent dopants maintain their $E_a$ because there are not new structural changes. Raman measurements also confirm the $E_a$ behavior, since they predicted the highest oxygen vacancy disorder, and thus the lowest $E_{ass}$, for the longest cell parameters (Figure 3.7b) and the exceptional behavior of Pr doped samples.

By fitting to Eq. 3.4, a steady $E_0$ value of 0.70±0.08 eV (0.68±0.16 eV considering Tb and Pr) for Co-free series and 0.77±0.08 eV (0.76±0.13 eV considering Tb and Pr) for Co-containing samples have been obtained. The higher activation energy of Co-doped samples may result from the location of Co along the grain boundary as mentioned above. [42] It is worthy to note the opposite behavior of the Tb doped sample, whose $E_a$ decreases when Co is added. It may be attributed to the partial incorporation of Co in the ceria solid phase and increasing in oxygen vacancy concentration as will be studied in detail in Chapter 5. The minimum cell parameter that allows oxygen diffusivity, $B$, was found to be close to 5.4±0.9 Å. As the accuracy of $B$ parameter is not high enough, it can be found in the literature a minimum cell parameter of 5.3987 Å corresponding to Lu, the smallest rare earth dopant, which presents the minimum ionic conductivity. [59] Finally, $E_a \approx E_0$ is found for non-distorted cell materials and longer cell parameters i.e., Gd.
The influence of the structural parameters on the transport properties is also illustrated in Figure 3.11 (bottom), which represents the total conductivity of all the samples (with and without Co) measured in air 700 ºC as a function of the cell parameter at 700 ºC in air. Predominant ionic conductivity is expected and, confirming reported studies, there is a maximum conductivity for Gd and Nd doped ceria due to the low association energy between the oxygen vacancies and the trivalent dopants and the lowest migration energy, minimized to those materials with cell parameters close to pure ceria, which represents the less distorted cell as well. [1, 59] Furthermore, it should be pointed out that electronic transport begins to contribute to the total transport for the Tb and Pr doped materials with Co addition, which could explain the dispersion of the data for these materials.

### 3.3. Conclusions

A large series of partially-substituted cerias has been prepared and characterized by XRD, μ-Raman spectroscopy and conductivity measurements in order to have an overall understanding on the relationship of structural and transport properties. Several lanthanides (Gd, La, Tb, Pr, Eu, Er, Yb and Nd) have been dissolved in the fluorite structure. XRD and HT-XRD shows that the doping produces modifications in the cell parameter of the pure ceria, which follows a linear dependency on the trivalent ionic radii of the dopants. Tb and Pr are out of this tendency due to their mixed 3+/4+ oxidation state. The amount of each dopant in each oxidation state has been estimated finding that Eu and Yb remain always trivalent but the amount of trivalent Pr and Tb depends on temperature and atmosphere. Raman spectroscopy allowed studying the
substitution effect and Co-doping on the final fluorite structure and estimating the vacancy disorder.

The lowest $E_a$ value was found for materials with long cell parameters, which is reached by the non distorted cell (Gd) and by cells with longer lattice parameters. This is ascribed to the decreasing Coulombic interactions combined with the lowest $E_{\text{ave}}$ due to the higher vacancy disorder, which was predicted by Raman. Once the cell parameter of the non distorted cell is overwhelmed, the increase in ionic radii is not joined by the ionic conductivity improvement. Vacancy-dopant concentration and distribution affect the total conductivity, giving rise to the highest ionic conductivity for the non distorted cells (Gd-doped ceria), which represent the optimum balance between Coulomb interactions, steric effects and vacancy distribution.

This first screening permits the reorientation of the next steps on the research of new electronic and/or ionic materials. Questions about macrostructure optimization of the sintering aids, the possibility of the electronic promotion in multivalent doped ceria and potential in their redox reactions or the possibility to enhance or introduce new properties by co-doping the present materials will be formulated and discussed in the upcoming chapters.

### 3.4. References


Structural-transport properties relationships on Ce$_{1-x}$Ln$_x$O$_{2-δ}$ system


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Chapter 4. Microstructure

optimization of $Ce_{0.9}Pr_{0.1}O_{2-\delta} + Co \, 2\text{mol\%}$

Abstract

Praseodymium doped ceria shows a combination of mixed ionic and electronic conductivity, redox catalytic properties and chemical compatibility with water and carbon dioxide at high temperatures. Minor additions of cobalt oxide have been demonstrated to be a sintering aid and an electronic conduction promoter. However, an excess of sintering temperature causes cobalt aggregation into the grain boundaries, which is demonstrated by scanning electron microscopy (SEM) and temperature programmed desorption (TPD). In this chapter, the evolution of the microstructure has been investigated in order to understand the role of Co and to optimize both electronic and oxygen ion transport properties.
4.1. Introduction

By doping with multivalent cations, mixed conduction in ceria may be further extended to higher oxygen partial pressure range ($10^{-5}-10^{-1}$ atm). In the previous chapter, terbium and praseodymium doped ceria have shown MIEC properties arising from their possible $3+/4+$ valences at these $pO_2$ thus being suitable for the development of oxygen permeation membranes. [1-3] However, the low electronic conductivity is still the bottleneck for the oxygen permeation. The addition of small amounts of cobalt oxide is a strategy to improve the electronic conductivity and the material sintering. Since this addition modifies the microstructure, the sintering conditions need to be optimized.

For that purpose, a systematic study of $\text{Ce}_{0.9}\text{Pr}_{0.1}\text{O}_{2-\delta} + \text{Co 2mol\%}$ synthesized by co-precipitation and impregnation and sintered at different temperatures was carried out. XRD, FE-SEM and TEM were employed on the structural characterization. The electrochemical properties were studied by means of DC-conductivity, which results allow finding a correlation between the grain boundary length, composition and the conduction properties. The aim is to engineer the grain boundary contribution to the total conductivity by promoting the electronic conductivity and to optimize the compound micro and macrostructure for oxygen membrane applications.
4.2. Results and discussion

The initial powders of Ce$_{0.9}$Pr$_{0.1}$O$_{2-\delta}$ (CP) and Ce$_{0.9}$Pr$_{0.1}$O$_{2-\delta}$+Co 2mol% (CPCo) after co-precipitation were calcined at 800 °C, since this temperature is sufficient to ensure the elimination of the precursor nitrates and carbonates (Chapter 3). Subsequently, rectangular probes (4 x 0.4 x 0.2 cm$^3$) of the powders fired at 800 °C were uniaxially pressed at 125 MPa during 1 minute and subsequently sintered 5 hours at 900, 1000, 1100, 1200, 1300 and 1400 °C in air atmosphere. Besides, one probe of CP was calcined at 1300 °C for comparison with CPCo.

In first place, the effect of the Co addition on CP is considered. XRD patterns of CP and CPCo calcined at 1300 °C show a cubic fluorite structure, which is depicted in Figure 4.1a. [4] The obtained patterns indicate that praseodymium forms a solid solution and incorporates in the lattice of ceria. Diffraction peaks corresponding to any precursor or Co oxides were not observed on the initial powder to the limit of the apparatus. As previously reported, CoO$_x$ is a sintering aid that enables lowering the densification temperature with respect to the Co-free specimen. [5] Thus, CPCo sample shows sharper and more intense diffraction peaks for the same calcination temperature as CP. This is a consequence of the grain size enlargement, which suggests that CoO$_x$ presence accelerates the densification rate and grain growth. As it has been reported and mentioned in Chapter 3, the addition of Co oxides may reduce the activation energy of grain growth of CeO$_2$, causing larger grains for CPCo with respect to CP. [6, 7] SEM images in Figure 4.1a illustrate this effect for CP and CPCo materials sintered at 1300 °C.
Figure 4.1. a) XRD patterns and SEM images and b) temperature dependence of cell parameter of CP and CPCo sintered at 1300 °C.

CPCo cell parameter, calculated from the XRD measured in air at room temperature, was 5.412 ± 0.001 Å, slightly higher than that obtained for CP (5.407 ± 0.001 Å). This fact points out that (i) the Co is not incorporated in the ceria lattice, as it would result in the shrinkage of the cell parameter (the ionic radius of Co$^{2+}$ in 8-fold coordination is 0.9 Å, smaller than that of Pr$^{3+}$ and Ce$^{4+}$ cations, 1.126 and 0.97 Å, respectively) and (ii) the catalytic effect of Co on the reduction of part of Pr$^{4+}$ to Pr$^{3+}$, with higher ionic radius and thus increasing the cell parameter. [8]

On the other hand, different redox processes dependent on temperature and $pO_2$ occur at the most interesting operating conditions for this kind of materials as oxygen separation membranes and SOFC cathodes. In order to check the evolution of the cell parameter and associated vacancy concentration at high temperature, XRD measurements were performed on CP and CPCo sintered at 1300 °C by heating the specimens in air up to 900 °C and cooling...
them down in N₂. The results plotted in Figure 4.1b show a linear dependence of cell parameter with temperature and an increase of the slope at 560 °C. The first slope corresponds to the thermal expansion coefficient in air. Above 560 °C, the increase in slope is caused by the additional contribution of chemical expansion due to the reduction of Pr⁴⁺ to Pr³⁺, the consequent expansion of the ionic radius and the associated oxygen release. From these results, a linear chemical expansion coefficient can be estimated from 830 to 900 °C. After stabilization at the highest temperature the atmosphere is switched to N₂ and the sample is cooled down. Both samples in N₂ have similar cell parameters and expansion coefficients due to the achievement of totally reduced Pr³⁺ concentration at this pO₂. The material is not re-oxidized upon cooling due to the lack of molecular oxygen, and only thermal expansion/shrinkage can be observed. All the coefficients are summarized in Table 4.1 considering an experimental error of 1%. It should be noted that the linear thermal expansion coefficient coincides for both oxidized and reduced material.

<table>
<thead>
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<th>Material</th>
<th>α_L, AIR₁ (x10⁶), K⁻¹</th>
<th>α_L, AIR₂ (x10⁶), K⁻¹</th>
<th>α_L, N₂ (x10⁶), K⁻¹</th>
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</thead>
<tbody>
<tr>
<td>CP</td>
<td>11.4</td>
<td>11.4</td>
<td>12.3</td>
</tr>
<tr>
<td>CPCo</td>
<td>11.6</td>
<td>10.4</td>
<td>12.7</td>
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Table 4.1 Thermal expansion coefficient of Ce₀.₉Pr₀.₁O₂₋ and CPCo 2% sintered at 1300 °C.

Temperature programmed reduction in H₂ (H₂-TPR) measurements were done in order to analyze the redox properties of pure CeO₂ and CP. The effect of Co addition is studied as well for the two bulk compositions. The profiles depicted in Figure 4.2 demonstrate the higher reducibility of the Co containing samples in terms of lower temperature reduction and H₂ consumption. H₂-TPR Figure 4.2a shows the H₂-TPR plot for the pure CeO₂ and CP samples calcined in air at 1300 °C. The CeO₂ sample show peaks at 440 and
860 ºC, which are ascribed to the easily reducible surface oxygen and that in the bulk ceria, respectively. Doping with Pr, however, shows multiple reduction peaks. The first one around 400 ºC overlaps several peaks attributed principally to Pr$^{4+}$ reduction (380 ºC) and in a minor degree to Ce$^{4+}$ surface reduction (shoulder at 440 ºC as observed in pure ceria). The hydrogen consumption at this low temperature range corresponds to an oxygen substoichiometry of 0.05. Consequently, a large percentage of Pr cations in ceria are originally expected in Pr$^{4+}$ oxidized state. The second peak is attributed to the bulk Ce$^{4+}$ reduction temperature, which is substantially diminished up to 720 ºC. This shift in temperature has been attributed in other works to the oxygen vacancies provided by the Pr, which increases the oxygen mobility in the lattice. [10]

Figure 4.2. a) TPR profile of CeO$_2$ and CP and b) TPR profile of CeO$_2$ + 2mol% Co and CPCo. All the samples are samples sintered at 1300 ºC. The flow gas is 10% H$_2$ in Ar; heating ramp was 5 ºC/min.
Figure 4.2b represents the effect of Co on the reduction temperature. The spectrum of CeO$_2$ presents a new reduction peak centered at 347 °C related to reducible species, Co$^{3+/2+}$ (which masks the surface Ce$^{4+}$ peak). With respect to the bulk ceria, the reducibility temperature decreases around 100 °C. The system becomes more complicated when the cobalt is added to Pr doped ceria. Up to four overlapped peaks can be found CPCo reduction in the range from 300 to 350 °C. These peaks are related to the reduction of Co$^{3+/2+}$, Pr$^{4+/3+}$ and in a minor extent of surface Ce$^{4+}$. Bulk ceria reduction occurs also at lower temperature, i.e., the maximum reduction peak is observed at 675 °C. Cobalt has a catalytic effect on the reduction of ceria which shifts bulk ceria reduction temperature in all cases. Moreover, the cobalt addition allows increasing the total amount of ceria reduced, i.e., the final oxygen substoichiometry. Specifically, the final vacancy concentration is increased as follows: 15% (CeO$_{2-x}$), and 30% (CP). From the point of view of the material application (as oxygen membranes or SOFC electrodes), the improvement in the reducibility would bring the possibility of reducing the operation temperatures.

The following section focuses on the optimization of the CPCo sintering conditions. The variation of the sintering temperature may induce substantial changes in the microstructure and consequently in the redox and transport properties. [11] The SEM pictures of CPCo exposed in Figure 4.3 show how the grain size evolves as a function of the sintering temperature (from 900 to 1450 °C). The grain size grows from the 60 nm of sample sintered at 900 °C up to 7 µm at 1450 °C. The density is improved as sintering temperature rises.
Changes in composition of the different grains and element distribution can be distinguished by backscattered electron (BS) FE-SEM. The pictures in Figure 4.4a and b show two different phases for both CPCo samples sintered at 1000 and 1450 °C, respectively. An EDX mapping analysis (Figure 4.4d) over the BS image of Figure 4.4c indicates that the dark particles correspond to CoOₓ and spots its distribution along the grain boundary (GB). [14] Further, the sintering temperature influences the size of these CoOₓ particles. The sample sintered at 1000 °C shows CoOₓ nanoparticles (below 50 nm). However, at 1450 °C these nanoparticles appear spread over the grain boundary surface next to aggregates of 1.5-2 µm in a non-homogeneous distribution.
Figure 4.4. Back scattering SEM of CPCo sintered at a) 1000 ºC and b) 1450 ºC. c) Less magnified image of CPCo at 1450 ºC and d) EDS Co mapping indicating Co rich zones. Arrows indicate the presence of the CoO$_x$ particles.

Indeed, the TEM images of a focused ion beam (FIB) lamella of Figure 4.5 show clean grain boundaries, with no reacting layers. The Co appears well distributed in the sample at 1000 ºC, with fine grain interface and it may be detected by EDS (Figure 4.6). On the other hand, the interface between grains of PCCo at 1450 ºC is very fine and no traces of Co have been found by EDS, even in the fringes (Figure 4.7). This proves that the CoO$_x$ particles are out from the lattice, and they agglomerate as a secondary phase that was not possible to find in such a small lamella.
Figure 4.5. FIB-lamella TEM of CPCo sintered at 1000 °C (left) and 1450 °C (right).

Figure 4.6. TEM-EDS on a 1000 °C CPCo grain. Co appears dispersed along the grain surface, while fine CP-CP grain interface is shown.
Chapter 4

Figure 4.7. TEM-EDS on a 1450 °C CPCo grain boundary. A fine CP-CP grain interface is shown. Co was not detected either in the grain boundaries or in fringes and surfaces.

XRD patterns of CPCo samples sintered at 1000, 1200 and 1450 °C are plotted in Figure 4.8a. All compositions demonstrate the cubic defect fluorite structure. More accurate XRD measurements of CPCo at 1000, 1200 and 1450 °C have revealed some low intensity peaks when they are represented in logarithmic scale. These peaks are due to the steel sample holder and not to the CoO$_x$ related species observed by FE-SEM, which indicates than the 2 mol% of cobalt is below the resolution of the diffractometer.

On the other hand, although all samples are fluorite structured and they have the same nominal composition, they may show different redox properties related to the microstructure. In order to check the oxygen release and simultaneous cation reduction and oxygen vacancy formation of the material, temperature programmed desorption (TPD) of CPCo sintered at 1000, 1200 and 1450 °C have been carried out in Ar as plotted in Figure 4.8b. For comparison,
TPD of CP was also measured and plotted in Figure 4.8b (bottom). All compounds have a similar oxygen desorption peak at around 600 °C. As Ce⁴⁺ reduction initiates at oxygen partial pressures lower than 10⁻¹⁰ atm [1, 8, 12, 13] and keeps nearly constant at higher oxygen contents (in the present study \( pO_2 > 10^{-5} \) atm), the observed reduction is most likely ascribed to the reduction of Pr⁴⁺→Pr³⁺ and would cause the chemical expansion of the lattice previously observed by HT-XRD (Figure 4.1b). A second peak appears at around 790 °C only for the three Co containing samples, which indicates a different reduction process corresponding to [Co₃O₄] Co³⁺→Co²⁺[CoO]. [14] The relative intensity of this peak decreases as sintering temperature rises. It can be ascribed to the better Co distribution at lower sintering temperatures, as showed by SEM analysis.

Figure 4.8. a) XRD patterns and b) TPD of oxygen in Ar for CPCo sintered at 1000, 1200 and 1450 °C. TPD of CP is also depicted for comparison.

The oxygen-release thermal behavior, i.e., the specific oxygen non-stoichiometry, is directly related to the solid-state transport properties (both
ionic and electronic transport). In this regard, the total conductivity in air at 800 °C and 400 °C is represented as a function of the sintering temperature in the Figure 4.9a. The highest total conductivity value observed at 400 °C is found for the sample sintered at the lowest temperature (i.e., 900 °C). At 800 °C, the sample sintered at 1000 °C shows the highest total conductivity. The different behaviour observed at different operating temperatures suggests that the dominating transport mechanisms strongly depends on the operating temperature and sintering temperature of the sample.

Figure 4.9a Total conductivity in air at 800 °C and 400 °C as a function of the sintering temperature and b) total conductivity of CPCo sintered at five different temperatures in air. For comparison, conductivity at $5 \times 10^{-5}$ atm for CPCo sintered at 1000 °C is plotted.

Figure 4.9b shows the conductivity ln(σT) as a function of the reverse temperature for the CPCo samples sintered at five different temperatures and measured in air. Changes in activation energy ($E_a$) are observed at high $pO_2$ as temperature drops due to oxygen re-incorporation. This process has been observed in Chapter 3 as a consequence of the thermal reduction of dopants that
may be present in more than one oxidation state, as the case of Pr. Actually, $E_a$ change of CPCo in air is produced around 560 °C, which matches temperature range of the oxygen desorption/incorporation shown by HT-XRD Figure 4.1b and by TPD (Figure 4.8). For comparison, the graph also plots the conductivity at $5 \times 10^{-5}$ atm for CPCo sintered at 1000 °C. At this $pO_2$ (lower than $10^{-3}$ atm), the oxygen incorporation in cooling down does not take place and only one $E_a$ is observed. The similar behavior of all the samples suggests that differences in conductivity are not caused by disparities in the composition.

In order to shed further light on the specific effect of the cobalt addition and minimize the influence of the microstructure change, the following comparison is done. Figure 4.10a plots the conductivity in air of (i) CPCo and Co-free CP sintered at 1300 °C and (ii) CPCo sintered at 1000 °C, which shows same grain size as CP sintered at 1300 °C. As aforementioned (Figure 4.1), the addition of Co to CP, causes the grain size growth. The conductivity at high temperatures is lower for CPCo with respect to CP, possibly due to the grain size. However, at low temperatures, CPCo sintered at 1300 °C presents higher total conductivity, which is attributed to electronic contribution. When comparing samples with similar grain size, \textit{i.e.}, CPCo sintered at 1000 °C and CP sintered at 1300 °C, the conductivity of the former is higher over the whole temperature range. This is ascribed to the homogeneous spreading of Co-enriched nanoparticles over the grains (observed in the BS/FE-SEM image Figure 4.4), which promotes the preferential percolation of electronic carriers through the grain boundaries.
Figure 4.10. a) Arrhenius plot of the total conductivity of CPCo sintered at 1000 and 1300 °C and CP sintered at 1300 °C, in air. b) \( pO_2 \) dependence of the total conductivity of CPCo sintered at different temperatures, 400 °C (open symbols) and 800 °C (solid symbols).

The results described Chapter 3 show that different electronic and ionic charge carriers contribute to the total conductivity in multivalent doped ceria. The specific contribution depends on the dopant content, operating temperature range and \( pO_2 \). Pr reduction (from Pr\(^{4+}\) to Pr\(^{3+}\)) results in a \( pO_2 \) dependent ionic conductivity (-1/6 dependency) as well as in the introduction of electronic levels within the energy gap of ceria. [15, 16] On the other hand, the electronic conductivity improvement by the addition of Co has been reported. [17] Figure 4.10b depicts the \( pO_2 \) dependency of the total conductivity measured at 400 and 800 °C, in a log-log representation. The conductivity at 400 °C of the CPCo specimen sintered 1450 °C follows a \( pO_2^{-1/6} \) dependence up to \( pO_2 = 10^{-2} \) atm, when it reaches a plateau. This dependence is stepwise attenuated as sintering temperatures are lower, up to the nearly independent-like behavior of CPCo sintered at 900 °C. Such dependence mitigation may be due to the \( p \)-type electronic contribution, which follows a counter 1/6 slope.
Since all the samples have the same nominal compositions, these behaviour differences should be most likely caused by microstructural issues. The conductivity enhancement observed for the low temperature sintered CPCo is related to their smaller grain size. Small grain size distribution produces the homogenous spreading of the Co additive, observed along the grain boundary (GB) area by BS FE-SEM (Figure 4.4a). As the sintering temperature rises, not only the fluorite grain size grows but also the Co-enriched particles aggregate and grow substantially in isolated positions among three or more fluorite grains (see Figure 4.4). Under this scenario, the electronic percolating network provided by the CoOₓ nanoparticles is not continuous and the contribution to the electronic conductivity along GB becomes negligible. In fact, the lowest conductivity values are obtained for the sample sintered at 1450 °C, for which mainly ionic bulk transport through large grains. At 800 °C the conductivity for this sample is apparently pO₂ independent, due to the prevailing contribution of ionic carriers, particularly considering that at this temperature most of the Pr cations are reduced to Pr³⁺ and the oxygen vacancies concentration is the highest. Under these conditions very large grains result in a very low electronic conductivity and the negligible effect of grain boundaries. Additionally, the transport in CPCo samples with very small grains will be more influenced by the grain boundary mechanisms, which in this case turns to be favorable for the mixed conductivity thanks to the well-distributed CoOₓ electronic percolating network. As a result of this microstructure, the most performing material is that sintered at 1000 °C. This conductivity results suggest that by tailoring grain sizes and Co distribution along the GB, it is possible to modify not only total conductivity but also the nature of the main transport mechanism.
4.3. Conclusions

This chapter summarizes the effect of the Co addition in praseodymium doped cerias: (1) as sintering aid that increases the grain size; (2) as reduction promoter, increasing the proportion Pr$^{3+}$ versus Pr$^{4+}$ at high $pO_2$ and (3) as promoter of the total conductivity, most likely by enhancing the electronic transport.

The variation of the sintering temperature allowed the structural and transport properties of Co-doped Ce$_{0.9}$Pr$_{0.1}$O$_{2-\delta}$ to be optimized. Specifically, important differences in fluorite grain size, grain boundary nature and composition as well as the oxygen release capacity have been found depending on the sintering temperature. The sample sintered at 1000 ºC showed the highest oxygen release upon thermal treatment in argon and this is ascribed to the very high dispersion on the grain boundary and the nanoscale of cobalt-rich particles. Furthermore, the sample sintered at 1000 ºC exhibited the highest total conductivity at high temperatures, which is directly related to the enhancement of the electronic conductivity. In summary, the mixed ionic electronic conductivity of the nanocomposite Co-doped Ce$_{0.9}$Pr$_{0.1}$O$_{2-\delta}$ could be optimized by engineering the microstructure and grain boundary.

4.4. References


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Chapter 5. Study of the bulk transport properties and oxygen surface exchange of the mixed ionic electronic conductor

\[ \text{Ce}_{1-x}\text{Tb}_x\text{O}_{2-\delta} + \text{Co} \ (x=0.1, 0.2, 0.3, 0.5) \]

Abstract

Structural, bulk ionic and electronic transport properties and surface operational of Tb-doped ceria have been evaluated as a function of Tb concentration, aiming to assess potential use of the materials as high-temperature oxygen-transport membrane. The materials were synthesized by the co-precipitation method. Cobalt oxide (2 mol%) was added in order to improve sinterability and conductivity. XRD measurements suggest that a part of the cobalt is incorporated in the ceria lattice. \( \text{Ce}_{1-x}\text{Tb}_x\text{O}_{2-\delta} \) materials showed predominantly ionic conductivity, but the mixed ionic electronic conductivity could be tuned by modifying Tb (and Co-doping) concentration and temperature. Low Tb-content materials (\( x =0.1 \) and 0.2) are predominant ionic conductors, but the materials with 50 mol\% Tb show both \( p \)-type electronic and ionic conductivity. The enhanced electronic conduction in \( \text{Ce}_{0.5}\text{Tb}_{0.5}\text{O}_{2-\delta} \) is associated with narrowing of the band gap upon doping ceria with Tb. The ambipolar conductivity was determined by galvanic method coupled with gas permeation. These results support the applicability of these materials as oxygen transport
membranes at high temperature. \( \text{Ce}_{1-x} \text{Tb}_x \text{O}_{2.4}^{+} \text{Co 2mol}\% \) membranes are \( \text{CO}_2 \) stable and yield oxygen fluxes that can compete with reported perovskite materials.

In addition, the surface chemistry of the samples was investigated by means of X-ray photoelectron spectroscopy (XPS) and pulse isotopic exchange (PIE). The surface exchange rate was found to increase by increasing the level of Tb doping. The highest surface exchange rates in this study were found for materials doped with 50 mol\% Tb.
5.1. Introduction

As it was studied in chapter 3 the redox properties of Pr and Tb change depending on the temperature and oxygen partial pressures. The possibility of electron-hole hopping between dopant cations allows extending the mixed ionic-electronic conduction to higher oxygen partial pressures ($10^{-5}$-$10^{-1}$ atm), which enables its application as oxygen permeable membrane. [1, 2]

Basic requirements for oxygen permeation are fast kinetics for bulk oxygen transport, e.g., via oxygen defect sites, and electron and/or electron hole transport, as well as fast surface oxygen exchange kinetics, involving dissociation of molecular O$_2$, its reduction to oxygen adatoms, and successive incorporation in the oxide lattice as O$^{2-}$ anions. These features have been exhaustively studied in the present chapter, where different compositions based on undoped and cobalt-doped Ce$_{1-x}$Tb$_x$O$_{2-\delta}$ ($x=0.1, 0.2, 0.3, 0.5$) have been prepared and characterized to assess its applicability as an oxygen separation membrane. The influence of the dopant concentration is also examined.

Several bulk-related properties were investigated to gain insight into the transport properties of terbium-doped ceria. These were investigated by means of X-ray diffraction (XRD), temperature programmed desorption and reduction (TPD and TPR), thermogravimetry (TG), DC-conductivity in high $pO_2$ range and UV-vis spectrophotometry, which enabled to gain insight into the conduction mechanisms dependent on $pO_2$ and temperature. The surface chemistry of the samples was studied by X-ray photoelectron spectroscopy (XPS). Pulse isotopic exchange (PIE) experiments were conducted to measure
the rate of oxygen surface exchange. Subsequently, CO$_2$ stability, oxygen permeation and ion pumping experiments were done.

### 5.2. Results and discussion

**Structural characterization**

XRD patterns of Ce$_{1-x}$Tb$_x$O$_{2.5}$ (CTx, x in percentage) and Ce$_{1-x}$Tb$_x$O$_{2.5}$+ Co 2 mol% (CTxCo) sintered at 1450 ºC are shown in Figure 5.1. All the powders are single phase with cubic fluorite structure. Within experimental error, diffraction peaks corresponding to any precursor or secondary phase related to Tb or Co oxides could not be detected. The obtained patterns indicate that terbium concentrations remain below the solubility limit of Tb in ceria compounds forming a solid solution within the lattice of ceria. [3, 4] However, reported studies on the microstructure of (Co free) Tb-doped ceria showed the presence of nano-sized precipitates in the fluorite-structured matrix, which are Tb$^{3+}$ and Ce$^{3+}$ rich. [5] As previously reported [6] and observed in Chapters 3 and 4, cobalt enables lowering the densification temperature and enhances the sintering. SEM pictures of CTx and CTxCo membranes calcined at 1200 ºC showed in Figure 5.2 show the grain size growth difference caused by the addition of Co. [7] Besides, dark CoO$_x$ aggregates are observed, similar to that observed for Pr-doped ceria and other reported doped cerias, as Gd-doped. [8-10]
Figure 5.1. XRD patterns of CTx and CTxCo sintered at 1450 °C recorded at room temperature.

Figure 5.2. Scanning microscope pictures taken by backscattered electrons of CTx and CTxCo membrane cross sections, at 1200 °C.
Inside the solubility limits of a solid solution the lattice parameter of a fluorite crystal follows the Vegard’s rule, i.e., a linear relationship exists at constant temperature between crystal lattice parameter and the concentration of the solute or dopant. [4, 11] For the ceria particular case the general relation is:

\[
a (Å) = 5.4113 + \frac{4}{\sqrt{3}}[r_M - 1.024] \cdot x
\]  
(Eq. 5.1)

where \(r_M\) is the ionic radius of the dopant cation, \(x\) is the dopant amount in \(\text{Ce}_{1-x}\text{Tb}_x\text{O}_{2-δ}\), and the constant \(a\)-intercept belongs to the non doped ceria cell parameter.

![Figure 5.3 Cell parameter dependence on the dopant amount for CTx and CTxCo and theoretical Vegard’s slope for Tb\(^{3+}\) and Tb\(^{4+}\) total substitution.](image)

The cell parameters of CTx and CTxCo depicted in Figure 5.3 as a function of the concentration of Tb \((x=0, 0.1, 0.2, 0.3, 0.5)\) are calculated from the XRD patterns measured at room temperature of samples sintered at 1450 °C. The slope of this line is known as Vegard’s slope. Also the theoretical Vegard’s slope for Tb\(^{3+}\) and Tb\(^{4+}\) total substitution is drawn following Eq. 5.1. The mentioned general equation is recalculated taking into account the experimental
cell parameters for Tb undoped ceria \( a_{\text{CeO}} = 5.411 \text{ Å} \) and \( a_{\text{CeO} \cdot \text{Co}} = 5.408 \text{ Å} \), which underwent the same treatment and came from the same raw precursors than doped specimens. They give a corrector factor from the perfect fluorite \( a = 5.427 \text{ Å} \) of 0.9971 and 0.9965 for CTx and CTxCo, respectively. [3, 11]

\[
a(x_{\text{Tb}}, r_M) = 5.411 + 2.293 \cdot (r_M - 1.024) \cdot x_{\text{Tb}} \quad \text{(Eq. 5.2)}
\]

\[
a(x_{\text{Tb}}, r_M) = 5.408 + 2.292 \cdot (r_M - 1.024) \cdot x_{\text{Tb}} \quad \text{(Eq. 5.3)}
\]

Where the ionic radii are \( r_M(Tb^{4+}) = 0.88 \text{ Å} \) and \( r_M(Tb^{3+}) = 1.04 \text{ Å} \) for the cations in eight-fold coordination according to Shannon. [12] The reduction of Ce cations has been ruled out in the calculations since the non-stoichiometry of ceria and doped ceria materials has been exhaustively studied demonstrating that Ce\(^{4+}\) reduction initiates at oxygen partial pressures lower than \( 10^{10} \text{ atm} \) and keeps nearly constant at higher oxygen contents. [13-16] Consequently, in the present study where \( pO_2 > 10^{-5} \text{ atm} \), the changes observed in oxygen non-stoichiometry are most likely related to the reduction of Tb\(^{4+}\)/Tb\(^{3+}\) rather than Ce\(^{4+}\).

For each series, Figure 5.3 shows the experimental cell parameters obtained in oxidizing atmospheres and at room temperature. They follow a nearly linear trend with the Tb concentration that lay inside the defined theoretical limits. Deviations from the Vegard’s slope frequently indicate non-solubility of the dopant. [3, 11, 17] However, single phase fluorite without impurities was observed for all the samples denoting that the experimentally observed deviations rather indicate that Tb is in a mixed valence. Considering the theoretical and experimental slopes for Co-free (solid circles) and Co-containing compounds (open circles), a rough estimation of the Tb\(^{4+}\)/Tb proportion at room temperature can be done, i.e., 38% (Co-free) and 48% (Co-containing). The first value agrees with that obtained by
Chatzichristodoulou et al. for CT20 measured by XANES. [16] The lattice parameter decreases as Tb concentration rises and the calculated ratio Tb$^{4+}$/Tb remains constant within the studied Tb concentration range.

On the other hand, the cell parameters of Co-containing compounds are slightly shorter than that of Co-free samples. This can be attributed to the partial incorporation of Co cations into the ceria lattice while the rest should be in the grain boundary. Specifically, incorporation of 1.05 mol% of Co$^{2+}$ would give rise to the cell parameter change observed in CeO$_2$. Moreover, the Co addition causes the increase in the Vegard’s slope, i.e., it produces a slight decrease in the cell parameter, when compared to Co-free samples. This is an indication of the higher average oxidation state of Tb cations in the Co-containing compounds previously calculated. It should be pointed out that such a shrinkage in cell parameter cannot be explained by a higher amount of incorporated Co in the lattice, since almost twice the Co nominally available (Co$^{2+}$ in eight-fold coordination is 0.90 Å) would be needed. Otherwise, it is well-known that cobalt cations/species have high redox activity and this enables to increase the reduction and re-oxidation rate of bulk material. [18] The improved reduction activity of cobalt on the overall material has been studied by temperature programmed oxygen desorption (TPD) in argon and temperature programmed reduction (TPR) in hydrogen. TPR measurements in Figure 5.4 show that introduction of the Tb dopant decreases the reduction temperature of Ce$^{4+}$ to Ce$^{3+}$. The addition of cobalt does not cause any change in the Ce reduction peak but it lowers the reduction temperature of Tb$^{4+}$ to Tb$^{3+}$, and causes higher overall reducibility of the material, i.e., a greater consumption of H$_2$ due to the Co extra reduction peak.
Figure 5.4. a) H₂ TPR signal for CT10 and b) H₂ TPR for CTxCo (x=10 and 20).

Figure 5.5. Crystallite size dependence on the cell parameter for CT10, CT20, CT10Co and CT20Co after sintering at 1300 °C.
The evolution of the cell parameter with different compositions also agrees with the evolution of crystallite size of the samples calculated by Scherrer equation (Eq. 2.3, Chapter 2). Figure 5.5 shows that 10% Tb doped samples have a larger cell parameter and smaller crystallite sizes. This lattice expansion with decreasing grain size at room temperature is due to the larger concentration of oxygen vacancies associated to the presence of intrinsic defects of Ce$^{3+}$ and/or Tb$^{3+}$ as previously reported for other cerias. [19-21]

![Figure 5.5](image_url)

Figure 5.6. Normalized TPD measured in Ar of CTx and CTxCo. Inset: CeO$_2$ and CeO$_2$+Co. The intensity (ion current) corresponds to $m/z = 32$.

The evolution of cell parameters and associated vacancy concentration may predict the different redox processes occurring in a material dependent on temperature and $pO_2$. In the Figure 3.6 of Chapter 3, HT-XRD measurements performed on CT10 showed a linear dependency of cell parameter on the temperature and an increase of the slope at 480 °C. The first slope corresponds to the thermal expansion coefficient (ca. 12.1x10$^{-6}$ K$^{-1}$) in air while the increase in slope beyond 430 °C is the additional contribution of chemical expansion due...
to the reduction of Tb$^{4+}$ to Tb$^{3+}$ and the associated oxygen release (ca. 5.77x10$^{-6}$ K$^{-1}$).

The mentioned oxygen release can be observed in the TPD measurements depicted in Figure 5.6, where the ion current corresponding to the released oxygen from the solid oxide powder (as determined by mass spectroscopy) is plotted as a function of temperature. CT10 oxygen desorption starts at ~ 480 °C, approximately the same temperature at which the chemical expansion becomes tangible by HT-XRD. It reaches a maximum at 670 °C and decreases hereafter. This oxygen release implies the reduction of the material and the oxygen vacancies formation that, taking into account that the $pO_2 > 10^{-5}$ atm, are related to the reduction of Tb$^{4+}$/Tb$^{3+}$. Higher dopant content is found to shift the oxygen release to lower temperatures (CT20 and CT50), i.e., it increases the reducibility of the material.

Co-containing samples present two distinguished redox processes: (1) the oxygen release associated with Tb$^{4+}$ reduction, which starts at lower temperatures than their analogous Co-free samples; and (2) a new important well-defined reduction process that takes place at temperatures ranging from 730 to 880 °C and that reaches a maximum at 780 °C, 800 °C and 810 °C for CT10Co and CT20Co, CT50Co, respectively. This high-temperature reduction process can be ascribed to two different processes:

- reduction of Co itself (Co$^{3+}$→Co$^{2+}$), which is confirmed by the existence of this reduction in in CeO$_{2.4}$ +Co sample, while it is not present in CeO$_{2.4}$ as shown in the inset of Figure 5.6. [18] Note that the intensity of the Co-related peak changes due to the Tb background.

- further reduction of bulk Tb$^{4+}$ to Tb$^{3+}$ is confirmed by the existence of a broadened peak at higher temperatures.
In summary, TPD measurements reveal that addition of either Co or a higher amount of Tb increases the oxygen release \textit{i.e.}, the material reducibility.

Figure 5.7. (a) Oxygen non-stoichiometry (\(\delta\)) determined by TG in air and (b) \(\delta\) determined by TG in N\(_2\), for CTx and CTxCo. Temperature range include from 430 to 1000 ºC.

Figure 5.7 shows the evolution of the oxygen non-stoichiometry parameter \(\delta\) in air (a) and in N\(_2\) (b) with temperature for the different samples, as determined by thermogravimetry. The estimations were done by taking into account that the mass loss corresponds to oxygen release (after a drying step). To obtain the final non-stoichiometry, the total \(O_2\) mol per mol of fluorite has been added to the oxygen of the totally reduced fluorite in N\(_2\), following the equation:

\[
2 - \delta = \frac{n_{O_2}}{n_{fluorite}} + (2 - \frac{x}{2})
\]

(Eq. 5.4)

As expected, at high temperatures \(\delta\) is not affected by the Co addition while only minor changes in \(\delta\) are observed at low temperatures. On the other hand, a substantial change in \(\delta\) arising from the Co addition is observed for the
sample with 50% Tb (CT50Co). In fact, the lower oxygen release observed in CT50Co with respect to CT50 seems to be related to the higher concentration of Tb$^{3+}$ at lower temperatures, which also emerged from the cell parameter data of XRD at room temperature (Figure 5.3).

**Electrochemical characterization**

In order to analyze the complete conductivity mechanism in the studied range of temperatures and $pO_2$, it would be interesting to explore the defect chemistry of these compounds. The dominant defect equilibrium at high $pO_2$ in the system CTx is thereby suggested to be that of Tb reduction. Stefanik and Tuller [22] and Fagg et al [17] described a defect model including a variable valency acceptor dopant. The four regions plotted in Figure 5.8 delimit the main key defect reactions for a general fluorite. In this study, the considered limits are regions II and IV, which are listed below for the particular case of Tb doped ceria, starting with the consideration of pure CeO$_2$.

![Figure 5.8. Kröger-Vink diagram for acceptor doped fluorite. Four $pO_2$ regions delimit the main key defect reactions for a general fluorite.](image)
First, the formation of oxygen vacancies and interstitials is dominated by the Frenkel defect mechanism for pure CeO$_2$ as described in Eq. 5.5, while intrinsic electron-hole pair generation produced by thermal excitation over the band gap is governed by Eq. 5.6. On the other hand, molecular oxygen release causes oxygen vacancy generation (charge compensated by the formation of free electrons localized on Ce ions) and hopping conduction will occur over the Ce$^{3+}$/Ce$^{4+}$ from intrinsic defects of pure ceria sublattice following Eq. 5.7.

\[ O_0^x \rightarrow V_0^- + O_i^{x''} ; \quad [V_0^-] [O_i^{x''}] = K_F(T) \quad (Eq. 5.5) \]

\[ \text{nil} = e^+ + h^- ; \quad np = K_e(T) \quad (Eq. 5.6) \]

\[ O_0^x \rightarrow \frac{1}{2} O_2 (g) + V_0^- + 2e^- ; \quad [V_0^-] n^2 pO_2^{1/2} = K_R(T) \quad (Eq. 5.7) \]

In a Tb doped system it has to be taken into account the mass conservation law Eq. 5.8, the ionization of Tb cations Eq. 5.9, and the added charge defect $Tb_{ce}'$ in order to keep the electroneutrality condition Eq. 5.10.

\[ [Tb_{ce}'] + [Tb_{ce}^x] = Tb_{total} \quad (Eq. 5.8) \]

\[ Tb_{ce}^x \rightarrow Tb_{ce}^x + e^- ; \quad \frac{[n_{ce}^x]}{[Tb_{ce}]} = K_{Tb}(T) \quad (Eq. 5.9) \]

\[ 2[O_i^{x''}] + n + [Tb_{ce}'] = 2[V_0^-] + p \quad (Eq. 5.10) \]

Considering the region IV boundary, since Tb introduces extrinsic oxygen vacancies, the concentration of interstitial oxygen will be suppressed $[O_i^{x''}] \rightarrow 0$, improving the ionic conductivity in the high $pO_2$ range.

Besides, in oxidizing conditions and low temperatures (below 480 °C) most terbium exists as Tb$^{5+}$ and then $Tb_{ce}^x \ll Tb_{ce}$ (n << $K_{Tb}$). Therefore:
Study of the bulk and surface transport properties of Ce$_{1-x}$Tb$_x$O$_{2-d}$ + Co

\[ n \propto pO_2^{-1/6} \quad \text{(Eq. 5.11)} \]

\[ [V^\cdot_0] \propto pO_2^{-1/6} \quad \text{(Eq. 5.12)} \]

\[ p \propto pO_2^{1/6} \quad \text{(Eq. 5.13)} \]

In the case of region II, when \( Tb'_\text{Ce} \gg Tb^x_\text{Ce} \) (\( n \gg K_{\text{Tb}} \)) but \( T_{\text{b total}} \gg n \); consequently:

\[ n \propto pO_2^{-1/4} \quad \text{(Eq. 5.14)} \]

\[ [V^\cdot_0] \propto pO_2^0 \quad \text{(Eq. 5.15)} \]

\[ p \propto pO_2^{-1/4} \quad \text{(Eq. 5.16)} \]

Thus, vacancy concentration is independent on \( pO_2 \) and proportional to the total terbium concentration.

On the other hand, XRD analysis suggests that the Co added in CTxCo samples is partially incorporated in the fluorite lattice as Co$^{2+}$, giving rise to one oxygen vacancy per substituted Ce. [13] Therefore, the electroneutrality balance (Eq. 5.10) must be modified with the addition of \( [Co^\cdot_{\text{Ce}}] \) in the left term. The rest of Co is likely placed along the grain boundaries and does not affect bulk defect chemistry in equilibrium.

Hence, equations above describe a way to make an evaluation of the predominant type of carriers from the conductivity behaviour with temperature and \( pO_2 \). Primarily, the influence of the temperature is considered.

Figure 5.9 shows the conductivity \( \ln(\sigma T) \) as a function of the reverse temperature for CTx (a) and CTxCo (b) measured in \( pO_2 \) of 0.21 and
5x10⁻⁵ atm. Given a sample composition, differences in conductivity at different $pO_2$ indicate a change in the level of reduction. Moreover, oxygen can be more easily incorporated at high $pO_2$ as temperature drops while it cannot be incorporated (reoxidization) at low $pO_2$. This causes an apparent change in activation energy ($E_a$) under oxidizing atmospheres ($pO_2 = 1$ and 0.21 atm), in CT10, CT10Co, CT20 and CT20Co, while reoxidization cannot be observed at $pO_2$ of 5x10⁻⁵ atm in the measured temperature range (800-400 °C). The variations in $E_a$ at given conditions indicate a temperature dependent change in the main conduction process contributing to the total conductivity. Therefore, changes in the $E_a$ in this kind of mixed conducting materials may denote variations in the proportion between ionic and electronic contributions. Taking into account that at these high $pO_2$ doped ceria materials have been reported as pure ionic conductors, the change in apparent $E_a$ could be analyzed in terms of vacancy concentration increase due to the release of oxygen (Tb reduction). Actually, $E_a$ change of CT10 in air is produced at 470-480 °C, which matches the temperature range observed by XRD for oxygen desorption/incorporation and TPD (Figure 5.6).

On the other hand, the conductivity of CT50Co and specially CT50 is higher than for the cerias with lower Tb concentration. In this case, the change in $E_a$ occurs in the low oxygen content environment and at higher temperature (ca. 680 °C). At this temperature, TPD in Ar shows that the majority of Tb is already reduced. Accordingly, the increase in the ionic conduction that causes the $E_a$ change may be attributed to oxygen vacancies that form associates at lower temperatures.
Figure 5.9. Arrhenius plot of the total conductivity in air and in Ar (0.21 and $5 \cdot 10^{-5}$ atm, respectively) for a) CTx and b) CTxCo.

Table 5.1 summarizes apparent $E_a$ in the temperature ranges of 800-480 °C (high) and 480-400 °C (low) for 10 and 20% Tb doped ceria and 800-700 °C (high) and 700-400 °C (low) for 50% Tb. Experimental $E_a$ matches the typical values found for ionic conductors. As mentioned, at $pO_2 < 10^{-3}$ atm, there exists only one $E_a$ for CT10, CT10Co, CT20 and CT20Co, i.e., terbium is mainly reduced to Tb$^{3+}$, over the whole studied temperature range, thus behaving as a fixed valence dopant, which enhances principally the ionic conductivity. CT50 and CT50Co constant $E_a$ appears at $pO_2 > 10^{-3}$. 
Table 5.1. Apparent activation energies in the temperature ranges of 800-480 °C (high T) and 480-400 °C (low T) for CT10, CT10Co, CT20 and CT20Co and in the temperature range of 800-700 °C (high T) and 700-400 °C (low T) for CT50 and CT50Co. The conductivities at 800 °C under air and 5x10⁻⁵ atm is indicated.

<table>
<thead>
<tr>
<th></th>
<th>$E_a$ (eV)</th>
<th>$\sigma$ (S/cm) 800 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.21 atm High T</td>
<td>5x10⁻⁵ atm High T</td>
</tr>
<tr>
<td>CT10</td>
<td>1.08</td>
<td>0.80</td>
</tr>
<tr>
<td>CT10Co</td>
<td>1.06</td>
<td>0.80</td>
</tr>
<tr>
<td>CT20</td>
<td>0.87</td>
<td>0.71</td>
</tr>
<tr>
<td>CT20Co</td>
<td>0.82</td>
<td>0.77</td>
</tr>
<tr>
<td>CT50</td>
<td>0.51</td>
<td>0.78</td>
</tr>
<tr>
<td>CT50Co</td>
<td>0.51</td>
<td>0.83</td>
</tr>
</tbody>
</table>

Total conductivity at 800 °C under 0.21 and 5x10⁻⁵ atm $pO_2$ is also specified in Table 5.1. The increase of Tb content and the addition of cobalt enables increasing substantially the total conductivity and the best conductivity values have been obtained for 50% Tb content. Figure 5.10a depicts the dependency of total conductivity on Tb dopant concentration under air. The total conductivity in air increases with the Tb dopant concentration, which is interpreted as a superior concentration of oxygen vacancies at higher Tb doping levels having into account that only the dopant is reduced ($\text{Tb}^{4+} \rightarrow \text{Tb}^{3+}$). [14] When Co is added to both compositions, the reduction of the sample is enhanced as shown by TPD (Figure 5.6) and TPR measurements (Figure 5.4). This allows increasing the oxygen vacancy concentration at the same temperature with respect to the Co-free samples. Thus, the ionic conductivity increases and electronic conductivity is promoted as confirmed later on by oxygen permeation results.
Figure 5.10. Total conductivity dependency on Tb dopant concentration in (a) air, $pO_2 = 0.21$ atm and (b) Ar, $pO_2 = 2 \cdot 10^{-5}$ atm.

Figure 5.10b shows the dependency of the total conductivity on Tb dopant concentration under Ar ($pO_2 = 2 \cdot 10^{-5}$ atm). The total conductivity in Ar reaches a maximum at about $x = 30$ mol% Tb. For CT50 and CT50Co, the total conductivity is found to decline with decreasing $pO_2$. This observation can be attributed to a mainly $p$-type electronic conductor behavior. In this case, the conductivity diminishes with the decrease in $pO_2$ since the formation of oxygen vacancies diminishes the electron-hole density. Besides, defect association between dopant Tb$^{3+}$ cations and oxygen vacancies can occur, according to Eq. 5.15,

$$Tb^{3+}_Ce + V_0^- \leftrightarrow (Tb^{3+}_CeV_0^-)$$  \hspace{1cm} (Eq. 5.17)

This new type of defect occurs at high concentrations of oxygen vacancies due to the electrostatic interaction among dopant Tb$^{3+}$ and the oppositely charged oxygen vacancy. These associated defects prevent oxygen vacancies to move and, consequently, the ionic conductivity diminishes. [15, 23]
Figure 5.11. Conductivity of CTx and CTxCo systems as a function of $pO_2$ measured at (a) 400 °C and (b) 800 °C.

Figure 5.11 plots the conductivity logarithm at 400 °C (a) and 800 °C (b) versus $pO_2$ logarithm for the different compositions. The plots show that (1) in general, the total conductivity is enhanced by cobalt addition, and (2) the conductivity-$pO_2$ behavior depends on the Tb content.

At 400 °C, over the whole range of $pO_2$, CT10 shows $pO_2^{-1/6}$ dependency, which could be characteristic of both $n$-type electronic conductivity and ionic conductivity via oxygen vacancies for this material and conditions, as described by Eq. 5.11 and Eq. 5.12, respectively. At 800 °C the $pO_2^{-1/6}$ dependency exists but there is a conductivity plateau at $pO_2 < 10^{-3}$ atm, characteristic of ionic transport as illustrated in Eq. 5.15. This plateau is related to a maximum reduction of Tb reachable at these temperatures in this $pO_2$ range that causes the material to achieve a steady vacancy concentration independently on the $pO_2$ and proportional to the oxygen vacancy concentration ($\sigma \propto [V_0^{\cdot\cdot}]$). The absence of a continuous slope of -1/6 indicates that, even at $pO_2$ higher than $10^{-3}$ atm, the total conductivity should be predominantly ionic because $n$-type electronic conductivity would result in a continuous increase.
when $pO_2$ decreases, which is not observed in the experimental results. Hence, conductivity is mainly ionic in these oxidizing conditions and the multiple oxidation state of the dopant makes the ionic conductivity $pO_2$ dependent (region not observed in single-oxidation-state acceptor doped systems as Gd).

The previous behaviour can be extended to CT10Co even though the increase in conductivity is not only due to the oxygen vacancies but also to enhanced densification of the sample. It has been reported that the use of Co as a sintering aid may result in the decrease of grain boundary resistance. [13, 24] By XRD the maximum solubility of the Co into the ceria lattice was determined to be 1.05 mol%, which increases the $p$-type contribution shifting the transition to the ionic plateau to higher $pO_2$. The remaining Co appears as a cobalt-rich phase at the grain boundary (observed by SEM but below the detection limit of the XRD device). This cobalt-rich phase might act as a percolating network, increasing the electronic conductivity as observed for other doped cerias. [24] A beneficial effect is that the catalytic activity for the oxygen reduction reaction at the surface may change by the presence of Co, which will be discussed below. As a result, the $pO_2$ slope is smoothed and an extended plateau range is observed.

As aforementioned, CT20 introduces more oxygen vacancies than the 10% doped material broadening the electrolytic region over that of CT10. The $pO_2$ dependency of the conductivity for CT20 is lower than -1/6 at both temperatures, and reaches the ionic plateau at higher $pO_2$ than for CT10. $pO_2$ dependency smaller than -1/6 sets the behaviour of the material in the diffuse region between the vacancy concentration $pO_2$ independent and the -1/4 dependence (boundary between regions II and III). This shift to region III with respect to CT10 may be the consequence of the introduction of energetic levels of Tb into the ceria band gap, which permits electron-holes to hop from Tb$^{3+}$ to
Tb\(^{4+}\) and the concurrent enhancement of the electronic conductivity. The small addition of Co oxide allows further increasing of the \(p\)-type electronic conductivity. The slight increase in conductivity observed for these samples with decreasing \(pO_2\) is still attributed to Tb\(^{4+}\) reduction with concomitant formation of oxygen vacancies.

The total conductivity of CT50 is found proportional to \(pO_2^{1/6}\), which is ascribed to predominant \(p\)-type conductivity. For highly Tb-doped ceria, the impurity band contributing to the electron-hole hopping and causing additional electronic transport is more evident. This small polaron hopping requires the mixed valent Tb, since electrons on Tb\(^{3+}\) sites can only hop to an adjacent Tb site if it is empty, \(\text{e.g.}\) in the Tb\(^{4+}\) state. The maximum contribution to the electronic conductivity is, therefore, \(pO_2\) dependent, and it will be reached when the concentration of the two valence states are equal. [22] These results confirm that transport of electron-holes is the dominant conduction mechanism at high \(pO_2\) for CT50. Finally, Co addition to CT50 provides the highest conductivity values obtained in this study. However, the slightly positive slope observed at 400 °C suggests that the enhancement of vacancy formation by Co is such that causes defect interaction typical for highly doped specimens, \((\text{Tb}^{4+}_{\text{Ce}}V_0)\). [15, 22] This effect is more appreciable at low temperatures since there is not enough energy to exceed the defect dissociation barrier and oxygen vacancies are not free and able to account on the ionic transport process.

On the other hand, UV-vis absorption experiments were carried out in order to study the influence of the dopant on the band gap and thus, the \(p\)-type electronic conductivity. The band gap measures the energy difference between the top of the valence band occupied with electrons and the bottom of the conduction band unfilled of electrons, thus being related to the extent of electronic conductivity of the materials. In fact, the optical band gap energy is
 comparable to the thermal band gap related to the formation of electron-hole pairs. Kubelka-Munk (K-M) theory has been used to estimate the forbidden energy gap $E_g$ of the material. [25, 26] This consists of plotting the obtained reflectance $h\nu F(R\infty)^2$ as a function of the photon energy, $h\nu$. The band gap $E_g$ can be obtained by extrapolating to zero a tangent line drawn in the point of inflection of the curve, i.e., the point of intersection with the $h\nu$ horizontal axis.

![Figure 5.12](image)

**Figure 5.12.** UV-vis absorption spectra $(h\nu F(R))^2$ versus photon energy of (a) CTx and (b) CTxCo.

Figure 5.12 shows the room temperature UV-vis absorption spectra of the Tb-doped series. CeO$_2$ and Tb$_4$O$_7$ were taken as reference (see inset in Figure 5.12a). The energy gaps for CeO$_2$ and Tb$_4$O$_7$ are found to be 3.3 eV and 1.94 eV, respectively. The band gap energy of all doped samples decreases from the pure ceria to be close to that of Tb$_4$O$_7$. This fact suggests that Tb doping makes it easier electron hop and increases electronic-hole conductivity. For low Tb content materials ($x=0.1, 0.2$) the addition of Co results in even smaller values for $E_g$. 
Surface characterization

The overall oxygen transport through a dense oxide is governed not only by the electron (or hole) and oxygen ion in the bulk, but also by the surface exchange reaction at the device interfaces. Near-surface defect chemistry plays an important role in determining the chemical surface chemistry and gas exchange activity. Therefore, X-ray photoelectron spectroscopy (XPS) has been performed to gain insight into both (1) valence bands and (2) core levels of the atoms present in the first layer of the material surface. Samples of CT20 and CT20Co were quenched from 800 °C under air stream to liquid nitrogen in order to have the similar material state as for the conductivity and PIE measurements.

![Figure 5.13. a) Ce 3d XPS spectra of CT20 and CT20Co quenched in air from 800 °C and b) Ce3d XPS separation of peaks of CT20 and CT20Co spectra. Pass energy was 30 eV.](image)

The Ce 3d XPS spectra of CT20 and CT20Co quenched in air from 800 °C are plotted in Figure 5.13. Both of them show three peaks, separated by several eV for each of the spin–orbit split 5/2;3/2 components (Figure 5.13b). Thus, the 3d_3/2 and 3d_5/2 contributions are represented as u and v, respectively. The u''
component at 915.5 eV, which is a fingerprint of the Ce$^{4+}$ state, presents a considerable intensity and does not overlap with other peaks. This peak is coupled with $v''$ (897.2 eV) and they both come up from an almost pure 4$f^0$ electron configuration. [27] Besides, good separation between $v$ (881.3 eV) and $v''$ (888.1 eV) peaks can be observed, which is a characteristic of only Ce$^{4+}$ state. [28] The overall spectrum appears around 1 eV lower than positions found in the literature for Ce3d, ascribed to charge shifting of the device. There are slight differences between peak separation from CT20 to CT20Co, which are attributed to accumulated experimental errors rather than a mixture of oxidation states. Ce$^{3+}$ peaks represented as $v_0$, $u_0$ should be expected in these spectra at around 879.6 and 898.3 eV, respectively (marked by dotted vertical lines). [29] Here, it can be seen that the latter peaks do not appear, which confirms that there is a negligible amount of Ce$^{3+}$ in the samples. Furthermore, no meaningful changes in the cerium oxidation state have been found due to the influence of cobalt for the studied conditions.

Figure 5.14. XPS of the Tb 3$d$ region of a) CT20 and b) CT20Co, quenched in air from 800 °C, acquired with a pass energy of 30 eV.
The most intense core level for terbium is Tb 3d at the high binding energy between 1230 and 1290 eV. However, it is difficult to distinguish accurately the chemical states of Tb 3d line as binding energies of the different oxidation states are very similar. Figure 5.14 shows the Tb 3d XPS patterns of quenched samples. Two principal components are commonly distinguished related to Tb 3d_{3/2} and the Tb 3d_{5/2} binding energy peak position. The binding energy peak position is shifted from that earlier published for pure TbO_{1.5} (Tb^{3+}) where the Tb 3d_{5/2} and Tb 3d_{3/2} peak are reported at about 1239.1 and 1274 eV. Tb^{4+} related peaks should appear at 1244 and 1278 eV. [30] In addition to the main peaks, terbium shows relatively intense satellites in the Tb 3d_{5/2} component at 1251.5 eV for the tetravalent state and 1250.6 eV for the trivalent state, and at 1264 eV for the Tb 3d_{3/2} component of Tb^{4+}. [31] The spectra in Figure 5.14 show two peaks at 1241.6 and 1276.3 eV, and 1240.9 and 1275.7 eV for CT20Co and CT20, respectively. Also two satellites appear between the main peaks. These results suggest that the Tb introduced in the position of Ce in the CeO_{2} lattice as an extrinsic dopant appears in more than one oxidation state i.e., 3+ and 4+. [32] The CT20Co spectrum is more shifted to higher energies, which indicates a higher oxidation state, i.e. higher amount of Tb^{4+}. From the deconvolution of the spectra it is possible to estimate 60.5% and 54% of Tb^{3+} in the surface of CT20 and CT20Co respectively. The surface oxidation state of Tb differs from the bulk oxidation state estimated by TG measurements in air (800 ºC), ca. 88.5% for CT20 and 81% for CT20Co. Nevertheless, XPS results support the presence of Tb in both 3+ and 4+ oxidation states inferred from lattice parameter extracted from XRD of CT20 and CT20Co.
The presence of Co atoms on the grain surface is confirmed by the XPS. The binding energy (BE) values of the Co $2p_{3/2}$ core-level are close to 780 eV for Co$^{2+}$ on CoO. The main peak is accompanied by a relatively intense satellite peak at 787.5 eV characteristics of Co$^{2+}$ species and its interaction with other metal. [33] Spin–orbital splitting upon ionization between $2p_{3/2}$ and $2p_{1/2}$ is 16 eV. XPS spectra of the synthesized CT20Co material in the Co 2p region are shown in Figure 5.15. The low intensity of the Co 2p region is due to the very low Co content (2 mol%). The BE of Co $2p_{3/2}$ peak is ca. 780 eV and the satellite peak at 787.5 eV. These binding energies correspond to cobalt oxide species highly dispersed with a strong interaction with the structure, which may indicate the partial incorporation of Co in the lattice of CeO$_2$. Then, the BE is shifted to lower values (about 0.2 eV) compared to the BE of Co $2p_{1/2}$ on Co$_3$O$_4$ or CoO (780.2 eV). [34] Nevertheless, the proximity of BE for both oxide states, i.e., Co$_3$O$_4$ or CoO, makes it very difficult to indentify the Co state at the surface of these materials.
The oxygen exchange kinetics in the CTx and CTxCo surface has been studied by pulse response $^{18}O,^{16}O$ isotope exchange (PIE) measurements. Figure 5.16 displays Arrhenius plots of the overall exchange rate ($\mathcal{R}_0$) for all materials investigated in this study as evaluated from data of PIE measurements. $\mathcal{R}_0$ is found to increase profoundly with increasing Tb concentration. The data listed in Table 5.2 support a relationship between the surface exchange rate, at 700 °C, and the corresponding concentration of oxygen vacancies in the Tb-doped ceria samples.

Figure 5.16. Arrhenius plot of the exchange rates of CT10, CT10Co, CT20, CT20Co, CT50, CT50Co.

Lane et al reported that the rate of oxygen surface exchange might be influenced by the surface concentration of oxygen vacancies and by the supply of electrons, which are necessary for charge transfer. [35] Note from Figure 5.16 that Co addition to either CT10 or CT50 does not significantly improve $\mathcal{R}_0$. However, the surface exchange rate exceeds that of the CT10 at temperatures above 750 °C. Above this temperature, the [$V_{O}^-$] reaches a maximum, i.e. Tb$^{3+}$ is the prevailing Tb cation, as inferred from XRD, DC conductivity and TG
experiments. Below 700 °C surface exchange appears to be limited by the \([\nu O^-]\), partly due to the annihilation of oxygen vacancy associated with the partial oxidation of Tb from 3+ to 4+. However, when the Tb^{4+} starts to reduce at higher temperatures, \(\mathcal{R}_0\) is improved through Co addition, which slightly boosts \(p\)-type conductivity.

CT20Co increases the surface exchange rate of oxygen with respect to the Co-free counterpart at any tested temperature. This behavior suggests that for CT20, \([\nu O^-]\) was enough to ensure the reaction and that the rate limiting step is the electronic contribution. As a consequence, the increase in \(p\)-type conductivity by Co-addition alleviates the electronic limitation and raises the overall surface exchange rate. Table 5.2 shows the apparent \(\mathcal{R}_0\) activation energy obtained for all six compounds. The \(E_a\) observed for CT20Co decreases with respect to CT20, suggesting a possible change in the rate limiting step, presumably from limitation by insufficient electronic supply (CT20, 118 kJ/mol) to limitation by deficient \([\nu O^-]\) (CT20Co, 101 kJ/mol). Indeed, the \(E_a\), seemingly ascribed to \([\nu O^-]\) limitation coincides with the \(E_a\) observed for CT10 and CT10Co.

![Figure 5.17. Surface exchange rate as a function of the oxygen nonstoichiometry at 700 °C.](image)
Table 5.2: Oxygen non-stoichiometry and surface exchange rate, at 700 ºC, and apparent activation energies for the surface exchange rate of the different compounds.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\delta_{700\degree C}$</th>
<th>$\mathcal{R}<em>0$ (mol O·m$^{-2}$·s$^{-1}$)$</em>{700\degree C}$</th>
<th>$E_a$ (kJ/mol) [temperature range]</th>
</tr>
</thead>
<tbody>
<tr>
<td>CT10</td>
<td>0.042</td>
<td>$6.71 \times 10^{-5}$</td>
<td>105 [600 – 900 ºC]</td>
</tr>
<tr>
<td>CT10Co</td>
<td>0.036</td>
<td>$4.18 \times 10^{-5}$</td>
<td>136 [700 – 900 ºC]</td>
</tr>
<tr>
<td>CT20</td>
<td>0.08</td>
<td>$2.42 \times 10^{-4}$</td>
<td>118 [600 – 900 ºC]</td>
</tr>
<tr>
<td>CT20Co</td>
<td>0.073</td>
<td>$4.02 \times 10^{-4}$</td>
<td>101 [500 – 900 ºC]</td>
</tr>
<tr>
<td>CT50</td>
<td>0.179</td>
<td>$1.96 \times 10^{-3}$</td>
<td>96 [600 – 700 ºC]</td>
</tr>
<tr>
<td>CT50Co</td>
<td>0.174</td>
<td>$1.52 \times 10^{-3}$</td>
<td>97 [500 – 750 ºC]</td>
</tr>
</tbody>
</table>

Similar $\mathcal{R}_0$ increase through Co addition is not observed for the case of CT50Co. $E_a$ is not altered, which may indicate that there is no change in the rate limiting step when cobalt is added. Both CT50 and CT50Co showed $p$-type electronic conductivity (Figure 5.11). Additionally, the $E_a$ is similar to that observed for the samples limited by a process related to oxygen vacancies. Therefore, 50%-doped samples seem not to be limited by electronic supply but by a process involving oxygen vacancies. Accordingly, slightly higher exchange rates are observed for CT50, which can be explained by the CT50 higher non-stoichiometry than that for CT50Co (Figure 5.17). These observations support the conclusion that the rate of exchange in these materials is determined by i) the concentration of oxygen vacancies, which is related to the ionic conductivity, ii) the $p$-type electronic conductivity, and iii) the balance between these two parameters.
Figure 5.18. Inverse temperature dependence of the rates of dissociative adsorption ($\mathcal{R}_a$), oxygen incorporation ($\mathcal{R}_i$) of the terbium doped studied series.

PIE experiments also provide information about the surface exchange mechanism. [36] The distribution of oxygen isotopomers ($^{18}\text{O}_2$, $^{16}\text{O}^{18}\text{O}$, $^{16}\text{O}_2$) in the effluent pulse can be modeled by adopting a two-step exchange mechanism, in which isotopic scrambling is determined by the rate of dissociative adsorption of $\text{O}_2$ molecules at the surface ($\mathcal{R}_a$) and that of successive incorporation of adsorbed O adatoms in the oxide lattice ($\mathcal{R}_i$). Thus, overall exchange rate is considered to occur as a series process. Figure 5.18 plots the rates of dissociative adsorption, $\mathcal{R}_a$, and oxygen incorporation, $\mathcal{R}_i$, as a function of the inverse temperature. The overall rate of surface exchange on both CTx and CTxCo samples appears to be controlled by the dissociative adsorption reaction,

$$\frac{1}{\mathcal{R}_o} = \frac{1}{\mathcal{R}_a} + \frac{1}{\mathcal{R}_i}$$
as was found for, e.g., 25 mol% erbia-stabilized bismuth oxide (BE25) [37] and Ba	extsubscript{1-x}Sr	extsubscript{x}Co	extsubscript{0.8}Fe	extsubscript{0.2}O	extsubscript{3-δ} (BSCF). [36] For all compositions studied, the rate of oxygen incorporation was found between one or two orders of magnitude higher (Figure 5.18). The observations made in this study can be reconciled by assuming that O	extsubscript{2} dissociative adsorption and, hence, overall oxygen exchange is rate determined by electron transfer to intermediate superoxide ions (O	extsubscript{2}⁻) at the surface:

\[ \text{O}_2^- + e^- \rightarrow 2\text{O}^- \]  

(Eq. 5.18)

with the notion that surface oxygen vacancies are assumed to be the sites onto which adsorption of O	extsubscript{2} molecules takes place.

Permeation measurements and stability tests

In this section, the oxygen permeation through different samples has been determined under certain pO	extsubscript{2} gradients, which may enable to assess the presence of mixed conductivity and the applicability as O	extsubscript{2} separating membranes. Oxygen permeation through CT10, CT10Co, and CT20Co membranes sintered at 1200 °C has been measured. Membranes made of CT20, CT50 and CT50Co were not measured due to manufacturability issues, i.e. no dense membrane specimens were achieved by the current sintering techniques. Figure 5.19 shows the oxygen flux as a function of temperature for two sweep gases: argon and diluted methane.
Figure 5.19. Oxygen permeation flux as a function of inverse temperature for CT10Co and CT20Co. The oxygen flux for similarly configuration BSCF and LSCF membranes is shown for comparison.

Figure 5.19 illustrates also the flux obtained with benchmark perovskite-based membranes as BSCF and LSCF in a similar configuration (thickness and sample history). CT10 membrane shows nearly negligible permeability (not shown) and this confirms the main ionic conductivity of this composition. However, 0.9 and 1.2 mm-thick membranes made of CT10Co and CT20Co present permeation values at 1000 ºC around 0.02 and 0.08 mL·min⁻¹·cm⁻² for argon sweeping, respectively.

The combination of conductivity and permeation measurements using imposed current allowed determining the ambipolar conductivity at high temperature for CT20Co. CT20Co transference numbers were obtained across a 1.2 mm thick membrane under a $pO_2$ gradient 0.21/5x10⁻⁵ atm (air/Ar). The transference numbers showed in Figure 5.20a indicate that CT20Co performs electronic conductivity although ionic conduction prevails. This study is performed in the temperature range from 750 to 1020 ºC, which is suitable for
application as oxygen transport membranes. Ionic conductivity is predominant in the whole temperature range but electronic contribution becomes more important as the temperature drops. Figure 5.20b presents the ionic, electronic and total conductivity values obtained by using the transference number. The $O_2$ flux obtained agrees perfectly with the expected value calculated considering the ambipolar conductivity, $\sigma_{amb} = \frac{\sigma_1 \sigma_e}{(\sigma_1 + \sigma_e)}$, and assuming a constant mean value in the experimental $pO_2$ gradient, e.g. at 860 °C, the obtained $J(O_2)$ is $2.8 \cdot 10^{-8}$ mol·s$^{-1}$·cm$^{-2}$, and the $\sigma_{amb}$ is 0.0085 S·cm$^{-1}$. By using Wagner’s equation (Eq. 1.26), the $J(O_2)$ is given by Eq. 5.19

$$J(O_2)_{(calculated)} = \frac{RT}{16F^2} 0.0085 ln \frac{0.21}{0.00019}$$

(Eq. 5.19)

which finally gives $j_{O2} = 3.1 \cdot 10^{-8}$ mol·s$^{-1}$·cm$^{-2}$.

Figure 5.20. a) Ionic transport number and b) total, ionic and electronic conductivity of CT20Co measured across a 1.2 mm thick membrane under a nominal gradient of $pO_2$ 0.21 || 5x10$^{-5}$ atm (air/Ar). $J(O_2)$ measured using chromatography analytical data.

In more reducing conditions the permeation can be enhanced since the driving force is augmented. For methane sweeping CT10Co sample is able to achieve 0.5 mL·min$^{-1}$·cm$^{-2}$. This flux increase is additionally attributed to the
Study of the bulk and surface transport properties of $\text{Ce}_{1-x}\text{Tb}_x\text{O}_{2-d} + \text{Co}$

substantial rise of $n$-type conductivity achieved preferentially through Ce$^{4+}$ reduction. The permeation values are also interesting when compared to some perovskite type permeation membranes, especially due to the high stability under reducing and CO$_2$ containing atmospheres at high temperatures, where broadly applied perovskite materials are prone to decompose. Specifically, these harsh operation conditions are encountered in the targeted applications in oxyfuel processes using flue gas as sweep and high-temperature catalytic membrane reactors for hydrocarbon conversion into syngas and added-value products. The stability of CT10 and CT10Co has been studied by thermogravimetry under continuous flow of air with 5% CO$_2$. Figure 5.21 shows the mass evolution of these samples and the evolution of a highly permeable perovskite, i.e., BSCF. Carbonates are formed (mass increase) in the range from 700 to 900 °C and decompose (mass decrease) at higher temperatures, as it can be observed for the BSCF measurement. Ceria sample presents a negligible formation of carbonates while it is also detected the mass loss due to oxygen release at temperatures above 480 °C.

Figure 5.21. Thermogravimetric analysis under CO$_2$ flow of CT10 and CT10Co in CO$_2$, and BSCF.
5.3. Conclusions

Tb doped ceria shows predominant ionic conductivity or mixed ionic-electronic conductivity depending on the Tb and Co doping level and temperature range, which proves MIEC properties of these compounds. The extent of mixed ionic-electronic conductivity is a function of temperature and can be tuned by modifying the Tb- (and Co-doping) concentration. Low Tb content materials (x = 0.1) show predominantly ionic conductivity. Higher doping levels (>20%) introduce new Tb electronic levels in the ceria bandgap that allow the small polaron hopping between Tb ions of different oxidation state (since Ce does not reduce under the studied pO$_2$ range). As a consequence, the electronic conductivity of these compositions in oxidizing conditions is not negligible and will be limited by the number of available sites for electron hops. CT50 and CT50Co materials show p-type electronic conductivity. XRD results showed that Co addition does affect the oxidation state of Tb, increasing the ratio Tb$^{4+}$/Tb$^{3+}$. However, it is not entirely incorporated into the ceria lattice, but in the grain boundary. Consequently, the addition of cobalt oxide enables the improvement of the sintering and densification of specimens and the significant enhancement in total conductivity due to (i) the improved material reducibility at high temperatures and the associated rise in electronic conductivity and (ii) the increased concentration of oxygen vacancies.

XPS measurements proved the existence of Tb$^{3+}$ on the CT20 and CT20Co surfaces and thus the availability of oxygen vacancies. The oxygen exchange rate of cobalt-free and cobalt-containing CTx materials was measured using PIE. The highest surface exchange rates were obtained for CT50 and CT50Co samples. On the whole, the results indicate that the exchange rate exhibited by CTx is limited by the dissociative adsorption of molecular oxygen.
at the oxide surface, and its magnitude is bounded by the relative magnitudes of the ionic and electronic conductivities in the materials.

Oxygen permeation was negligible for cobalt-free samples while CT10Co and CT20Co showed an oxygen flux at 1000 °C and 1 mm thickness of ca. 0.02 and 0.08 mL·min\(^{-1}\)·cm\(^{-2}\) for argon sweeping, respectively. CT10Co oxygen permeation flux was increased until reaching 0.5 mL·min\(^{-1}\)·cm\(^{-2}\) by using methane as sweep gas at 1000 °C. Permeability results confirm the higher ambipolar conductivity of the CT20Co composition and the possibility to achieve suitable mixed conductivity values by the appropriate dopants and doping level.

Finally, thermogravimetric tests carried out in 5% CO\(_2\)–containing air confirmed the stability of ceria material in contrast with the instability observed for Ba- and Sr-containing perovskites. The combined oxygen permeation and stability under reducing and carbonated atmospheres suggests the application of Ce\(_{1-x}\)Tb\(_x\)O\(_{2.5+}\) + Co 2% as membrane materials in catalytic membrane reactors for hydrocarbon and syngas conversion and oxygen separation using flue gas sweeping.

From the conductivity and surface results, a good membrane would be expected from CT20, CT50 and CT50C0 materials. However, new manufacturing methods need to be developed to obtain dense monoliths. New sintering strategies that allow obtaining dense specimens may be intended in the future, as spark plasma sintering or the combination of the material in dual-phase membranes.
5.4. References

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Chapter 6. LSM-ceria composite cathodes for SOFC

Abstract

Ceria-based materials have been incorporated into LSM conventional cathodes in order to improve electrochemical performance. Gadolinium and other elements (Mg, Cr and Bi) have been used as ceria co-dopants for enhancing conductivity and electrocatalytic properties of several LSM-ceria composites. Among different tested LSM-ceria cathodes, Ce$_{0.8}$Gd$_{0.1}$Mg$_{0.1}$O$_{2-\delta}$-based composite offers the lowest electrode polarization resistance in the 700-900 °C temperature range. However, at the lowest temperatures these results are similar to those for the Ce$_{0.8}$Gd$_{0.2}$O$_{2-\delta}$ binary system, thus the selection of Gd-co-doping elements should principally address economic issues. Since LSM-based cathodes are pointed for high temperature applications (>800 °C) this magnesium-co-doped ceria-gadolinia compound could be an alternative to widely used Ce$_{1-x}$Gd$_x$O$_{2-\delta}$ binary systems for composite SOFC cathodes.
6.1. Introduction

Strontium doped lanthanum manganite \((\text{La}_{1-x}\text{Sr}_x\text{MnO}_3 - \text{LSM})\) perovskites are the principal component of the state-of-the-art materials for SOFC cathodes. LSM-based compositions show interesting electrochemical properties for oxygen activation at high temperatures while they are thermochemically compatible with the widely used YSZ electrolyte materials. However, the lack of ionic conductivity and the decreasing performance of LSM cathodes at temperatures below 800 °C require mixing LSM cathodes with electrolyte materials. These LSM-electrolyte composites allow increasing the global ionic conductivity of the cathode while augments the three phase boundary (TPB) region thus enhancing the cathode performance. In order to increase the electrochemical performance of LSM-based composite cathodes, it is desirable that the electrolyte-like component possesses high oxygen ion conductivity, some electronic paths and/or enhanced catalytic activity for the oxygen reduction reaction at the operating temperature.

Rare earth-doped cerias studied in the previous chapters have demonstrated mainly ionic conductivity. Among them, gadolinia doped cerium oxide (CGO) shows the highest ionic conductivity, which is also higher than that of the YSZ electrolyte. Hence, this material seems to be an interesting electrolyte material, specifically at low temperatures. [1, 2] In addition, thermochemical properties of CGO material fit much closer with LSM than YSZ electrolytes. [3] The presence of gadolinium favors the oxygen ion conductivity while the addition of other doping elements may promote
electronic conductivity. Moreover, the use of doped ceria materials in composite cathodes allows incorporating additional redox electrocatalytic activity. [4, 5]

In order to increase the ionic conductivity and improve other related properties of the materials, co-doping approach has been proved to be effective. Co-doping ceria with alkaline earth and rare earth ion shows significantly higher conductivity in air than singly doped materials with the same dopant concentration. [6-8] The possibility of adding mixed ionic-electronic conductivity is a plus for the use as a cathode, since the electronic conduction is not blocked by the electrolyte particles, thus boosting the TPB area.

With the premises exposed above, this chapter shows the electrochemical performance of composite SOFC cathodes, fabricated with LSM and different co-doped gadolinium-cerium oxides based on the Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ (X = Gd, Cr, Mg, Bi, Ce) nominal formula.

6.2. Results and discussion

Structural characterization

The La$_{0.8}$Sr$_{0.2}$MnO$_3$ (LSM) commercial powders are obtained from Fuel Cell Materials (USA). Pure fluorite compounds with the nominal formula Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ (X = Gd, Cr, Mg, Bi, Ce) were synthesized by means of co-precipitation method after an 800 °C firing step under air. Table 6.1 summarizes the short nomenclature of the compounds that is used along the text. The compositions comprise both CG82 and CG91 parent materials and also the corresponding magnesium-, bismuth- and chromium-co-doped materials.
Figure 6.1 plots the XRD patterns of the as-sintered powders showing the expected cubic fluorite structure. Diffraction peaks corresponding to any other oxide or any precursor are not observed, which means that the different dopants are fully dissolved into the ceria lattice. Crystallographic lattice parameters ($a$) for the ($h k l$) planes were calculated using the corresponding equation for a cubic system and the results are summarized in Table 6.1. Crystallite sizes were extracted by fitting the (4 2 2) plane reflection to a Gaussian function. [9] The small amount of the co-dopant has small influence on the lattice size, calculated from the XRD patterns (Table 6.1), but it seems to have some effect on the ceria crystallite size as compared from Table 6.1. Indeed, by using the same synthesis method and sample processing, it is possible to obtain a range of crystallite sizes depending on the co-doping added element. Further investigations should be focused on this fact, studying the
nucleation and crystal growth during precipitation and later sintering process, and its influence on particle size.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Nomenclature</th>
<th>Lattice parameter (Å)</th>
<th>Crystallite size (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ce$<em>{0.8}$Gd$</em>{0.2}$O$_{2-δ}$</td>
<td>CG82</td>
<td>5.4224 (1)</td>
<td>529</td>
</tr>
<tr>
<td>Ce$<em>{0.9}$Gd$</em>{0.1}$O$_{2-δ}$</td>
<td>CG91</td>
<td>5.4152 (1)</td>
<td>535</td>
</tr>
<tr>
<td>Ce$<em>{0.8}$Gd$</em>{0.1}$Bi$<em>{0.1}$O$</em>{2-δ}$</td>
<td>CGBi</td>
<td>5.4153 (3)</td>
<td>255</td>
</tr>
<tr>
<td>Ce$<em>{0.8}$Gd$</em>{0.1}$Cr$<em>{0.1}$O$</em>{2-δ}$</td>
<td>CGCr</td>
<td>5.4143 (9)</td>
<td>169</td>
</tr>
<tr>
<td>Ce$<em>{0.8}$Gd$</em>{0.1}$Mg$<em>{0.1}$O$</em>{2-δ}$</td>
<td>CGMg</td>
<td>5.4159 (6)</td>
<td>477</td>
</tr>
</tbody>
</table>

Table 6.1 Cerium-based compositions synthesized as cubic fluorite structure with the nomenclature used along the text, and the corresponding lattice parameter and crystallite size extracted from XRD patterns.

Irrespective of differences in crystallite size among co-doping elements (Table 6.1), co-precipitation synthesis method permits obtaining powders nanometric particles, as observed in Figure 6.2a. Comparing, for instance, the particle size of the ceria-based CG82 material with the as-received LSM powder ($d_{50} = 0.6–0.3 \, \mu m$) it can be observed by SEM imaging (Figure 6.2a and b) that particle size is much smaller for the ceria-based co-precipitated powders.
Figure 6.2. Comparison of the initial particle size of a) CG82 (as-sintered) and b) LSM (as-received) and the resulting powders after calcination at 1150 °C: c) CG82 and d) LSM.

In addition and despite the fact that initial sintering temperature for the ceria-based compounds is lower than the commercial LSM (1000 °C), after a 1150 °C calcination step, LSM powder has bigger and more sintered grains than ceria-based CG82 powder (Figure 6.2c and d) indicating the higher sintering activity of LSM material compared to the co-precipitated ceria compound. On the contrary, ceria-based CG82 material seems to produce smaller grains, with respect to LSM, after treatment at high temperature even when the initial synthesis temperature is low (800 °C).

Regarding this particle size evolution it should be assumed that the bigger LSM particles will become covered by smaller ceria-based ones in a
non-well distributed composite. However, SEM images of several LSM-ceria composite cathodes (Figure 6.3) revealed that particle size is similar for both materials (LSM and ceria-based) after sintering together at 1150 ºC. Therefore, the sintering evolution of composite components that have been together calcined is not as different as when materials are sintered separately. This fact enables that mixtures of these materials have well distributed and connected particles and the fabricated composite cathodes can maintain proper conducting paths and enough porosity for SOFC cathode purposes. Spatial constraints favored by the good mixing of LSM and doped-ceria phases in the composite cathode allow preventing particle agglomeration and coarsening of the electrode materials. Therefore, both materials achieve similar particle size when calcined together. In addition, the presumably presence of small nanoparticles of ceria compounds (Figure 6.2c) on the surface of bigger LSM grains might be beneficial from the point of view of the cathode performance, as reported for both LSM-YSZ [10] and GDC-impregnated LSM. [11] This could be a further optimization step, which would entail a final infiltration step of the previously sintered cathode.

Figure 6.3. SEM images of several composite cathodes sintered at 1150 ºC: a) CG82-LSM, b) CG91-LSM and c) CGBi-LSM.
**DC-conductivity characterization**

As an extension of the Chapter 3, metallic dopants are introduced in combination with the Gd in order to improve the transport properties. Electrical conductivity of ceria-based materials under air is shown in Figure 6.4a at several temperatures. As described for several experimental works on Ce$_{1-x}$Gd$_x$O$_{2-\delta}$ system, it can be observed that the total conductivity for CG91 is lower than for CG82 (Figure 6.4a). [12, 13] Conductivity enhancement on the ceria fluorite lattice obtained by Gd$^{3+}$ addition arises from the ionic conductivity introduced by extrinsic vacancies formed to maintain the electroneutrality in the structure. Nevertheless, the best ratio for CeO$_2$-Gd$_2$O$_3$ solid solution is still not clearly defined for applications as solid oxide fuel cells, since it depends on the proposed limiting conductivity mechanism. [14-17] Co-doped materials with the general formula Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ have a cerium and gadolinium molar content in between CG82 and CG91. Conductivity of these co-doped ceria materials (Figure 6.4a) lies in between the values observed for 20% and 10% Gd-doped ceria, i.e., CG82 and CG91. The total conductivity of these co-doped materials is high enough for SOFC cathode applications since CG82 and CG91 conductivity fits with the SOFC application requirements.
Among the different co-doped materials, those based on bismuth (CGBi) and magnesium (CGMg) have similar conductivities. These conductivity values are close to those obtained for CG82 material at low temperatures (Figure 6.4a). On the other hand, CGCr composition has lower conductivity than the other two co-doped materials and it is even lower than CG91 compound when temperature decreases. Therefore, by using Mg or Bi, it is possible to reduce gadolinium and/or cerium content in ceria-gadolinia materials without significantly decreasing the total electrical conductivity. This partial replacement of the gadolinium or cerium by other elements would make possible to reduce production costs whereas both electrochemical and catalytic activity are not significantly affected.

In addition to the conductivity measurements performed under air, different oxygen partial pressure atmospheres were used for measuring the total conductivity of these materials (Figure 6.4b) at 800 °C. The studied range of $pO_2$ includes both oxidant (0.21 - 1 atm) and moderately reducing atmospheres (10^{-5} - 0.21 atm), in where Ce^{4+} does not reduce to Ce^{3+}. [18-21] Conductivity
values in this oxygen partial pressure range (Figure 6.4b) seem to be independent on \( pO_2 \) for all the materials, and consequently \( p \)-type conductivity is negligible. Therefore, conductivity of these ceria-based materials is mainly ionic in these oxidizing conditions. It seems not possible to promote the electronic conductivity either in the bulk or in the grain boundary through the chosen doping strategy on the ceria-based fluorite structure.

**Cathode testing on symmetric cells**

The enhancement of electrocatalytic properties of LSM cathode after mixing with ceria-based materials is shown in Figure 6.5. Electrode polarization resistance (Rp) values were obtained from electrochemical impedance spectroscopy (EIS) measurements on symmetrical cells, fabricated with the different LSM-ceria-based composite cathodes and using CG82 cathode that used Co as a sintering aid (CG82-Co). The chosen LSM to ceria ratio (in weight) is 50/50 because it has been reported as the optimum for SOFC LSM-based cathode purpose. Irrespective to the ceria-based composition, the corresponding Rp of the resulting composite electrode is lower than single LSM electrode at any tested temperature. This confirms that the addition of ion conducting paths (ceria-based percolating network) allows extending the TPB area into a certain thickness of the electrode. Indeed, the presence of ionic conducting phase on LSM-YSZ state-of-the-art composite cathode also introduces ionic paths and increases TPB sites thus reducing the electrode polarization resistance compared to single LSM cathode. As previously explained, the use of ceria-compounds instead of YSZ materials could also increase the composite cathode electroactivity. [4] As a consequence, obtained electrode polarization resistance for ceria-based composites is lower than for LSM-YSZ cathodes as observed in Figure 6.5.
Figure 6.5 shows that the electrode polarization resistance values for the LSM-co-doped-ceria composites lie in between LSM-CG82 and LSM-CG91 electrodes. This is in agreement with the previously observed electrical conductivity measurements of single ceria-doped materials (Figure 6.4a). The best composite electrode regarding $R_p$ values (Figure 6.5) is the LSM-CGMg cathode at any tested temperature. It also presents lower electrode polarization resistances than the LSM-CG82 cathode at higher temperatures (800-900 °C). Nonetheless, electrode resistance of LSM-CGMg increases with decreasing operating temperatures, i.e., this cathode presents high activation energy. This performance worsening may be ascribed to the less mobile oxygen vacancies at lower temperatures. Chromium- or bismuth-co-doping do not improve the electrochemical reaction for oxygen reduction with respect to the LSM-CG82 composite.
Moreover, it can be inferred from Figure 6.5 that all ceria-based material composites exhibit higher polarization resistances than the well-established high-performing LSFC (tested on same symmetrical cell support) cathode. Blending ionic materials (CGMg, CGBi, CGCr and Gd-doped ceria) with LSM allows increasing the concentration of TPB sites. However, the extension of the active zone for the oxygen reduction reaction in these composites is still restricted to TPB points and not to the whole electrode surface area, as for LSFC mixed ionic-electronic conducting (MIEC) electrode. Despite that fact, composite microstructure of the present ceria-based electrodes could be further optimized in order to approach the corresponding performance of MIEC cathodes. [11, 12, 22]

![Figure 6.6. Polarization resistance as a function of conductivity at three different temperatures (700, 750 and 800 °C)](image)

Figure 6.6 represents the polarization resistance of LSM-ceria-based composites as a function of ceria-based material conductivity at three temperatures (700, 750 and 800 °C) for all tested electrodes. In general, there
exists a trend suggesting that the resistance decreases with increasing conductivities for selected cathode composites. However, all data points do not fall in the same (straight) trend line, which suggests that other factors are affecting the polarization resistance such as surface reaction, interfacial resistance, electrode microstructure, etc.

Figure 6.7. Cross section SEM images of a) CGCr-LSM, b) CG82-LSM and c) CG91-LSM electrodes on a dense CG82-Co electrolyte. The upper layer seen in a) corresponds to the gold current collecting mesh.

The high density of CG82-Co electrolyte and the thickness of all porous electrode layers (~35 µm) can be inferred from the SEM images recorded on symmetrical cells fracture cross-section in Figure 6.7. [9] Electrodes are apparently very well attached to the electrolyte upon sintering in air at 1150 ºC. Moreover, the grains of the two materials of the composite are well connected and distributed. Despite the initial particle size of the different doped-cerias varies significantly (Table 6.1 and Figure 6.2a), a similar particle size distribution is observed for all electrodes, i.e., when they are mixed and sintered together with LSM (Figure 6.3 and Figure 6.7).

Impedance spectroscopy modeling

Previous results were analyzed in terms of the electrode polarization resistance obtained from EIS measurements. As an example of such EIS
obtained data, Figure 6.8a represents the Nyquist plot for the different LSM-ceria-based composites while Figure 6.8b shows the corresponding imaginary Bode plot, both at 800 °C under air. Spectra in Figure 6.8a correspond to the sum of both identical electrode contributions from symmetrical cells and from it the electrode polarization resistance can be calculated. In addition, data presented in Figure 6.8 could also be fitted to an equivalent circuit comprising three ZARC elements, as named by J. R. McDonald, [23] in series with an inductance (from wiring) and a pure resistance, which comes mainly from current collection and electrolyte resistance. Indeed, in Figure 6.8a the values corresponding to the pure resistance (R) vary among samples because of current collecting and contact resistances.

Figure 6.8. Electrochemical impedance spectra of symmetrical cells with different LSM-doped-ceria electrodes at 800 °C under air: a) Nyquist and b) imaginary impedance Bode plots. Logarithm of selected frequencies is provided on the Nyquist plot.

Fitting values of spectra in Figure 6.8, regarding to resistance, constant phase element parameters (Q₀ and n) and corresponding summit frequency of ZARC elements, are presented in Table 6.2, in which the area-corrected resistance values for single electrode contribution are presented. Taking into account the calculated summit frequency (as the inverse product of the resistance times the equivalent capacitance of the constant phase element [24]),
two processes occur at similar relaxation times at high frequencies (A and B) and the other one at relatively low frequencies (C).

<table>
<thead>
<tr>
<th>Fit Units</th>
<th>LSM-CG91</th>
<th>LSM-CG82</th>
<th>LSM-CGMg</th>
<th>LSM-CGCr</th>
<th>LSM-CGBi</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_A$ $\Omega \cdot cm^2$</td>
<td>0.21</td>
<td>0.10</td>
<td>0.09</td>
<td>0.21</td>
<td>0.12</td>
</tr>
<tr>
<td>$Q_{0A}$ $S \cdot s^n$</td>
<td>$5.26 \cdot 10^{-5}$</td>
<td>$4.99 \cdot 10^{-5}$</td>
<td>$1.08 \cdot 10^{-5}$</td>
<td>$6.34 \cdot 10^{-5}$</td>
<td>$1.21 \cdot 10^{-4}$</td>
</tr>
<tr>
<td>$n_A$ $-$</td>
<td>0.91</td>
<td>0.96</td>
<td>1.00</td>
<td>0.90</td>
<td>0.86</td>
</tr>
<tr>
<td>$\omega_A$ Hz</td>
<td>$8.10 \cdot 10^4$</td>
<td>$1.08 \cdot 10^5$</td>
<td>$1.74 \cdot 10^5$</td>
<td>$8.00 \cdot 10^4$</td>
<td>$1.10 \cdot 10^5$</td>
</tr>
<tr>
<td>$R_B$ $\Omega \cdot cm^2$</td>
<td>0.77</td>
<td>0.48</td>
<td>0.41</td>
<td>0.69</td>
<td>0.64</td>
</tr>
<tr>
<td>$Q_{0B}$ $S \cdot s^n$</td>
<td>$9.62 \cdot 10^{-4}$</td>
<td>$1.01 \cdot 10^{-3}$</td>
<td>$5.80 \cdot 10^{-4}$</td>
<td>$1.30 \cdot 10^{-3}$</td>
<td>$1.28 \cdot 10^{-5}$</td>
</tr>
<tr>
<td>$n_B$ $-$</td>
<td>0.72</td>
<td>0.75</td>
<td>0.75</td>
<td>0.70</td>
<td>0.71</td>
</tr>
<tr>
<td>$\omega_B$ Hz</td>
<td>$4.65 \cdot 10^3$</td>
<td>$6.10 \cdot 10^4$</td>
<td>$1.47 \cdot 10^4$</td>
<td>$4.47 \cdot 10^3$</td>
<td>$4.49 \cdot 10^3$</td>
</tr>
<tr>
<td>$R_C$ $\Omega \cdot cm^2$</td>
<td>0.29</td>
<td>0.06</td>
<td>0.05</td>
<td>0.36</td>
<td>0.08</td>
</tr>
<tr>
<td>$Q_{0C}$ $S \cdot s^n$</td>
<td>$8.23 \cdot 10^{-2}$</td>
<td>$6.45 \cdot 10^{-1}$</td>
<td>$4.26 \cdot 10^{-1}$</td>
<td>$5.11 \cdot 10^{-2}$</td>
<td>$4.52 \cdot 10^{-1}$</td>
</tr>
<tr>
<td>$n_C$ $-$</td>
<td>0.54</td>
<td>0.54</td>
<td>0.58</td>
<td>0.56</td>
<td>0.61</td>
</tr>
<tr>
<td>$\omega_C$ Hz</td>
<td>$1.21 \cdot 10^2$</td>
<td>$5.12 \cdot 10^1$</td>
<td>$9.82 \cdot 10^1$</td>
<td>$1.63 \cdot 10^2$</td>
<td>$3.47 \cdot 10^1$</td>
</tr>
</tbody>
</table>

Table 6.2. Fitting parameters obtained for each LSM-ceria-based composite at 800 ºC under air. $R$ is the resistance, $Q_0$ and $n$ are the constant phase element parameters and $\omega$ the summit frequency of each ZARC element, respectively. Capital letters (A, B, C) indicate the process according to the proposed electrical equivalent circuit.

Electrochemical impedance response of electrodes based on LSM-YSZ composites has been described to possess up to five contributions. [25] Nonetheless, three different electrode processes named A, B and C seem to be more predominant in Figure 8 spectra. [26] According to the possible processes occurring in ceramic-ceramic composite cathodes for oxygen reduction reaction and the proposed model in Figure 6.9 and results (Table 6.2), [25-27] the two high frequency processes (named A and B) can be related to (1) transference of the oxygen intermediates/oxide ions or electrons [28-30] between ceria and LSM particles; and (2) the transport through the ceria-based particles of the composite towards the electrolyte. On the other hand, the low frequency
contribution (named C) is related to the dissociative adsorption, dissociation and transfer of species at TPB coupled with the surface diffusion, similar to the processes occurring in MIEC cathodes. The last assignment is supported on the exponent values close to 0.5 of the constant phase elements \( (n_C) \) (Table 6.2). Therefore the ZARC configuration could also have been modeled by a diffusion element [31] like a Gerischer or a Warburg. [32, 33] However, in order to directly analyze the contribution of the resistance of each process to the total electrode resistance, it was decided to model this contribution using a ZARC element.

![Figure 6.9](image)

Figure 6.9. Resistances corresponding to the different processes obtained by fitting the spectra of symmetrical cells at 800 °C under air according to the equivalent circuit \( LR(Q_A)(R_0Q_B)(R_0Q_C) \) shown in the inset. The corresponding logarithm of summit frequency is indicated on top of each bar.

In Figure 6.9 the resistance values for each fitted process are compared. Composite cathodes based on CG82 and CGMg showed resistance values lower than CG91- and CGCr-based composites for all three (A, B, C) processes. For
LSM-CGBi cathode, the resistance assigned to process B seems to limit the whole cathode performance (Figure 6.9) even when the other two contributions (A and C) have relatively low values. CGMg-based cathode presents the lowest LF resistance of all and this improvement of the surface processes should be related to both (1) the higher surface concentration of oxygen vacancies/active (adsorption) sites in the oxygen reduction mechanism; and (2) the alkali character of the Mg cation, which improves the molecular oxygen adsorption. On the contrary, the highest LF contribution is observed for CG91 and CGCr. This could be caused by the lower surface concentration of oxygen vacancies and, in the particular case of CGCr, some detrimental effect of Cr species on the ceria surface and TPB sites.

The nature of limiting processes at the highest frequency range (A and B) cannot be unambiguously established. However, assuming that total conductivity at 800 °C for all doped cerias is similar (Figure 6.4a), high frequency limitations should be originated principally from cathode microstructure, i.e., effective paths for oxygen ions/intermediates or electrons transfer are limited. [26] Additionally, the potential role of grain boundary oxygen resistances of these ceria-based compounds cannot be excluded, especially at lower temperatures. [18]

On the whole, high amount of trivalent/divalent dopants (e.g. Gd$^{3+}$ and Mg$^{2+}$), in the ceria lattice benefits the oxygen vacancy formation and the resulting processes, at different frequency ranges, are enhanced due to the promotion of adsorption/dissociation steps, but also by better surface and bulk diffusion processes. On the contrary, multivalent chromium or gadolinium-isovalent bismuth have a detrimental effect on the global polarization resistance of the electrode. As a consequence, all three limiting processes analyzed and modeled here appear to be highly related one to each other.
6.3. Conclusions

Several ceria-based materials have been properly synthesized by co-precipitation method. Total electrical conductivity of Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ (X = Gd, Cr, Mg, Bi, Ce) materials has been proved to be sufficient for SOFC applications. Specifically, the electrochemical properties of several LSM-Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ composite cathodes have been analyzed in symmetrical cells. In general, polarization resistance decreases with increasing ceria-based materials total conductivities. Among different co-dopants, Mg seems to enhance both the oxygen ion transferring and the surface processes for oxygen reduction. Addition of chromium and bismuth to the ceria-gadolinia-based materials seems not to improve the LSM-composite cathode performance. Nevertheless, performance and conductivity of these new co-doped composite materials as SOFC cathodes is not significantly higher than Ce$_{0.8}$Gd$_{0.2}$O$_{2-\delta}$ material. The use of the Mg-Gd-co-doped ceria may be viable due to economic reasons, regarding the price of cerium and gadolinium raw materials. Microstructural issues should be focused in order to further improve cathode performance.

6.4. References


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Chapter 7. Ionic transport
improvement on CaTiO$_3$ based perovskites

Abstract

The second family of materials under research comprises CaTiO$_3$-based perovskites. CaTiO$_3$ perovskite is a $p$-type electronic conductor whose ionic conductivity may be increased by doping with lower valent cations. However, the stability of the structure may be drastically affected by the guest cations turning temperature or atmosphere dependent or developing a different non-perovskite structure. The series Ca$_{1-x}$A$_x$Ti$_{1-x}$(M1$_{1-y}$M2$_y$)$_x$O$_{3-δ}$ has been prepared by solid state reaction to evaluate the structural stability and ionic conductivity improvement of CaTiO$_3$ perovskites. The first screening shows that even the structural predictions are contrary, single phase perovskite structure is not achieved for all the compositions. DC conductivity indicates that all the dopants allow increasing the conductivity, especially CaTi$_{0.8}$Fe$_{0.2}$O$_{3-δ}$ and CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3-δ}$. The former has been well reported in literature and compared with the second one in the present chapter.

Bulk transport and surface properties have been characterized by means of DC conductivity up to 1000 °C, thermogravimetry (TG) and temperature programmed desorption (TPD), X-ray photoelectron spectroscopy (XPS) and pulse isotope exchange (PIE). The present study presents the preparation of a membrane made of CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3-δ}$. The systematic study of the
membrane operation variables on the oxygen flux is shown, *i.e.* inlet gas flow rate at sweep membrane compartment, temperature and oxygen feed gas nature. Moreover, membrane stability is proved under CO$_2$ as sweep gas.
7.1. Introduction

Perovskite-structured compounds with the formula ABO$_3$ take their name from the mineral CaTiO$_3$. [1] The ideal perovskite possesses the cubic symmetry in which A is a large cation sited on the corners in 12-fold oxygen coordination (as a lanthanide, alkali or alkali earth), while the body center B is a small cation in 6-fold oxygen coordination, as transition metals or small lanthanides. The oxygen anions occupy the faces center. Similar to the fluorites, the structure is able to host cations with different valence and ionic radii to achieve high concentrations of oxygen vacancies and ionic conductivity. This affects the crystal stability, which is predicted by the tolerance factor established by Goldschmidt

$$t = \frac{r_A + r_O}{\sqrt{2}(r_B + r_O)} \quad \text{(Eq. 7.1)}$$

where $r_A$, $r_B$ and $r_O$ are the ionic radii of the cations in A and B site and the oxygen anion, respectively. The A cation in an ideal perovskite has the same size as the oxygen anion (1.4 Å) forming a well packed cube that hosts the B-cation. Cubic perovskites will be obtained for values close to $t = 1$, while other distorted perovskites are obtained provided $0.8 < t < 1$. Goldschmidt tolerance condition is necessary but not sufficient for the stability of the perovskite. Another governing parameter for the formability of perovskites is known as octahedral factor (o.f.), the ratio of radii of the small cation B over the radii of anion O, which is related to the stability of BO$_6$ basic octahedron. The lowest limit for the o.f. to deliver a stable perovskite is 0.425, which means that the minimum $r_B$ is 0.53 Å. [2]
Perovskites as LaGaO$_3$ are mainly ionic conductors, while in perovskites containing Fe, Co, Ni or Mn the electronic conductivity is usually orders of magnitude larger than the ionic conductivity. [3, 4] The ionic conductivity takes place by the hopping of oxygen ions from one oxygen vacant site to neighbouring vacant sites, so that the oxygen deficiency ABO$_{3-\delta}$ is needed. The A site is commonly used to tune the oxygen vacancy concentration while the B site is basically responsible for overall electronic conductivity through $B^{n+}-O-B^{(n+1)+}$ conduction pairs. [5] For example, the ionic conductivity can be enhanced by the substitution of A-site cations with lower valent cations. On the contrary, higher valent ions may increase the phase stability, although the annihilation of oxygen vacancies is expected to decrease the ionic conductivity. Partially substituting the A site by larger isovalent cations may improve the chemical and structural stability in reducing environments while keeping the oxygen flux levels. [4] On the other hand, the decrease of average A cations radii may reduce the oxygen flux. [4]

Figure 7.1. CaTiO$_3$ orthorhombic perovskite from the family A$^{II}$B$^{IV}$O$_3$. 

![CaTiO3 orthorhombic perovskite from the family A$^{II}$B$^{IV}$O$_3$.](image-url)
As mentioned in Chapter 1, iron and cobalt-based perovskites typically possess high electron conductivity as well as good ionic conductivity, so that intensive research devoted to the preparation and characterization of MIEC perovskite membranes has been developed in the last decades. [6-9] The most extensively used perovskite materials are large A-site BSCF and LSCF, which despite their high oxygen fluxes still present fundamental disadvantages for a proper use, as temperature dependent phase transitions. [4, 10-12] On the other hand, CaTiO₃ based materials are stable perovskite in a vast range of temperatures and pO₂. [13] 

The perovskite CaTiO₃ represented in Figure 7.1 is slightly distorted from cubic structure and belong to orthorhombic symmetry with *Pnma* spatial group. [1, 13] CaTiO₃ is a MIEC at low pO₂, but the electrolytic range of pO₂ may be extended to higher oxygen content atmospheres by doping, since CaTiO₃ is able to keep the perovskite structure also at high defect concentrations. When acceptor dopants are substituted in the Ti site, both ionic and electronic conductivities may be enhanced. The ionic conductivity can be also enhanced by the substitution of Ca cations with lower valent cations, while higher valent dopants create A-site deficiencies. [14, 15] In this study, different dopants have been introduced in the mainly electronic CaTiO₃, attending to criteria as ionic radii affinity and aliovalence or multivalence of the cations with the aim of improving the ionic and the electronic conductivity of the material. A first approach has been done to obtain the perovskite structure discarding such compositions that were not single phase or perovskite at all. The compounds showing the perovskite structure were tested by DC conductivity with the aim of selecting MIEC materials suitable for oxygen transport.
7.2. Results and discussion

The series Ca_{1-z}A_{z}Ti_{1-x}(M_{1-y}M_{2y})_{x}O_{3-\delta} has been prepared by solid state reaction. The synthesis has been done at 1200 °C during 14 hours to favour the formation of the perovskite phase. Several dopants have been selected on their ionic radii and aliovalences of the substituted element. Table 7.1 lists the combination of dopants and the ionic radii of A and B positions, as well as an identification number. A preliminary prediction of the formation of stable perovskite phase has been tried on the basis of the Goldschmidt and octahedral factors, and the experimental results are also exposed. The crystalline phase(s) of the materials were determined by XRD. Even though all the compounds exhibit 0.8 < t < 1 and o.f. < 0.425 (except CaTi_{0.8}Fe_{0.2}O_{3-\delta} and CaTi_{0.6}Fe_{0.4}O_{3-\delta}), several of them did not deliver pure perovskite structure. Therefore, they were not further investigated. Several reasons may contribute to the instability, as that the dopant does not substitute in the expected site or with the expected oxidation state (so that expected ionic radii) and thus the predictions are not valid.
### Ionic transport improvement on CaTiO₃-based perovskites

Table 7.1. Ionic radii of constituent ions A and B, tolerance factor, octahedral factor (r_B/r_A) and symmetry of different compounds (O=orthorhombic, X=not single phase)

<table>
<thead>
<tr>
<th>Material</th>
<th>#</th>
<th>r_A(Å)</th>
<th>r_B(Å)</th>
<th>t</th>
<th>o.f.</th>
<th>Symmetry</th>
</tr>
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<td>CaTiO₃</td>
<td>0</td>
<td>1.34</td>
<td>0.605</td>
<td>0.966</td>
<td>0.432</td>
<td>O</td>
</tr>
<tr>
<td>CaTiₓ₂Feₓ₂O₄₋δ</td>
<td>1</td>
<td>1.34</td>
<td>0.594</td>
<td>0.972</td>
<td>0.424</td>
<td>O</td>
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<td>1.34</td>
<td>0.6</td>
<td>0.969</td>
<td>0.428</td>
<td>O</td>
</tr>
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<td>0.960</td>
<td>0.441</td>
<td>O</td>
</tr>
<tr>
<td>CaTiₓ₂Gaₓ₂O₄₋δ</td>
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<td>0.608</td>
<td>0.965</td>
<td>0.434</td>
<td>X</td>
</tr>
<tr>
<td>CaTiₓ₂Mgₓ₂O₃₋δ</td>
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<td>1.34</td>
<td>0.617</td>
<td>0.961</td>
<td>0.440</td>
<td>O</td>
</tr>
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<td>0.966</td>
<td>0.432</td>
<td>O</td>
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<td>0.606</td>
<td>0.966</td>
<td>0.432</td>
<td>X</td>
</tr>
<tr>
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<td>1.34</td>
<td>0.610</td>
<td>0.964</td>
<td>0.436</td>
<td>X</td>
</tr>
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<td>1.34</td>
<td>0.613</td>
<td>0.963</td>
<td>0.438</td>
<td>X</td>
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<td>CaTiₓ₂Crₓ₁Mgₓ₁O₃₋δ</td>
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<td>0.960</td>
<td>0.441</td>
<td>O</td>
</tr>
<tr>
<td>CaTiₓ₂Nbₓ₁Mgₓ₁O₃₋δ</td>
<td>9</td>
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<td>0.443</td>
<td>O</td>
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<td>0.970</td>
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<td>0.612</td>
<td>0.963</td>
<td>0.437</td>
<td>X</td>
</tr>
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<td>CaTiₓ₂Coₓ₁Mgₓ₁O₃₋δ</td>
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<td>0.963</td>
<td>0.436</td>
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</tr>
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<tr>
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<td>0.963</td>
<td>0.437</td>
<td>X</td>
</tr>
<tr>
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<td>0.960</td>
<td>0.441</td>
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<td>0.950</td>
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<td>O</td>
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<tr>
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<td>0.966</td>
<td>0.433</td>
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<td>1.34</td>
<td>0.601</td>
<td>0.968</td>
<td>0.429</td>
<td>X</td>
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<tr>
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<tr>
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<td>0.969</td>
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<td>0.612</td>
<td>0.963</td>
<td>0.437</td>
<td>X</td>
</tr>
</tbody>
</table>
The lattice parameters of the succeeded perovskites, with pure perovskite crystalline phase, have been determined from XRD and summarized in Table 7.2.

<table>
<thead>
<tr>
<th>Material</th>
<th>#</th>
<th>Cell volume (Å³)</th>
<th>a (Å)</th>
<th>b (Å)</th>
<th>c (Å)</th>
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<td>7.640 (2)</td>
<td>5.440 (1)</td>
<td>5.382 (1)</td>
</tr>
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<td>CaTi₀.₉Fe₀.₁₂O₃₄</td>
<td>T1</td>
<td>223.35</td>
<td>7.637 (4)</td>
<td>5.425 (3)</td>
<td>5.391 (3)</td>
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<td>7.65 (1)</td>
<td>5.435 (7)</td>
<td>5.372 (5)</td>
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<td>7.640 (2)</td>
<td>5.440 (1)</td>
<td>5.382 (1)</td>
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<tr>
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<td>T18</td>
<td>224.18</td>
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<td>5.447 (5)</td>
<td>5.379 (5)</td>
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<td>225.32</td>
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</tr>
<tr>
<td>Ca₉₈Bi₂₆Ti₉₈Fe₂₆O₃₄</td>
<td>T26</td>
<td>223.82</td>
<td>7.70 (2)</td>
<td>5.436 (9)</td>
<td>5.349 (2)</td>
</tr>
</tbody>
</table>

Table 7.2. Cell volume and cell parameters of formed perovskite compounds (fitting errors in brackets).

DC electrical conductivity measurements have been conducted on sintered rectangular bars, both as a function of temperature and in different oxygen containing atmospheres. Figure 7.2 shows the total conductivity dependence on the $pO_2$ of doped calcium titanates that present pure perovskite structure. Samples T3, T5 were not measured since they reacted with the holder upon calcination. T26 is not dense even uprising the sintering temperature, which explains the low conductivity.
All the introduced dopants are able to increase total conductivity with respect to pure CaTiO$_3$. The majority of these materials diminish conductivity as the $pO_2$ decreases indicating that they are mainly $p$-type electronic conductors, even though doping with lower valence ions produces oxygen vacancies and ionic conductivity was also expected. On the other hand, CaTi$_{0.95}$Mo$_{0.05}$Mg$_{0.1}$O$_{3-\delta}$ (T10), CaTi$_{0.9}$Al$_{0.1}$O$_{3-\delta}$ (T11) and CaTi$_{0.9}$Mn$_{0.1}$O$_{3-\delta}$ (T13) are not $pO_2$ dependent, which suggests mainly ionic conduction.

Among all of them, the best conductivity is found when doping the B-site with 20% Fe, which has been widely studied in literature and used here as a reference. [13, 16-19] Iron oxide is an acceptor dopant that enhances the total conductivity in oxidising conditions by simultaneously increasing the $p$-type and the ionic conductivity. In a similar composition Mg$^{2+}$ is added with the aim of further increase the oxygen vacancy concentration, i.e.
CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3.6}$ (T6). The preliminary conductivity screening shows that this is the only composition with a comparable electric behaviour with respect to the reference CaTi$_{0.8}$Fe$_{0.2}$O$_{3.6}$ (T1), although the total conductivity is not improved. Therefore, further studies on this composition have been carried out in order to determine the role of the ionic carriers and possible effects of the Mg on the overall properties of CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3.6}$.

Characterization of CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3.6}$ (CTFM)

XRD patterns of initial CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3.6}$ (CTFM) calcined at 1200 ºC show perovskite structure with only orthorhombic distorted symmetry depicted in Figure 7.3 (a=7.65 (1) Å, b=5.419 (6) Å, c=5.30 (6) Å). The obtained patterns indicate that a solid solution is formed and the dopant elements are mainly dissolved in the lattice of CaTiO$_3$, since no additional phases were observed. Goldschmidt tolerance factor, and octahedral factor were found to be 0.96 and 0.432 respectively, optimum values to form a perovskite. Figure 7.3b depicts the SEM picture of the membrane, where a closed packing is observed. The grain size is estimated to be in the range of 3 to 6 µm. Based on the literature optimization of the microstructure of CaTi$_{0.8}$Fe$_{0.2}$O$_{3.6}$ (CTF), these large grains may favor the ionic conductivity as the grain boundary becomes negligible. [17] Besides the large grains, small grains with a cubic shape are glimpsed in the SEM picture despite XRD does not clearly indicate a secondary phase. They could be related to the segregation of any of the elements on a minor separated phase, although the XRD or EDS do not give any concluding proof on their nature.
Ionic transport improvement on CaTiO$_3$ based perovskites

Figure 7.3. a) Room temperature XRD pattern of orthorhombic CTFM calcined at 1200 °C and b) SEM picture of the formed membrane.

Figure 7.4. TG analysis under 5 % CO$_2$/air of a) CTFM and CTF and b) BSCF. Heating rate was 10 °C·min$^{-1}$. 
Figure 7.4a shows the TG analysis of CTFM and CTF materials in CO$_2$ containing atmosphere. The analysis of the BSCF is also depicted in Figure 7.4b for comparison. The BSCF gain of mass around 580 °C evidences the formation of carbonates and the later decomposition when temperature keeps increasing. [20, 21] On the other hand, CTFM and CTF do not undergo any visible CO$_2$ uptake under CO$_2$ containing atmospheres to the limits of this technique. The observed weight loss is attributed to the sample dehydration up to 200 °C and latter oxygen release from the perovskite lattice and subsequent oxygen vacancy formation.

Figure 7.5. a) DTG in air, b) TPD of oxygen, m/z=32, in He (pO$_2$=10$^{-5}$ atm), c) cell parameters extracted from HT-XRD in air and d) cell parameters extracted from HT-XRD in N$_2$, for CTFM. Heating rate was 10 °C·min$^{-1}$. 
Figure 7.5 confirms the oxygen release both in high (air) and low ($10^{-5}$ atm) oxygen partial pressures. Figure 7.5a plots the differential thermogravimetry (DTG) in air of CTFM. Two maxima in oxygen desorption are observed at 361 and 557 °C. The starting desorption temperatures (220 and 453 °C) correspond to changes in slope observed in Figure 7.5c, which depicts the evolution of the cell parameters with increasing temperature extracted from HT-XRD. CTFM undergoes chemical expansion as oxygen desorbs and the cations are reduced (larger ionic radius), which occurs at different temperatures for each lattice constant. Figure 7.5b plots mass spectrometry results on TPD of CTFM performed in He, which settles that oxygen is the released specie (both 16 and 32 a.m.u.). Under low $pO_2$ (e.g. He or N$_2$), lattice parameters evolve differently than in air, as lattice parameter, c, is shortened.

Figure 7.6 shows the conductivity, $\ln(\sigma T)$ of CTFM as a function of the reverse temperature measured at 0.21 atm. In the preliminary screening, a change in behaviour is glimpsed at temperatures close to 800 °C. Thus, the temperature range is extended from 400 to 1000 °C and compared with the reference CTF. The addition of dopants to the perovskite results in an increase of the total conductivity with respect to the non-doped CaTiO$_3$. The equilibrium of this oxide in air favours the loss of oxygen and subsequent formation of oxygen vacancies under increasing temperature, which explains the $E_a$ changes. Below 800 °C, total conductivity of CTFM is slightly lower than the corresponding to CTF, which has been reported to have maximum conductivity on the 20% Fe dopant concentration as a consequence of the balance between moderate oxygen vacancy concentration and a relatively disordered Fe$^{3+}$ structure. [13]

On the contrary, above 800 °C, CTFM exceeds the conductivity observed for CTF. This behaviour can be explained by the trapping of the
oxygen vacancies into ordered microdomains that avoid the mobility in CTFM. [13] By increasing the temperature, the large energy required to break down the pairs of these associated defects is achieved and the conductivity of CTFM increases over that of CTF.

Figure 7.6. a) Arrhenius plot of the conductivity of CTFM, CTF and CaTiO$_3$ and b) logarithm of the total conductivity versus $pO_2$ of CTFM and CTF at 1000, 800 and 400 ºC.

The dependence of conductivity on the oxygen partial pressure for CTFM and CTF is presented in Figure 7.6b. The drop in conductivity as $pO_2$ decreases indicates $p$-type electronic conductor behaviour for both CTFM and CTF. The ionic character is enhanced as temperature increases, as can be observed from the weaker dependence on the $pO_2$. The results confirm the mixed ionic-electronic conductivity of these compositions. The high temperature bulk transport properties comprise a major ionic conductivity while electronic contribution becomes more important as the temperature drops. Particularly, at 400 ºC, over the whole range of $pO_2$, CTFM shows 1/2 dependency. This unexpected behaviour is attributed to the coupling of
reduction processes (as inferred from TPD results). At this temperature the formed oxygen vacancies have low mobility and may be trapped as associates ($V_0^\cdot \cdot \cdot \cdot Mg^{\cdot\cdot\cdot}_{Ti}$). In such a case both the $p$-type electronic and ionic conductivities will be decreased providing the observed dependence. At 1000 °C the $pO_2$ dependency is 1/6 and a plateau starts to appear at $pO_2$ lower than $10^{-3}$ atm, characteristic of ionic transport. This plateau should be related with the steady vacancy concentration independent on the $pO_2$ and $\sigma \propto [V_0^\cdot\cdot\cdot]$, and indicates the higher ionic conductivity of the CTFM over the CTF.

On the other hand, reported studies on the oxygen permeability of CaTi$_{1-x}$Fe$_x$O$_{3-\delta}$ dense ceramic membranes with different thicknesses demonstrate significant oxygen surface exchange limitations. [22] The study of the surface properties of CTFM is of interest to evaluate the suitability as oxygen permeation membrane and catalyst. First, the composition of the surface has been studied by XPS. Also the oxygen exchange rate has been measured by PIE and compared with that of CTF.

![Figure 7.7. Ca 2p spectrum of CTFM quenched from 800 °C in Ar. Pass energy was 30 eV.](image-url)
Figure 7.7 shows the deconvoluted Ca 2p XPS spectrum of CTFM quenched from 800 °C in Ar. The Ca 2p band-like spectrum exhibits two features: the 'bulk' one as Ca$^{2+}$ oxidation state in oxide at 347.1 eV (Ca 2p$_{1/2}$) and 350.7 eV (Ca 2p$_{3/2}$) binding energies and a 'surface' feature for Ca$^{2+}$ bonded in the perovkite structure (345.5 eV + 348.1 eV). The last two small peaks are the characteristic plasmon features in the Ca 2p XPS spectrum. [23]

Figure 7.8. Ti 2p spectrum of CTFM quenched from 800 °C in Ar and deconvolution of the spectrum. Ti$^{4+}$ and Ti$^{3+}$ components are highlighted. Pass energy was 30 eV.

Figure 7.8 shows the fitting results of the XPS spectrum of Ti 2p for CTFM. According to characteristic BEs and the split-orbit parameter ($\Delta \sim$6 eV), the Ti 2p main oxidation state is Ti$^{3+}$. A percentage of surface Ti$^{4+}$ was highlighted after deconvolution (~17%), whose peaks appear at binding energies of 458.4 eV (Ti2p$_{1/2}$) and 464.1 eV(Ti2p$_{3/2}$). [23, 24]
Figure 7.9. a) Fe 2p XPS spectrum and b) Mg 2s and Fe3s XPS spectra of CTFM quenched from 800 °C in Ar. Pass energy was 30 eV.

Figure 7.9a shows the XPS spectrum of Fe 2p. According to the BEs and to the presence of the characteristic satellite at 719.2 eV, the oxidation state is identified as Fe$^{3+}$. However, the presence of FeOOH confined on the top of the surface cannot be completely ruled out. [25, 26] The XPS spectrum of Mg 2s for CTFM is shown in Figure 7.9b. The peak observed at 89.3 eV belongs to Mg$^{2+}$ in a non-stoichiometric environment. A second peak in the Mg2s deconvoluted spectrum is observed at 92.5 eV. It corresponds to Fe3s component and confirms the Fe$^{3+}$ oxidation state. [27, 28]

Figure 7.10. O1s XPS spectra of CTFM quenched from 800 °C. Pass energy was 30 eV.
The XPS spectrum of Figure 7.10 shows O1s peaks at binding energies of 528.5 eV, 530.2 eV and 531.5 eV, which are attributed to the chemical state $O^{2-}$ bonded in the lattice. The last two components at 532.9 eV and 534.3 eV can be assigned to strongly adsorbed oxygen as $O^-$, $\cdot OH$, or $H_2O$ (the surface is hygroscopic).

Thus, the XPS demonstrates the presence of oxygen vacancies in the CTFM surface, from the tri- and divalent oxidation states of Ti, Fe and Mg, respectively. These surface oxygen vacancies are expected to provide adsorption sites for the oxygen exchange.

The interaction between gaseous $O_2$ and the oxide surface may be evaluated by pulse isotope exchange (PIE). The overall oxygen exchange reaction in the surface can be expressed as,

$$O_2 + 2V_O^0 + 4e^- \rightarrow 2O^X_0$$

which indicates that both lattice oxygen vacancies and electronic carriers at the oxide surface participate in the rate of surface exchange. Figure 7.11a plots the PIE results of oxygen surface exchange rate, $\Re_o$, of CTF and CTFM obtained by PIE at $pO_2 = 0.21$ atm. Up to 650 ºC $\Re_o$ is the same for both materials, so that a similar surface limitations are expected on the oxygen permeation. Below 550 ºC an abrupt drop in CTFM $\Re_o$ is observed, which is attributed to the decrease of free oxygen vacancies as temperature decreases, also observed by conductivity and TPD. The range of temperature at which PIE measurements could be performed limits the direct comparison, as the measurement is enabled as long as the effective time constant associated with the isotopic exchange is of similar order as the time scale of the experiment, i.e. the residence time of the packed bed reactor. Thus, higher temperature experiments were not possible.
Figure 7.11b shows the temperature dependence of the rates of dissociative adsorption $\mathcal{R}_a$, and oxygen incorporation $\mathcal{R}_i$, of CTFM. The overall rate of surface exchange appears to be controlled by the dissociative adsorption reaction, as the rate of oxygen incorporation is found between one or two orders of magnitude higher than the adsorption one, similar to reported studies of BSCF and Bi$_{0.75}$Er$_{0.25}$O$_{2-\delta}$. \[29, 30\]

Figure 7.11. Arrhenius plot of CTF and CTFM (a) the overall surface exchange rate $\mathcal{R}_o$ and (b) partial $\mathcal{R}_a$, $\mathcal{R}_i$ and $\mathcal{R}_o$.

Figure 7.12 collects the Arrhenius plots of the overall oxygen surface exchange rate, $\mathcal{R}_o$ of some reference oxide materials measured by PIE (YSZ, CGO10, CGO20, CT50) or isotopic exchange followed by deep profiling (IEDP) as La$_{0.8}$Sr$_{0.2}$CoO$_{2-\delta}$ (LSC20) and BSCF. CTF and CTFM show a comparatively high surface exchange rate with regard to YSZ and CGO10, known as solid pure ionic conductors, and LSC20 perovskite. $\mathcal{R}_o$ is comparable to that of CGO20, CT50. On the other hand CTF and CTFM $\mathcal{R}_o$ are lower than BSCF. \[30-32\] The limits in oxygen surface rate may be less significant by applying a catalytic layer over the membrane surface.
Figure 7.12. Comparative Arrhenius plot of the CTFM and CTF $\mathcal{R}_0$ with literature data of widely used oxides measured by PIE as YSZ, CGO10, CGO20, CT50 or IEDP as LSC20 and BSCF. [30-32]

Oxygen permeation test

Permeation measurements have been performed on a CTFM membrane 900 $\mu$m thick. Both disc sides are coated by a Pt ink 20 $\mu$m layer aiming to improve surface catalytic behaviour. Oxygen is separated from synthetic air flow using argon as sweep gas. The permeation fluxes (mL·min$^{-1}$·cm$^{-2}$) are calculated by dividing the permeation rates by the effective surface area of the membranes.

First, oxygen permeation flux is represented in Figure 7.13 as a function of reciprocal temperature using 100 mL·min$^{-1}$ of synthetic air and pure oxygen as feed and 400 mL·min$^{-1}$ Ar as sweep flow. It can be observed that in both feeding conditions, oxygen flux increases with temperature following Arrhenius behaviour. The oxygen permeation flux increases 2-fold by using oxygen as
feed gas, and reaches values up to 2 mL·min⁻¹·cm⁻² at 1000 ºC. This increase is a consequence of the higher driving force, which corresponds to the oxygen chemical potential gradient along the thickness of the membrane. Based on the magnitude of the membrane thickness (0.9 mm), which is over the characteristic length expected for perovskites and on the catalytic activation of the surface achieved by depositing a Pt porous layer, the bulk diffusion is expected to limit the oxygen permeation rate. [4, 33] In such a case, the relationship between the oxygen potential gradient and the oxygen flux is postulated by the Wagner equation:

\[
J(O_2) = \frac{RT}{16F^2L} \int_{pO_2^1}^{pO_2^2} \sigma_{amb}(pO_2) d\ln pO_2
\]  
(Eq. 7.3)

where \(J(O_2)\) is the oxygen permeation rate, \(R\) and \(F\) are gas and Faraday constants, \(L\) is the membrane thickness, \(\sigma_{amb}\) is the ambipolar conductivity, and \(pO_2^1\) and \(pO_2^2\) are the oxygen partial pressure at the high pressure side and the low pressure side, respectively.

Constant apparent activation energies for oxygen transport are observed in the whole temperature range (1000-400 ºC), regardless of \(pO_2\) in the feed stream. This suggests that the ion diffusion through the bulk and grain boundary is the rate limiting step. Values of the apparent \(E_a\) are 1.05 eV (57.58 kJ/mol) and 0.96 eV (43.89 kJ/mol) for air and pure oxygen feed, respectively. Obtained \(E_a\) values are in the order of those obtained for thick BSCF membranes at high temperatures, where oxygen permeation process is limited by the bulk oxygen ions diffusion. [34] Otherwise, the \(E_a\) for oxygen surface exchange is expected to be higher than that for the oxygen ion diffusion. [4, 7] The porous Pt layer on the surface enhances the catalytic activity and the oxygen uptake due to the increase of the available area for the surface exchange reactions. Thus, this
surface modification avoids the surface limitation leading to an improvement of surface exchange rate and the subsequent increase in the $J(O_2)$. \[11, 35\]

Figure 7.13. Oxygen permeation flow for CTFM membrane as a function of the temperature, with a sweep flow of 400 mL·min$^{-1}$ of Ar. Synthetic air and pure oxygen at a flow rate of 100 mL·min$^{-1}$ are used as feed gas.

Figure 7.14a shows the oxygen permeation fluxes in the temperature range of 700-1000 °C for different flow rates of sweep gas and synthetic air used as feed gas. The increase of the gas flow rate causes an increase of the driving force by the reduction of $pO_2$ in the sweep stream, as the permeated oxygen is diluted. As predicted by the Wagner's equation, it leads to an improvement in the oxygen permeation flux. On the other hand, the enhancement can be further produced by the improvement in the fluid dynamics in the permeate chamber that avoids the accumulation of species on the permeate side surface. This allows reducing concentration polarization resistance and decreasing substantially the local $pO_2$ at the membrane surface. \[8\] The influence of the gas flow rates is more pronounced in the high temperature range. It may be the consequence of the improvement in ionic conduction for
CTFM achieved above 800 ºC (increasing the ambipolar conductivity) together with the influence of the higher \( pO_2 \) gradient.

Figure 7.14b depicts the permeation results for pure \( O_2 \) and air as feed gas. An important increase in the \( J(O_2) \) is observed by the growth of the chemical potential across the membrane (provided by the pure \( O_2 \)). Specifically, an oxygen flux of ca. 1.6 \( \text{mL} \cdot \text{min}^{-1} \cdot \text{cm}^{-2} \) is achieved at 1000 ºC by the highest studied \( pO_2 \) gradient.

![Figure 7.14. a) Oxygen permeation flux for CTFM membrane as a function of the Ar sweep flow at three different temperatures using synthetic air as feed and b) using air and \( O_2 \) as feed gas at 1000 ºC.](image)

The oxygen flux obtained for the reference CTF 1 mm membrane is reported to achieve 7.08 \( \times 10^{-9} \) mol s\(^{-1}\) cm\(^{-2}\) at 950 ºC, which uses a CTF porous layer as catalyst on the both sides, and is fed by air. [22] This value is more than one order of magnitude lower than that obtained for CTFM, which was 4.6 \( \times 10^{-7} \) mol s\(^{-1}\) cm\(^{-2}\) at 950 ºC using air as feed gas and 400 mL\( \cdot \)min\(^{-1}\) of Ar as sweep gas. The improvement is ascribed to the enhanced ionic conductivity observed above 800 ºC.
Paying attention to the most probable operating conditions of an oxygen transport membrane, the stability of CTFM against CO\textsubscript{2} has been checked. The sample has been exposed to 15\%CO\textsubscript{2}/Ar atmosphere during 24 hours at 750 °C, since the carbonates formation, if any, would occur at this temperature. After this treatment, it has been cooled down to 650 °C, and the oxygen permeation flow is measured from 650 °C to 1000 °C in the same atmosphere. In Figure 7.15a, it can be observed that the permeation values of heating up and cooling down measurements are slightly different. Since TG of CTFM in 5\% CO\textsubscript{2} containing Ar does not show any carbonation reaction, the flux gap among both processes in CO\textsubscript{2}/Ar sweep side can be ascribed to the CO\textsubscript{2}/O\textsubscript{2} competitive adsorption on the membrane surface. Actually, oxygen flux values obtained after CO\textsubscript{2} treatment using Ar as sweep gas are very similar to those measured before CO\textsubscript{2} treatment (Figure 7.15b). The oxygen flux drop when the membrane is exposed to CO\textsubscript{2} may be produced by a change in the Pt porous layer.

Figure 7.15. a) Oxygen permeation flow after 24 hours in CO\textsubscript{2}/Ar atmosphere. Oxygen flux using Ar as sweep gas is represented using open symbols, b) Oxygen permeation flux in Ar before and after of CO\textsubscript{2} treatment. \(Q_{\text{Ar}}=400 \text{ mL.min}^{-1}\).
Finally, CTFM membrane material was tested in harsh conditions. On one hand, the membrane was exposed to a continuous gas flow composed of 1000 ppm COS, 100 ppm HCN, 4% CO₂, 46% CO, 46% H₂ and 4% H₂O at 35 bar and a thermal treatment up to 400 °C (treatment 1). On the other hand, a treatment of two weeks on a continuous stream of CO₂ containing 5% O₂, 1% of SO₂ and 2.5% of H₂O, was carried out during 14 days at 800 °C (treatment 2).

From the XRD patterns of Figure 7.16, it can be inferred that materials do not suffer any degradation by the treatment 1, and the formation of carbonates, sulfides or sulfates phases is negligible up to the limit of XRD analysis. However, the SO₂ containing treatment 2 causes the reaction of the material and the loss of the perovskite structure. SO₂ preferentially reacted with Ca and Ti. The impurities founded are indexed as CaFeO₂ (ICSD 00-021-0917), CaSO₄.

Figure 7.16. XRD patterns of as sintered CTFM powder and membranes after the stability treatments. Reference pattern and XRD of sample holder has been plotted for comparison.
Note that diffraction peaks corresponding to the sample holder material are also observed.

7.3. Conclusions

The series $\text{Ca}_{1-x}\text{A}_x\text{Ti}_{1-x}(\text{M}1_{1-y}\text{M}2_y)_x\text{O}_{3-\delta}$ has been prepared by solid state reaction to evaluate the formability and ionic conductivity improvement of $\text{CaTiO}_3$ perovskites. Although the Goldschmidt and the octahedral factors anticipated the perovskite structure for the majority of the compounds, they are not experimentally achieved for all the compositions. It has been attributed to different sitting of the dopant or to different oxidation states than expected (so that expected ionic radii), but no more research has been developed yet on this regard.

DC conductivity indicates that all the dopants allow increasing the conductivity and particularly CTF and CTFM, which have been further characterized. DC conductivity up to 1000 ºC demonstrates an increase of conductivity of CTFM over that of CTF upon 800 ºC, which has been ascribed to the mobility of associated oxygen vacancies released at high temperature.

XPS shows the presence of $\text{Fe}^{2+}$ and $\text{Fe}^{3+}$, and $\text{Mg}^{2+}$ in CTFM surface, while PIE results do not show any surface exchange rate improvement with respect to CTF up to 750 ºC, so that the CTFM membrane is covered with Pt ink layer to avoid surface restrictions. The systematic study of the CTFM membrane variables on the oxygen flux and temperature delivers an oxygen permeation flux one order of magnitude higher than referenced CTF membranes, which is attributed to the temperature dependent increase of oxygen vacancies concentration and mobility. Moreover, membrane stability is
Ionic transport improvement on CaTiO$_3$ based perovskites proved under CO$_2$ as sweep gas, and after being submitted to harsh 1000 ppm COS, 100 ppm HCN, 4% CO$_2$, 46% CO, 46% H$_2$ and 4% H$_2$O at 35 bar environment up to 400 °C. However, the material is not stable against wet SO$_2$.

7.4. References


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Ionic transport improvement on CaTiO$_3$ based perovskites


Chapter 8. Ceria-based stable dual-phase membranes for fast oxygen separation

Abstract

As another approach of the new materials research, dual phase materials composed of spinel and fluorite phases have been developed. They have been selected on the basis of their electrical conductivities and the expected contributions to the overall composite material to enhance the mixed ionic and electronic conductivity. Fe$_2$NiO$_4$ is the spinel of choice as it is prone to form dense composite without reacting with the fluorites.

60 vol% Fe$_2$NiO$_4$ - 40 vol% Ce$_{0.8}$Ln$_{0.2}$O$_{2+\delta}$ (FNO-CLnO) composites have been successfully obtained by one-pot fabrication method showing both spinel and fluorite pure phases. Narrow grain size distribution centred around 1 $\mu$m and homogeneous distribution of grains is attained in the gastight membrane, as well as percolative pathways from side to side of the dual-phase membranes. Specifically, composite membranes with enhanced oxygen permeability and unprecedented stability in oxyfuel-like gas environments are reported. The conductivity of the composite is investigated as a function of temperature and $pO_2$. Oxygen separation in this kind of FNO-CLnO composite membranes occurs via the separate ambipolar transport through the two distinct percolating
networks. Further study of a 60 vol% Fe$_2$NiO$_4$ - 40 vol% Ce$_{0.8}$Tb$_{0.2}$O$_{2.8}$ (FNO-CTO) composite membrane has been developed. Oxygen permeation flux values of 0.17 mL·min$^{-1}$·cm$^{-2}$ and 0.20 mL·min$^{-1}$·cm$^{-2}$ are achieved at 1000 °C when Ar and pure CO$_2$ are used as sweep gas, respectively, through a 0.68 mm-thick membrane. Long term experiments at 900 °C show that the material is stable and effective in pure CO$_2$ atmospheres and the oxygen permeation is even improved after 76 h on CO$_2$ stream. Moreover, the composite resists a stability test in SO$_2$ and CO$_2$ containing gas at 800 °C for 170 h, which represents a step forward towards its potential use in oxyfuel power plants.
8.1. Introduction

Materials with perovskite structure are the MIEC materials showing the highest electrical conductivities but permeabilities are often limited by the ionic conductivity. Doping strategies to increase the oxygen ion conductivity normally affect the crystal and thermo-mechanical stability and increase their vulnerability when exposed to large oxygen concentration gradients and atmospheres containing CO₂, SO₂ and H₂O, the most probable operation conditions in the oxyfuel process and catalytic membrane reactors in different intensified industrial processes. [1-3]

In previous chapters, the difficulty of joining all the desired characteristics in a single material has been recognized, which points out to achieve them separately. Dual phase composite membranes try to combine the best characteristics of different compounds to achieve large oxygen permeability and relatively good chemical and mechanical stability at elevated temperatures. Thus, composite materials should consist of an electron conducting material, which allows the percolation of electrons, and an ionic conductor that transports the oxygen ions through the membrane. Due to the existence of two phases, the influence of the grain boundary between them plays an important role, since it can either promote or block the transport of ions and electrons across the membrane. Furthermore, large catalytic activity towards surface oxygen exchange is required.
The first reported dual phase membranes were ceramic-metal (cermet) composites consisting of a continuous oxygen ion-conducting oxide phase and a continuous electron-conducting noble metal phase, but the high costs arising from the use of noble metals prevent the industrial applications. [4-6] Therefore, attempts to decrease the total price of the membranes involve the substitution of the metallic phase by solid oxides with high electronic conductivity. Indeed, the use of ceramic oxide composites also brings the possibility of substituting the electronic or ionic phase by a MIEC phase that does not block the ionic or electronic transport. Ceramic–ceramic (cercer) dual-phase compounds have been successfully prepared since early 2000’s decade. [7] In a cercer, other important requirements besides the transport properties have to be taken into account, i.e. the stability of both phases against reaction with each other and the compatibility of thermal expansion coefficient (TEC) of both compounds. Following this composite strategy, several crystalline structures have been combined, e.g. fluorites, perovskites, spinels and rock salts. [8] Materials with perovskite and spinel (A$_2$BO$_4$) structure may be used as electronic phase. [9] Fluorite materials such as yttria stabilized zirconia (YSZ), stabilized bismuth oxide (SBO) or Ce$_{1-x}$B$_x$O$_{2+\delta}$ (B=lanthanide) play the oxygen conductor role in composites since they present mainly ionic conductivity. Moreover, the chemical stability of doped cerias is significantly higher than that of perovskites. [10]

On these premises, cercers composed of doped ceria fluorites and several spinels have allowed the development of CO$_2$ stable composite membranes. First suggested membranes joined Ce$_{0.8}$Gd$_{0.2}$O$_{1.9}$ (CGO20) with CoFe$_2$O$_4$ (CFO) and MnFe$_2$O$_4$ (MFO). [11] An optimum ratio for the percolation of both phases was found to be 30% volume of spinel phase. However, these compositions presented reaction between phases and showed a small amount of GdFeO$_3$, which reduces the Gd content in CGO20 and
consequently drops the ionic conductivity. Lower fractions of spinel (ca. 17%) are also oxygen permeating despite percolation of the electronic phase was not achieved. [11] To further improve the oxygen permeation properties, the same authors studied the composite formed by 85 vol% of Ce$_{0.8}$Sm$_{0.2}$O$_{1.9}$ (SDC20) and 15 vol% of Mn$_{(1.5-0.5y)}$Co$_{(1+0.5y)}$Ni$_{0.5}$O$_4$ (MCNO, 0$\leq$y$\leq$1) spinel oxide. The microstructural analysis found out that two spinel type oxides are formed, one consisted of Mn-Co-Ni and another of Co-Ni. The highest oxygen flux density was obtained when y=1, which is comparable to those for SDC–15MFO. [12]

Figure 8.1. Oxygen and electron transport scheme in a dual phase composite membrane composed of a) pure electronic and mixed ionic electronic conducting phases and b) pure electronic and ionic conducting phases.

Co-free spinel Fe$_2$NiO$_4$ (FNO) 40 wt% was mixed with Ce$_{0.9}$Gd$_{0.1}$O$_{1.95}$ (CGO10) both by powder mixing method and one pot. [9] No intermixing of cations between the two phases occurs while the dual phase structure at high temperatures is retained even in 50% CO$_2$ containing atmospheres. The enhancement of the oxygen flux was obtained by the one pot method due to the improvement in morphology and homogeneity. [9, 13] In fact, oxygen flux improves with decreasing grain sizes in composite materials while high density
is needed to avoid leakages. Therefore, sintering temperatures should be optimized for each composite composition. 20% Pr-doped CeO₂ (CPO) fluorite possesses redox properties that bring about both ionic and electronic conductivities, which is favorable for oxygen permeable ceramics. Oxygen permeation through Ce₁₋ₓPrₓO₂₋₅ membranes has been obtained under small pO₂ gradients, but the large expansion under reducing atmospheres such as methane causes severe strains in the membrane up to the final fail. [14, 15] CPO/ 40 wt% MFO composite has demonstrated several benefits over the single fluorite. [16, 17] From the fact that CPO is a MIEC conductor arises that even non percolative fractions of spinel phase tolerate the oxygen permeation. Although a small amount of PrFeO₃ is formed when MFO is over 10%, the optimum volume fraction of MFO was determined to be from 5 to 25%. [16]

The present chapter reports the research on dual phase membranes composed of the spinel and fluorite phases listed in Table 8.1. They have been selected on the bases of their electrical conductivities and the expected contributions to the overall composite material.

<table>
<thead>
<tr>
<th>Spinel</th>
<th>Fluorite</th>
<th>Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe₂NiO₄</td>
<td>Ce₀₈GdB₀₃O₂₋₅</td>
<td>Electronic + ionic</td>
</tr>
<tr>
<td>Mn₁CuO₄</td>
<td>Ce₀₈GdB₀₁Pr₀₁O₂₋₅</td>
<td>Electronic + mixed ionic-electronic</td>
</tr>
<tr>
<td>Fe₂CrO₄</td>
<td>Ce₀₈Tb₀₂O₂₋₅</td>
<td>Electronic + mixed mainly ionic</td>
</tr>
<tr>
<td>Co₂FeO₄</td>
<td>Ce₀₈Tb₀₂O₂₋₅+Co</td>
<td>Electronic + ionic-electronic + sintering aid</td>
</tr>
</tbody>
</table>

Table 8.1. Spinel and fluorite structured compounds selected for dual-phase membranes.

An optimization of phase ratio and synthesis method is carried out, as well as a screening of several materials combinations to develop dual-phase membranes. X-Ray Diffraction (XRD) and Scanning Electron Microscopy (SEM) carry out characterization of powders and sintered membranes. The
electrochemical study is performed by 4-point DC conductivity. The oxygen permeation of a planar disk dense membrane with both sides covered with a previously optimized catalytic composite layer has been tested by Ar and CO₂ sweeping gases. Finally, the stability of the material for 7 days against exposure to a 1 mol% SO₂-containing wet gas environment is checked.

8.2. Results and discussion

Component and microstructural optimization

In a first step, the synthesis of the spinels was checked separately. They were prepared by Pechini reaction method and sintered at 1300 °C, which is a normal temperature to obtain dense membranes. Initial identification of the crystalline phases by XRD is shown in Figure 8.2. Only Fe₂NiO₄ and Fe₂CrO₄ show pure spinel structure. Thus Mn₂CuO₄ and Co₂FeO₄ were not further considered. Mn₂CuO₄ was modified to Mn₁.₅Cu₁.₅O₄ and formed pure spinel phase. However, even though both Mn₁.₅Cu₁.₅O₄ and Fe₂CrO₄ form spinel and a stable composite with fluorites, they were not able to deliver dense monoliths and were not further considered, either.
Figure 8.2. XRD patterns of tested spinels: \( \text{Fe}_2\text{NiO}_4 \), \( \text{Cr}_2\text{FeO}_4 \), \( \text{Co}_2\text{FeO}_4 \), \( \text{Mn}_2\text{CuO}_4 \). Grey circles indicate impurity peaks.

In order to optimize the microstructure, the composite powder has been prepared by powder mixing method (PM) and one pot method (OP). The first method blends the oxides separately, i.e., the fluorite compounds formed by co-precipitation method and incipient wetness impregnation (if Co oxide is added) and the spinels prepared by Pechini, both calcined at 1300 °C. Otherwise, the OP method allows the composite to be synthesized by a single step Pechini method. [9, 18] Figure 8.3a shows the XRD pattern of 50\%\( \text{Fe}_2\text{NiO}_4 \)-50\%\( \text{Co}_{0.8}\text{Gd}_{0.2}\text{O}_{1.95} \) (FNO-CGO) sintered in air at 1400 °C. For comparison, separate CGO and FNO structure peaks are also plotted. Both structural phases are well identified in the composites with no evidence of impurities or reaction between them up to the limit of the XRD device, demonstrating the validity of both fabrication methods.
Figure 8.3. a) XRD patterns of the composites prepared by SS and OP, and the Fe$_2$NiO$_4$ and Ce$_{0.8}$Gd$_{0.2}$O$_{1.9}$ single phases. b) Arrhenius plot of total electrical conductivity for the 50/50 phase ratio composites prepared by SS and OP and for 60/40 phase ratio OP composite. c) SEM picture of 50/50 FNO-CGO prepared by PM and d) SEM picture of 50/50 FNO-CGO prepared by OP.

Figure 8.3c and d show the backscattered SEM images of the 50%Fe$_2$NiO$_4$-50%Ce$_{0.8}$Gd$_{0.2}$O$_{1.9}$ composite membranes prepared by PM and OP methods, respectively. While the specimens blended by PM form agglomerates of spinel phase (dark grey), the OP method enables the formation of homogeneous-nanometric sized grains in a well-distributed mixture of the two phases forming the composite. On the other hand, the lack of continuity of spinel phase observed in SEM pictures suggests that percolative pathways for electrons may be not completely achieved.
To confirm the importance of materials uniformity, Arrhenius plot of the total electrical conductivity in air for the differently sintered materials is shown in Figure 8.3b. Conductivity increases the more uniform grain size and homogeneous mixture. Besides, increasing the spinel content from 50 to 60% provides a continuous pathway for electrons across the bulk material. Therefore, 60 vol.% of electronic FNO and 40 vol.% of doped ceria ionic fluorite phases synthesized via OP method have been selected since this rate allows the electronic percolation and a proper microstructure.

![Figure 8.4](image)

Figure 8.4. XRD patterns of FNO-CLnO synthesized by OP and sintered in air at 1400 °C. Reference fluorite and spinel structure peaks are also indicated (ICDD 00-043-1002 and 00-044-1485, respectively).

Figure 8.4 shows the XRD patterns of FNO-CLnO synthesized by OP and sintered in air at 1400 °C. For comparison, reference fluorite and spinel structure peaks are also indicated (ICDD 00-043-1002 and 00-044-1485, respectively). Both structural phases are well identified in FNO-CGO,
FNO-CTO and FNO-CTOCo composites with no evidence of impurities or reaction between them up to the limit of the XRD device. Otherwise, FNO-CPGO has another phase identified as Pr$_7$O$_{11}$. This impurity is detrimental for the conductivity as it may block the diffusion of ions and electrons. Therefore, this composition is not further considered.

The conductivity in air of stable FNO-CLnO series is compared in Figure 8.5 as a function of temperature. At high temperatures, the FNO-CGO is the compound with higher conductivity, ascribed to the ionic conductivity given by the higher extrinsic nonstoichiometry of CGO. At low temperatures FNO-CTO shows an enhanced conductivity over that of FNO-CGO, which can be attributed to the improved electrical conductivity of the fluorite phase containing multivalent dopants. CTOCo showed the lowest conductivity, probably due to the increased particle size induced by the Co.

*Fe$_2$NiO$_4$ - Ce$_{0.8}$Tb$_{0.2}$O$_{2.4}$ composite characterization*

In the following, an exhaustive study on FNO-CTO composite material is carried out. It is compared in some points with with FNO-CGO. Different conduction mechanisms are expected from the CGO and CTO fluorite phases since CGO is an ionic conductor at high $pO_2$ and the CTO may provide electronic conduction, as deduced in Chapter 5.
Figure 8.5. Arrhenius plot of the total electrical conductivity in air of FNO-CLnO composite series.

Dense membrane and probes of FNO-CTO have been achieved by sintering the composite at 1400 °C. XRD pattern of FNO-CTO shown in Figure 8.4 indicate that pure spinel and cubic fluorite phases are obtained. The calculated cell parameters of FNO and CTO are \( a = 8.332 \, \text{Å} \) and \( a = 5.406 \, \text{Å} \). They are very similar to those obtained for single phases separately (\( a = 8.339 \, \text{Å} \) and \( a = 5.392 \, \text{Å} \), respectively), which point out that there is not inter-diffusion of the elements. Despite these differences, diffusion of Ce or Tb inside the spinel structure (if any) should cause an expansion of the cell parameter, but the experimental data shows a slight shrinkage. On the other hand, the cell parameter of the ceria, which is longer than that observed previously for the single phase, could suggest that Tb dopant is not fully solved in the fluorite lattice, so that the rest should appear as \( \text{Tb}_2\text{O}_3 \) impurity. However, \( \text{Tb}_2\text{O}_3 \) is not detected by XRD, up to the limits of the device. The composite has been calcined in atmospheric air. Under such condition not all the Tb is introduced as \( \text{Tb}^{3+} \) (i.r. 1.04 Å) but as \( \text{Tb}^{4+} \) (i.r. 0.88 Å), as studied in chapters 3 and 5. Both the sample thermal history and the influence of the spinel on the Tb reduction
properties may cause the slight difference in cell parameter by delivering higher ratio $\text{Tb}^{3+}/\text{Tb}^{4+}$. [19]

Figure 8.6. a) BS-SEM picture of the fractured cross-section of a FNO-CTO composite membrane sintered at 1400 °C in air; b) detail of the grain boundary between spinel (dark) and fluorite (light) phases; c) detail of two fluorite grains fringe; d) detail of two spinel grains fringe and; e) CGO-LSM-Pd catalytic layer on the composite.
SEM images allow the analysis of the membrane microstructure, e.g. grain size distribution, pores, grain boundaries as well as offering information about the possible solid state reaction between the two composite phases. SEM backscattered electrons detector (BS-SEM) provide images with compositional contrast related to the averaged atomic number of the elements of each phase, and therefore changes in composition of the different grains and element distribution can be distinguished. The images in Figure 8.6 show a dark phase that corresponds to the spinel and a bright one that belongs to the fluorite. In Figure 8.6a, a narrow grain size distribution (centred around 1 µm) and a homogeneous distribution of grains can be observed. The existence of percolative paths from side to side of the dual-phase membranes necessary for the oxygen permeation is ascertained. Image analysis confirms the expected 60/40-volume phase ratio (spinel/fluorite). Figure 8.6b presents a higher magnification image offering details about the grain boundaries between the two different phases, and apparently no phase reaction, impurities segregation or inter-diffusion take place during the membrane sintering at 1400 ºC. Further, Figure 8.6b-d reveal the very good bounding between grains made of the same phase (CTO-CTO or FNO-FNO) or different phases (FNO-CTO). Additionally, Figure 8.6e shows a detail of the 18 µm-thick CGO-LSM catalytic layer, which can be easily identified by its smaller grain size and higher porosity. The BS-SEM image (Figure 8.6e) also makes visible the two distinct phases of this catalytic layer, i.e., CGO corresponds to the bright phase and the LSM to the dark one. From this image, it is also visible a very clean interface between the membrane and the catalytic layer, with no reaction interfaces or cracks, which reveals the chemical compatibility of all the different materials and the good bounding between layer and membrane.
Figure 8.7. a) Total electrical conductivity recorded in air as a function of temperature for the single phase CTO, FNO and for the composite, b) Arrhenius plot of the total electrical conductivity of the composite in air and Ar atmospheres and c) total electrical conductivity at 800 °C as a function of the $pO_2$.

Once proved the compatibility between both crystalline phases and the appropriate microstructure of the sintered composite body, the transport properties are investigated. Due to the selected percentage of phases in the composite, preferential pathways for oxygen and electrons are expected, so that the total conductivity would result from the sum of CTO and FNO contributions. The plot in Figure 8.7a shows the electrical conductivity recorded in air as a function of temperature for the single phase CTO, FNO and for the composite. The composite conductivity is surprisingly higher than the equivalent sum of the single components, which is attributed to the improvement of densification for the composite in contrast to the FNO, with lower sintering activity. FNO sample displayed in Figure 8.8 is not totally dense.
(relative density ~90%) when sintered at 1400 °C for 10 h, while CTO and the composite are dense at the same conditions.

Figure 8.7. SEM image of Fe2NiO4 sample sintered at 1400 °C in air dwelling 10 h. ~90% relative density was achieved at these conditions.

The Figure 8.7b displays the total conductivity, ln(σT) as a function of reciprocal temperature measured in air and argon (pO2 =0.21 and 5x10^{-5} atm, respectively) for the composite. An apparent change in activation energy (E_a) is observed in air, which suggests a change in the dominant conduction mechanisms. E_a values of 0.45 and 0.71 eV have been calculated from the fitted data below and above 655 °C, respectively. The low temperature E_a may correspond to FNO phase and is characteristic for the hopping of electronic charge carriers, i.e., hopping of localized electrons between multivalent cations located in the octahedral Fe sublattice. [20]

\[ \text{Fe}^{3+} + e^- \rightarrow \text{Fe}^{2+} \]  \hspace{1cm} (Eq. 8.1)

For temperatures above 655 °C, oxygen is released from CTO phase (oxygen vacancy formation) and induce Tb^{4+} reduction to Tb^{3+}, which is charge compensated by the formation of electrons localized on Tb ions, as indicated by the following equations:
\[
\frac{1}{2} \text{O}_2 (g) + V_0^– \rightarrow O_0^{\times} + 2h^–; \quad [V_0^{\times}]^{-1} p^2 p\text{O}_2^{-1/2} = K_0(T) \quad (\text{Eq. 8.2})
\]

\[
\text{Tb}^{\times}_{ce} + e^– \rightarrow \text{Tb}^{\times}_{ce} \quad \frac{[\text{Tb}^{\times}_{ce}]}{[\text{Tb}^{\times}_{ce}]^n} = K_{\text{Tb}}(T) \quad (\text{Eq. 8.3})
\]

The oxygen vacancies concentration in combination with the strong thermal activation of the oxide ion mobility leads to the noticeable increase in the ionic conductivity. As a result, the contribution of the ionic transport to the total conductivity of the FNO-CTO composite becomes significant at temperatures above 655 ºC. This thermal process in CTO is responsible for the raise in the overall composite \(E_a\), which is related to the higher \(E_a\) associated to oxygen vacancy diffusion with regard to the electron hopping \(E_a\) observed at lower temperatures. \[21, 22\]

Figure 8.7b also presents the thermal evolution of the composite conductivity in Ar atmosphere upon equilibration of the samples in argon at 800 ºC for 2 h. At this low \(p\text{O}_2\) terbium and iron cations are mainly reduced to \(\text{Tb}^{3+}\), and \(\text{Fe}^{2+}\) valence state along the whole studied temperature range. Hence, in Ar, the concentration of oxygen vacancies in CTO remains constant with decreasing temperatures. Consequently a monotonic Arrhenius behaviour is observed, \(i.e.,\) the \(E_a\) remains constant. \(E_a\) is 0.38 eV, which suggests that the electronic conductivity of the partially-reduced FNO governs the composite conductivity. Additionally, the conductivity increases substantially at higher temperatures due not only to the further activation of electron hopping rate but also to the increased probability of the localized electrons in the spinel to transfer from one ion to another due to the lattice vibrations. Thus, these trapped electronic carriers in the ferrite may contribute to the conduction process leading to an overall decrease in resistivity.
Total electrical conductivity at 800 °C as a function of the $pO_2$ is depicted in Figure 8.7c. Nickel ferrite is a $p$ or $n$-type semiconductor depending on the oxygen deficiency determined by the sintering temperature and atmosphere. The present sintering conditions, i.e., high temperatures and oxygen partial pressures lower than 4 atm of oxygen, favour the $Fe^{2+}$ concentrations instead of $Ni^{3+}$ holes. [20, 23] Thus, $n$-type conductivity (-1/4 dependency with $pO_2$) is expected. However, the influence of the slight increase in the CTO ionic conductivity (<-1/6 dependency with $pO_2$) provokes the adjustment of conductivity-$pO_2$ dependency of both phases giving rise to -1/11 $pO_2$ dependency, as shown in Figure 8.7c. [20]

**Dual phase Fe$_2$NiO$_4$ - Ce$_{0.8}$Tb$_{0.2}$O$_{2.4}$ membranes**

The preceding results provide sufficient evidences about the participation of both ionic and electronic carriers in the ambipolar conduction. Consequently this stable composite presents the appropriate characteristics to work as oxygen permeation membrane. The catalytic layer consisted of CGO-LSM porous composite (Figure 8.6e) and Pd infiltration is selected to further promote oxygen activation. In chapter 6, this mixture demonstrated to be the optimum between LSM-doped ceria compositions with interesting electrochemical properties for oxygen activation at high temperatures, while these phases are known to be stable and effective in CO$_2$-rich atmospheres. CGO provides the ionic conductivity that LSM lacks and thus allow increasing the global three-phase-boundary (TPB) region of the oxygen activation and incorporating additional redox electrocatalytic activity. Further, no reaction between the layer and the membrane surface is observed.
Figure 8.9. a) Oxygen permeation flux as a function of temperature and sweep gas (Ar or CO₂) for FNO-CTO membrane and in Ar for single phase CTOCo; and b) flux stability with time on stream for FNO-CTO membrane in CO₂ sweeping at 900 °C.

Figure 8.9a shows the oxygen flux as a function of temperature when two different sweep gases are used, i.e., argon and pure CO₂. Using a 0.68 mm thick membrane, permeation fluxes of 0.17 mL·min⁻¹·cm⁻² and 0.20 mL·min⁻¹·cm⁻² for Ar and CO₂ sweeping are obtained at 1000 °C, respectively. These permeation values are twice the permeation values achieved in Chapter 5 using a single phase Co-containing Ceₐ.₈ₖ₂O₂₋₆ (CTOCo) membrane. This result stresses the importance of the electronic contribution provided by the spinel percolating network.

Oxygen permeation observed in Figure 8.9 represents a two-fold Arrhenius behavior with a change in the activation energy at 800 °C. This indicates that the nature of the processes limiting the oxygen permeation rate changes with the temperature. In the high temperature range (1000-800°C), the
bulk diffusion through the dense membrane controls the permeation. In the low
temperature range, the limiting step is the surface exchange reaction.

By using CO₂ as a sweep gas, the oxygen permeation is maintained and
even improved at high temperatures with respect to Ar sweeping. As observed
for broadly applied perovskite materials as BSCF or LSCF, the oxygen
permeation could be affected by either carbonate formation or O₂ competitive
adsorption on the membrane surface. [24-26] These two phenomena would lead
to a reduction in oxygen permeation rates, the first being the most dramatic.
Since terbium doped ceria is not prone to interact with CO₂ to form carbonates,
oxygen permeation diminution observed in Figure 8.9a at temperatures below
900 °C would be related with the process involving O₂-CO₂ competitive
adsorption on reaction sites. This process is more favorable at low temperatures,
as previously observed on several oxygen-ion conducting surfaces. [2, 27]
Figure 8.9a also indicates that above 900 °C, CO₂ utilization as sweep gas
improves the oxygen permeation. This could be related with the better sweeping
properties of CO₂ with respect to argon at these high temperatures and the
smaller CO₂ adsorption constant as temperature is increased, thus neglecting the
effect of competitive adsorption.

Figure 8.9b represents the evolution of the oxygen permeation as a
function of time in CO₂ sweeping conditions, at 900 °C. The flux kept stable
during 76 h and slightly increased with time on stream. The origin of this
intriguing activation with time remains unclear and may be related with surface
and/or grain boundary processes.
By increasing the separation driving force, \textit{i.e.}, the chemical potential across the membrane, the oxygen permeation can be enlarged depending on the $pO_2$ gradients. Figure 8.10 presents the permeation results when pure $O_2$ was fed instead of air and provides evidence about the important increase in the $J(O_2)$, specifically flux values up to 0.28 mL·min$^{-1}$·cm$^{-2}$ were achieved at 1000 °C. This improvement in the flux at high temperature may be related to bulk diffusion limited oxygen permeation, whereas other processes such as surface exchange reactions are more limiting below 800 °C. Alternatively, when methane is employed as sweep gas instead of Ar (and air feeding), $J(O_2)$ is strongly boosted, reaching flux values up to 8.71 mL·min$^{-1}$·cm$^{-2}$ at 1000 °C. This last increase in flux is not only related to the large increase in the transport driving force but also to the substantial rise in $n$-type conductivity achieved thanks to the higher reducing conditions in the permeate side. Such reduction
induces the lattice expansion and the generation of new strains, leading to the formation of cracks through the membrane inside the quartz setup. However, after measuring the cracked membrane, XRD pattern does not reveal the presence of reacted phases or impurities (Figure 8.11). Therefore, the membrane failure is attributed only to the mechanical stresses associated with the chemical expansion.

![XRD pattern](image)

*Figure 8.11. XRD pattern of both sides of the FNO-CTO membranes after 7 days of permeation experiments in Ar, CO\(_2\), CH\(_4\) sweeping.*

Finally, the membrane is compared with a reference composite made of FNO-CGO as this membrane is reported to be CO\(_2\) stable. [9] Figure 8.12 shows the permeation flux of both membranes. FNO-CTO composite membrane exhibits higher oxygen permeation and this difference becomes more important with decreasing temperatures, despite the lower ionic conductivity of CTO. The better behavior of the FNO-CTO membrane may originate from the higher \(p\)-type electronic conductivity of CTO, which increases the overall mixed
Ceria-based stable dual-phase membranes for oxygen separation

Ionic-electronic conductivity of the composite and makes possible to enlarge the surface reaction area from single TPB points, where the two solid phases and the gas phase converge, to the overall CTO grain surface. This is in contrast with the FNO-CGO composite since CGO behaves as a purely ionic conductor in the applied $pO_2$ environments and oxygen exchange is limited to single TPB points.

Figure 8.12. Temperature dependency of oxygen permeation of FNO-CTO and FNO-CGO composites. The membranes were fed by air and the sweep gas was 100% CO$_2$.

In addition to the permeation in CO$_2$, the stability of FNO-CTO composite has been studied by treating the membrane during 7 days at 800 °C under continuous flow of CO$_2$ with 5% O$_2$, 1% of SO$_2$ and 2.5% of H$_2$O. The membrane surface directly exposed to this gas environment was analyzed by XRD. The pattern plotted in Figure 8.13 reveals that only spinel and fluorite phases appear and neither new oxide phases nor minor impurities are detected up to the limits of the diffractometer. On the contrary, the membrane reference material La$_{0.8}$Sr$_{0.2}$Co$_{0.5}$Fe$_{0.5}$O$_{3-\delta}$ (LSCF) is completely destroyed when exposed
at the same treatment (Figure 8.14). This is a very remarkable result since these harsh environmental conditions are likely encountered in the targeted applications in (i) oxyfuel processes using recirculated flue gas as sweep and (ii) high-temperature catalytic membrane reactors for hydrocarbon conversion.

Figure 8.13. XRD pattern of FNO-CTO and FNO-CGO composites after 7 days of treatment under a continuous flux consisted of 1% SO₂-5% O₂ in CO₂, at 800 °C.
Summarizing, FNO-CTO composites seem appealing materials for oxygen separation devices for integration in oxyfuel processes using recirculated flue gas as sweep due to the particular combination of reasonable oxygen fluxes and unrecorded chemical stability at high temperatures. However, the oxygen fluxes are still orders of magnitude lower than the state of the art BSCF perovskites, which in addition has more economic components than the cercer composite. These issues should be taken into account. A step forward in the optimization of the membrane would imply the use of the cercer as a dense thin film, which is expected to increase the oxygen flux rate and decrease the material costs as the necessary amount of cercer would be substantially reduced.
8.3. Conclusions

One-pot fabrication of $60\% \text{Fe}_2\text{NiO}_4\cdot 40\% \text{Ce}_{0.8}\text{Ln}_{0.2}\text{O}_{2.5}$ (FNO-CLnO) composite has been proved to be the most suitable method to obtain both spinel and fluorite pure phase materials with a narrow grain size distribution and homogeneous distribution of grains, while the 60/40 phase ratio permits the existence of percolative paths from side to side of the dual-phase membranes. From DC-conductivity screening FNO-CTO composite has been selected having a comparable conductivity with FNO-CGO composite. An exhaustive study of this material has been carried out to assess the suitability as oxygen transport membrane.

FNO-CTO composite does not show evidence of impurities or reaction between phases up to the limits of the XRD. Additionally, SEM analysis reveals a narrow grain size distribution and homogeneous allocation of grains. Furthermore, the existence of percolative paths from side to side of the dual-phase membranes necessary for the oxygen permeation can be ascertained whilst the packing and bounding among the different grains in the membrane is appropriate.

The conductivity of both FNO-CTO phases is investigated as a function of temperature and $pO_2$. The ambipolar transport involves the concomitant (i) hopping of charge carriers, i.e., hopping of localized electrons between multivalent cations located in the octahedral Fe sublattice and (ii) the ionic transport in though the doped ceria lattice. Additionally, CTO presents significant $p$-type electronic conductivity although the ionic conductivity prevails.
Oxygen permeation flux values of 0.17 mL·min\(^{-1}\)·cm\(^{-2}\) and 0.20 mL·min\(^{-1}\)·cm\(^{-2}\) when argon and pure CO\(_2\) are used as sweep gas, respectively, are measured through a 0.68 mm-thick membrane at 1000 °C. These values are higher than those achieved using the reference composite FNO-CGO. Imposing higher \(pO_2\) gradients across the membrane leads to a substantial increase in the oxygen flux. Long term experiments at 900 °C showed that the material is CO\(_2\) stable and that oxygen permeation is even improved after 76 h on stream. Moreover, the composite was stable against SO\(_2\) and CO\(_2\) gas environments, as can be inferred from the stability test at 800 °C. However, permeation tests on SO\(_2\)-CO\(_2\) containing sweep gases would be necessary.

Conclusively, FNO-CTO composites seem appealing materials for oxygen separation devices for integration in oxyfuel processes since they combine reasonable oxygen fluxes and unrecorded chemical stability at high temperatures. A step forward in the optimization of the membrane would imply the use of the cercer as dense thin films.

8.4. References


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Chapter 9. Mixed protonic-electronic conductors based on lanthanum chromites

Abstract

A second block of materials on the research of mixed protonic-electronic conductors is presented. Perovskite-structured LaCrO$_3$ has been doped with the aim of modifying the electronic and catalytic properties in hydrogen-containing atmospheres, where these materials turn out to be proton conductors.

The results have addressed to a new generation of anodes for PC-SOFCs based on catalytically-promoted La$_{0.87}$Sr$_{0.13}$CrO$_3$ (LSC), La$_{0.85}$Sr$_{0.15}$Cr$_{0.9}$Ni$_{0.1}$O$_{3-\delta}$ (LSCN10), La$_{0.85}$Sr$_{0.15}$Cr$_{0.8}$Ni$_{0.2}$O$_{3-\delta}$ (LSCN) and La$_{0.75}$Ce$_{0.15}$Sr$_{0.1}$CrO$_{3-\delta}$ (LSCCe) chromite based materials are a good alternative to the widely used NiO, as they show good compatibility with LWO electrolyte in contrast to the highly reactive NiO. LSCCe is selected as electrode backbone structure, due to its superior total conductivity with respect to LSC, while LSCN10 and LSCN are selected to evaluate the role of the Ni in the structure. The infiltration of nickel nanoparticles in the sintered LSCCe and LSCN electrodes boosted the surface limiting reactions while preserving the compatibility. Metallic Ni nanoparticles become catalytically active centres for the H$_2$ oxidation reaction improving the anode performance regarding to pure LSC, previously limited by low frequency surface related processes.
9.1. Introduction

LaCrO$_3$ is a stable perovskite in both reducing and oxidizing atmospheres at high temperatures. Substituting alkali earth and transition metal elements on the A and B sites may modify the electronic and catalytic properties. Typically, Sr is introduced in A site, while other transition elements into the B-site of La$_{1-x}$Sr$_x$Cr$_{1.5}$M$_y$O$_3$ (M = Mg, Mn, Fe, Co, Ni) may improve the catalytic properties for, as example, methane fed SOFC. [1-3] These materials are traditionally known for their oxide ion conductivity, however, the conductivity experiments systematically conducted in different atmospheres evidence that these materials might turn out to be proton conductors in hydrogen-containing atmospheres.

Therefore, this kind of compounds is proposed to their potential application as hydrogen-separation membranes and catalytic membrane reactions. Hydrogen separation at high temperatures participates in the integrated gasification combined cycle (IGCC) of power plants. Thus, these membranes are the step to separate the H$_2$ and the CO$_2$ after fuel gasification, providing a moisturized CO$_2$ exhaust, ready to store.

On the other hand, recent interest in solid oxide fuel cells (SOFC) is focused on research and development of proton conducting SOFCs (PC-SOFC) that allow lowering the operation temperature and achieving very high fuel utilization, since it is not diluted by water that is formed in the air electrode. In this way, it emerges the possibility of simpler and cheaper cell designs. [4, 5] However, there still exists the need of developing novel protonic electrolytes.
and electrode materials, which combine excellent proton transport properties, compatibility with other cell components and thermochemical stability.

\( \text{La}_{6-x}\text{WO}_y \) (LWO, 0.4\(<x<0.7\)) \[6\] is a promising proton conductor, with high and predominant proton conductivity, and stable in wet CO\(_2\) under typical fuel cell operating conditions. \[6-10\] Therefore, compatible electrodes with the LWO electrolyte are currently being investigated. \[11, 12\] Recently, the development of specific LSM-LWO composite cathodes for LWO-based cells has been addressed. \[11\] Concerning the anodes, strong requirements entail high electrical conductivity, sufficient protonic conductivity, fast gas transport and exchange through a porous scaffold and fine-grained homogeneous microstructure, besides of high catalytic activity concerning hydrogen oxidation. Several nickel-based cermet combinations (Ni/YSZ, Ni/CGO, Ni/Ba(Ce\(_{0.8-x}\text{Zr}_x\))Y\(_{0.1}\)O\(_3\)) and mixed conductor Sr\(_{0.94}\)Ti\(_{0.9}\)Nb\(_{0.1}\)O\(_3\) (STN) have been proposed and investigated as SOFC anode materials. \[13-16\] However, these systems react with the LWO electrolyte, the reaction between LWO and NiO typical anode being especially dramatic. \[17\]

The present chapter focuses on the use of chromite based perovskites as PC-SOFC anode materials. Indeed, doped \( \text{La}_{0.85}\text{Sr}_{0.15}\text{CrO}_3 \) has already been tested as anode for conventional SOFCs, due to its particular combination of redox stability and sufficient \( p \)-type electronic and oxygen-ion conductivity under reducing conditions, while its electrochemical activity can be fine-tuned by incorporating catalytic doping elements, such as nickel and ruthenium. \[18-22\] Furthermore, doped \( \text{LaCrO}_3 \) compounds have been claimed as hydrogen permeation membrane materials and this particular function suggests the presence of mixed protonic-electronic conductivity in this material. \[23\]
Here, the initial LSC electrode has been improved stepwise by tackling the two major rate-limiting steps, *i.e.*, electrode total conductivity and surface processes. Firstly, the enhancement of the transport properties of the chromite has been tried by doping on A and B-perovskite positions (A =Sm, Ce, and Nd and B= Ni, Y, Al, Fe, Ti, Ga, Mn, Zn) and Ce and Ni-doped LSC have been selected for further investigation. Secondly, nickel infiltration into the sintered Ce and Ni-doped electrodes has been carried out in order to promote the surface reactions, since Ni is a well-known, highly active H₂ bond breaking catalyst.

Symmetrical cells made by La₀.₈₇Sr₀.₁₃CrO₃ (LSC), La₀.₈₅Sr₀.₁₅Cr₀.₉Ni₀.₁O₃ (LSCN10), La₀.₈₅Sr₀.₁₅Cr₀.₈Ni₀.₂O₃ (LSCN) and La₀.₇₅Sr₀.₁₅Ce₀.₁CrO₃ (LSCCe), both bare and infiltrated with Ni nanoparticles, are tested as anodes for LWO based PC-SOFCs aiming (1) to compare results of the new Ni and Ce doped LSC material with previously studied LSC based anodes and (2) to analyse the effect of the surface catalytic promotion due to the Ni infiltration.

9.2. Results and discussion

Screening of doped LSC

LSC and doped-LSC compounds have been prepared by the Pechini method explained in section 2.1. Powders of nanometric size are obtained after calcination of the dry precursors at 600 °C. Table 9.1 summarizes the nominal stoichiometry and short nomenclature of the A and B-site doped perovskite
compositions \( (A = \text{Sm, Ce, and Nd and } B = \text{Ni, Y, Al, Fe, Ti}) \). Initial XRD measurements evaluate whether the doping succeeded to form pure perovskite phase. As example, the patterns of LSC calcined at 900 °C, LSCN10 and LSCN20 at 1050 °C and LSCrF at 1000 °C are represented in Figure 9.1.

![XRD patterns of LSC calcined at 900 °C, LSCN10 and LSCN at 1050 °C and LSCrF at 1000 °C. The dotted lines represent the LSC reference pattern (ICSD 00-032-1240).](image)

Figure 9.1. XRD patterns of LSC calcined at 900 °C, LSCN10 and LSCN at 1050 °C and LSCrF at 1000 °C. The dotted lines represent the LSC reference pattern (ICSD 00-032-1240).

However, the chromites need very high temperatures to obtain the dense specimens required by the transport measurements, so that the structure was checked after pressing and firing the samples at 1600 °C.
Figure 9.2 shows the XRD patterns of A-site doped LSC beside their corresponding SEM microstructure images of SmSC, NSC and LSCCe. Only SmSC and LSCCe are pure phase, while NSC presents Nd$_2$O$_3$ and CrO$_2$ impurities. Regarding to the microstructure, the LSCCe shows the most homogeneous grain size distribution.

![XRD patterns and SEM microstructure images](image)

Figure 9.2. XRD patterns of A-site doped LSC calcined at 1600 °C and corresponding SEM microstructure images of (top to bottom) SmSC, NSC and LSCCe.

On the other hand, B-site doped LSC does not deliver single perovskites but a combination of hexagonal (LSC) and orthorhombic perovskite (LC) when they are submitted to 1600 °C. However, dense specimens are obtained, with
only closed porosity. In particular, LSCrF is very well packed, which is expected to be beneficial on the transport properties.

Figure 9.3. XRD patterns and corresponding SEM images of B-site doped LSC pressed and calcined at 1600 ºC: LSCY, LSCT, LSCA and LSCrF. The peak positions of LaCrO$_3$ (LC, ICSD 00-024-1016) and LSC (ICSD 00-032-1240) perovskites are represented for comparison as dotted vertical lines.

In order to test the possibility of both protonic and $p$-type electronic conductivity of the materials, the transport properties were measured by DC-conductivity in reducing atmospheres, i.e., 5% H$_2$ and 5% D$_2$ dry and wet (2.5% atm of H$_2$O and D$_2$O respectively). It is observed that the conductivity in wet atmospheres is higher than in dry atmospheres, especially at low temperatures. At higher temperatures (ca. 800 ºC), the exothermal oxide
Mixed protonic-electronic conductors based on lanthanum chromites

dehydration occurs, which drops the protonic carriers equalling the conductivity in wet and dry conditions. [28] Secondly, the conductivities in H₂ are higher than in D₂ in both dry and wet atmospheres, the so-called isotopic effect. The results at 700 °C indicate the mixed protonic-electronic character of doped LSC showing a clear isotopic effect. As direct comparison, the total conductivity in wet H₂ atmosphere at 700 °C, and the corresponding activation energy are written in the Table 9.1. The best conductivity results in wet H₂ are obtained for LSCCe. For a question of clarity, only further details about the transport properties of the selected lanthanum chromites are shown and explained below.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Nomenclature</th>
<th>Structure (at 1600 °C)</th>
<th>σ₇₀₀°C (H₂-H₂O) S·cm⁻¹</th>
<th>Eₐ eV</th>
</tr>
</thead>
<tbody>
<tr>
<td>La₈₇Sr₁₃CrO₃</td>
<td>LSC</td>
<td>Perovskite</td>
<td>6</td>
<td>0.28</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₉Ni₀.₁O₃</td>
<td>LSCN10</td>
<td>Perovskite</td>
<td>1</td>
<td>0.25</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₈Ni₀.₂O₃</td>
<td>LSCN</td>
<td>Perovskite</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₉Ce₀.₁O₃</td>
<td>LSCCe</td>
<td>Perovskite</td>
<td>11</td>
<td>0.28</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₉Fe₀.₁O₃</td>
<td>LSCF</td>
<td>Perovskite mixture</td>
<td>6.4</td>
<td>0.38</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₉Ti₀.₁O₃</td>
<td>LSCT</td>
<td>Perovskite mixture</td>
<td>1.1</td>
<td>0.51</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₉Al₀.₁O₃</td>
<td>LSCA</td>
<td>Perovskite mixture</td>
<td>1.9</td>
<td>0.43</td>
</tr>
<tr>
<td>La₈₅Sr₁₅Cr₀.₉Y₀.₁O₃</td>
<td>LSCY</td>
<td>Perovskite mixture</td>
<td>6.2</td>
<td>0.20</td>
</tr>
<tr>
<td>Sm₈₅Sr₁₅CrO₃</td>
<td>SmSC</td>
<td>Perovskite</td>
<td>4.1</td>
<td>0.72</td>
</tr>
<tr>
<td>Nd₈₅Sr₁₅CrO₃</td>
<td>NdSC</td>
<td>Perovskite+impurities</td>
<td>4.1</td>
<td>0.4</td>
</tr>
</tbody>
</table>

Table 9.1. Summary of doped lanthanum strontium chromites composition and short nomenclature. The main structural phase formation is indicated as well as the total conductivity in wet H₂ atmosphere at 700 °C, and the corresponding activation energy.

Although highly conducting materials have been obtained by doping LSC, the difficulty to obtain dense specimens in combination with a single phase structure limits the use of these materials for some applications, at least
with the fabrication approaches considered in this thesis. For hydrogen membrane application, other densification methods that do not employ such high temperatures, as spark plasma, or the development of a composite might be tried. Nevertheless, these materials may be used as anode in PC-SOFC, since a proper porous microstructure is a condition necessary for the gas current through the electrode to the electrolyte and vice versa.

Thus, further studies on the materials showing a single perovskite phase and good conductivity in wet H\textsubscript{2} are done. It is the case of LSCCe, which will be tested as anode backbone to improve the protonic transport through a proton conducting electrolyte, \textit{i.e.}, the LWO specified in the introduction. On the other hand, even though it does not improve the conductivity, the composition LSCN10 is tested as anode material due to the Ni structural loading, as the typical NiO anode material is incompatible with LWO under the oxidizing conditions necessary for the sintering and assembling of the fuel cell.

\textit{La}_{0.85}\textit{Sr}_{0.15}\textit{Cr}_{0.9}\textit{Ni}_{0.1}\textit{O}_{3.\delta} (LSCN10) anode

In first place, the behavior of a 10\% Ni doped-LSC, \textit{i.e.}, LSCN10, as anode has been tested and compared with pure LSC. Symmetrical cells have been prepared by screen printing LSC and LSCN10 inks on LWO electrolyte and fired at 1050 \textdegree C. The chemical compatibilities of LSC and LSCN10 anodes on LWO electrolyte have been tested by XRD analysis of both phases after sintering at high temperature. Figure 9.5 shows the XRD pattern of a mixture of LSC and LWO 50/50 vol.\% fired at 1400 \textdegree C. Single phase component materials are also plotted as reference. Both phases appear in the composite pattern without reaction evidences or impurities, confirming the material compatibility.
Figure 9.4. XRD patterns of (from bottom to top): LSC, LWO and their 50 vol.% mixture at RT, treated at 1400 °C in H₂ for 48 h and treated at 1150 °C in air for 5 h.

On the other hand, 10% Ni doping LSC aims to introduce catalytic properties on the anode without stability degradation. Figure 9.5 shows the XRD pattern of a mixture of LSCN10 and LWO 50/50 vol.% after being treated for 5 h at 1150 °C in air. No new crystalline phases appear whereas all diffraction peaks can be completely assigned to LSCN10 or LWO phases. This is in contrast with the high reactivity of LWO with NiO, which leads to the formation of several perovskite-related products (also shown in Figure 9.5). The confirmation of the lack of reactivity between LWO and metallic Ni can be inferred from the XRD pattern of the LWO+NiO 50 vol.% after treatment at 900 °C for 48 h in H₂, since no reaction products appear.
Figure 9.5: XRD patterns of (from bottom to top): LWO and NiO 50 vol.% mixture at RT, treated at 900 °C in H\textsubscript{2} for 48 h and treated at 1150 °C in air for 5 h; and LSCN10 and LWO 50 vol.% mixture after calcination in air at 1150 °C for 5 h.

SEM and TEM images of the LSCN10 anode on LWO electrolyte (Figure 9.6a and b, respectively) permit ascertaining that the integrity of the interface between both materials is preserved after sintering and no reaction layers appear. This is confirmed by an EDS analysis of the different phases observed in the TEM picture. In addition, anode porosity for gas exchange is confirmed, as well as relatively high surface area available for the electrochemical reaction. The LWO electrolyte is dense and only presents closed porosity.
Mixed protonic-electronic conductors based on lanthanum chromites

Figure 9.6: a) SEM image and b) TEM image of a FIB-lamella of a LSCN10 anode on LWO electrolyte sintered at 1150 ºC

On the other hand, TEM images of the reduced electrode have been taken in order to see the effect of the operation conditions in the microstructure, i.e., high temperature and reducing conditions. It is observed that LSCN10 splits the Ni dissolved into the structure distributing metallic Ni nanoparticles over the surface. Figure 9.7a and b show the TEM images of as sintered LSCN10 anode on LWO electrolyte and reduced LSCN10 powder, where Ni nanoparticles are visible, respectively. These Ni nanoparticles (19.7±3 nm when material is reduced at 900 ºC) turn into catalytic active particles boosting the anodic hydrogen oxidation reaction. [24-26] Furthermore, Ni particles are chemically compatible with LWO as concluded from Figure 9.5. Hence, the TEM analysis suggests that a higher Ni loading would increase the surface concentration of metallic particles, which will be studied below.
Figure 9.7: TEM images of a) as sintered LSCN10 on LWO electrolyte revealing the good bounding between electrolyte and porous electrode particles and b) LSCN10 powder after reduction at 900 °C.

The formation of the metallic Ni nanoparticles upon reduction in hydrogen is monitored by TPR measurements, as showed in Figure 9.8a. While no reduction takes place in the undoped LaCrO$_3$, a reduction peak appears centred at 380 °C when Cr is partially substituted by Ni. This reduction is related to two reduction processes, i.e., the reduction of Ni$^{2+}$ to metallic nickel, and the reduction of Cr$^{4+}$ to Cr$^{3+}$ and more reduced species (Cr$^{δ+}$) as supported by reported XPS measurements, which involves the concomitant formation of new oxygen vacancies. [27] This reduction peak increases substantially when Sr$^{2+}$ further substitutes La$^{3+}$, since the concentration of Cr$^{4+}$ increases to compensate the presence of Sr$^{2+}$ and Ni$^{2+}$ in oxidizing conditions. This additional reduction is attributed to (1) the formation of metallic Ni nanoparticles active in the anodic reactions and (2) the formation of oxygen vacancies, indispensable for the oxide hydration and protonic transport, through Cr$^{4+}$ reduction.

The kinetics of Ni particle formation has been studied using a LSCN10 porous electrode on LWO. The anode polarization resistance ($R_p$) was monitored at 900 °C upon feeding pure moist hydrogen. The total reduction of
the LSCN10 material with the consequent formation of metallic Ni lasted for 6.6 h, as can be inferred from Figure 9.8b, while the promotion of the electrochemical reactions through the Ni nanoparticles is confirmed.

![TPR analysis and Rp of LSCN10](image)

Figure 9.8: a) TPR analysis of LaCrO$_3$-$\delta$, LaCr$_{0.9}$Ni$_{0.1}$O$_3$-$\delta$, and LSCN10 materials and b) $R_p$ of LSCN10 as a function of time at 900 ºC in 100% H$_2$.

The transport properties of reduced LSCN10 in 5% H$_2$ and 5% D$_2$ dry and wet (2.5% atm of H$_2$O and D$_2$O respectively) have been studied by DC-conductivity. The results plotted in Figure 9.9 show that (i) the conductivity in wet atmospheres is higher than in dry atmospheres although the exothermal oxide dehydration at high temperatures drops the protonic conductivity and equals the conductivity in wet and dry conditions, e.g. at 800 ºC in wet and dry H$_2$; this effect is expected to occur at higher temperatures in D$_2$ due to slightly different oxide hydration equilibrium for D$_2$O; [28] (ii) the conductivities in H$_2$ are higher than in D$_2$ (very close to the theoretical $\sqrt{2}$ factor) in both dry and wet atmospheres. The isotopic effect in dry atmospheres is ascribed to the presence
of residual amounts of H$_2$O/D$_2$O (~30 ppm) in the so-called dry atmospheres; and (iii) the dependency of conductivity with $pO_2$ (D$_2$$<$H$_2$; D$_2$-D$_2$O$<$H$_2$-H$_2$O) indicates that the conductivity rises with increasing $pO_2$, specially at temperatures below 800 ºC. The results suggest that LSCN presents both protonic and $p$-type electronic conductivity under typical anode operating conditions. In fact, the conductivity data follow the expected hydration and isotopic effects for a proton conductor while electron-hole is affected by the corresponding $pO_2$ being attributed to $p$-type conduction. [11] At 600 ºC the mixed protonic-electronic character of LSCN is well evidenced showing a clear isotopic effect. However, at 800 ºC, the $pO_2$ dependency becomes minor and the isotopic effect is still visible, which suggests the growing contribution of oxide-ion conductivity at high temperatures.

Figure 9.9: Total conductivity of La$_{0.85}$Sr$_{0.15}$Cr$_{0.9}$Ni$_{0.1}$O$_{3-δ}$ in 5% H$_2$, 5% D$_2$, 5% H$_2$+H$_2$O and 5% D$_2$+D$_2$O.

Up to here, the compatibility of the LSCN10 with LWO, and the adequate transport properties and catalytic activity of LSCN material have been evidenced. Next, the suitability as PC-SOFC anode material is analysed by
electrochemical impedance spectroscopy (EIS) in symmetrical cells on LWO electrolyte.

Figure 9.10 a and b respectively represent Nyquist and Bode plots of the EIS analysis of LSC recorded in wet 100% H₂ at 750 ºC after subtracting electrolyte contribution. Figure 9.10c and d shows similar representation for LSCN10 recorded in wet 100% H₂ at different temperatures. The impedance of LSCN10 is decreased by some orders of magnitude with respect to LSC. This effect is attributed to surface activation when Ni is finely dispersed on the chromite grain surface.

Figure 9.10: a) Nyquist and b) Bode plots of impedance spectra recorded in wet (2.5% vol. H₂O) 100% H₂ at LSC at 700 ºC and c) Nyquist and d) Bode plots of impedance spectra recorded in wet (2.5% vol. H₂O) 100% H₂ at different temperatures for LSCN10.

Figure 9.10 show two main contributions that are fitted to the double R||CPE equivalent circuit illustrated in Figure 9.11, which comprises two
depressed arcs, one observed at low frequencies (LF, 10-100 Hz) and another at medium frequency (MF, 0.1-1 kHz), whose overlapping depends on the operating conditions. The depressing arcs need the use of these CPEs (Constant Phase Elements) instead of capacitors as corresponds to the porous structure of the electrode. The impedance of a CPE can be expressed as:

\[ Z = A^{-1}(j\omega)^{-n} \]  

(Eq. 9.1)

where \( A \) is a frequency independent constant, \( \omega \) is the angular frequency and \( j = \sqrt{-1} \). Note that there are two limiting cases, first when \( n = 1 \), the CPE operates as an ideal capacitor and second when \( n = 0 \), the CPE acts as a pure resistor. [29] The equivalent capacitance of the CPEs can be calculated according to the following equation. [30]

\[ C_{\text{CPE}} = R^{(1-n)/n}A^{1/n} \]  

(Eq. 9.2)

![Diagram](image)

Figure 9.11. Equivalent circuit used for EIS impedance modeling. CPS are used instead of perfect capacitors due to the porous nature of the anode.

LF and MF resistances (R) of LSCN10 measured in 100% moist H\(_2\) are represented as a function of temperature in Arrhenius arrangement (Figure 9.12a). Above 850 °C both LF and MF contributions are similar but below 850 °C the rate limiting mechanism corresponds to the LF associated processes, being twice the MF contribution. The resistance increases when reducing \( pH_2 \), and at 750 °C, both LF and MF associated mechanisms have a \( pH_2 \) dependency close to \(-1/4\) (Figure 9.12b). The LF process (with capacitances in the range of \( 10^{-2} \) F/cm\(^2\)) may be related to surface processes: catalytic reactions between chemisorbed species, adsorption/desorption reactions from/to the gas phase and
surface diffusion of intermediate species. [31] This important contribution to the anode impedance of the adsorption/desorption of H₂O and OH⁻ species has been previously observed in similar frequency ranges and activation energies for Ni-YSZ/Ni-CGO anodes. [32] These results anticipate that the further development of chromite-based PC-SOFC anodes should focus on the optimization of surface activity, either increasing the intrinsic activity of the catalyst or improving the catalyst dispersion, in order to diminish LF associated processes. Furthermore, MF processes (with capacitances in the range of 10⁻¹⁻¹⁰⁻² F/cm²) might be associated to charge transfer reactions, [33] i.e., (i) the charge transfer between Ni nanoparticles and the oxide and (ii) the diffusion charged species to the three phase boundary. [34]

Figure 9.12. a) Modeled LF and MF resistances as a function of the temperature and b) as a function of the pH₂.
Finally, the XRD in Figure 9.13 shows the structure pattern of the reduced LSCN10 anode on LWO after electrochemical cell testing. It demonstrates the stability of the material in working conditions. The perovskite and the fluorite phases are observed without any reaction between them. Also peaks corresponding to the metallic Ni in the surface after the reduction and the Au from the current collector are distinguished.

Figure 9.13. XRD pattern in logarithm scale of LSCN10 anode after electrochemical test. The materials maintains its pure perovskite structure.

**Catalytic surface promotion of La_{0.85}Sr_{0.15}Cr_{0.8}Ni_{0.2}O_{3-δ} (LSCN) anodes**

The LSC and LSCN10 anodes have been shown above to be specially limited by surface reactions. Therefore, further development of chromite-based PC-SOFC anodes should focus on the optimization of surface activity. Nickel infiltration into the electrodes has been considered, since metallic Ni
Mixed protonic-electronic conductors based on lanthanum chromites

nanoparticles have been demonstrated to be compatible with LWO under anode operating conditions and they are active catalyst in the \( \text{H}_2 \) bond breaking reaction. [20-22] Concerning the anode backbone, the first material of choice is 20% Ni doped in the B site, which was deduced in the previous section to increase Ni nanoparticles segregation.

The structure of the LSCN powder sintered at 900 °C can be observed from XRD pattern plotted in Figure 9.14a. The main diffraction peaks can be assigned to the orthorhombic \( \text{LaCrO}_3 \) perovskite phase, although some traces of \( \text{SrCrO}_4 \) spinel phase can also be observed. [35] These traces of \( \text{SrCrO}_4 \) were previously observed at this low sintering temperature but it was also proved that they disappear upon reducing conditions. SEM image of Figure 9.14b shows the microstructure of this powder, which presents a homogenous grain size distribution.

Figure 9.14. a) XRD pattern and b) SEM image of LSCN powder sintered at 900 °C

Symmetrical cells were prepared on LWO electrolyte by using LSCN ink, as explained in Chapter 2.1. Once the electrolyte was fired at 1050 °C, the infiltration of Ni in the LSCN electrode (LSCN+5Ni) was done by dropping a
5 M water solution of Ni nitrate on the LSCN electrode surface. It was allowed to penetrate into the pores for 30 min and calcined at 300 °C to decompose the nitrates. This process was repeated 4 times. XRD patterns of the different symmetrical cells, LSC, LSCN and LSCN+5Ni, are represented in Figure 9.15 (intensity in log scale) after the impedance measurements at high temperature and reducing conditions. The diffraction peaks of the LSC cell can be assigned to (1) the chromite perovskite phase, (2) the LWO electrolyte fluorite phase and (3) metallic Au, which is used as current collector in the symmetrical cell testing. As expected after the reducing conditions, no traces of SrCrO₄ are observed. When the anode material is doped with Ni (LSCN symmetrical cell) after reducing conditions diffraction peaks corresponding to metallic Ni are also visible. This metallic Ni is segregated from the LSCN compound due to the high temperature and the reducing conditions, as it has been previously proved. [24] The intensity of these Ni peaks is increased in the case of LSCN+5Ni symmetrical cell.

![Figure 9.15](image)

Figure 9.15. XRD patterns of LSC, LSCN and LSCN+5Ni symmetrical cells. The peaks corresponding to metallic Au from the current collector of the symmetrical cell and metallic Ni (infiltrated and structural) are indicated.
The Ni nanoparticle segregation from the LSCN material after reduction at high temperature is represented in the scheme of Figure 9.16a. However, from its real SEM image, showed in Figure 9.16b, it is not easy to distinguish the presence of these nanoparticles, due to the very small size and low amount of Ni. For this reason, in previous reported works, TEM was used for the characterization of these segregated particles. [24] The scheme of the LSCN electrode infiltrated by using 5 M water solution of Ni nitrate (LSCN+5Ni) is showed in Figure 9.16c aside with corresponding SEM images of the cross section anodes after measurements depicted in Figure 9.16d and e (two different magnifications). These SEM images show the covering by Ni nanoparticles on the LSCN+5Ni electrode backbone after reduction at 800 ºC. In fact, the surface of the LSCN+5Ni grains is fully covered with well distributed Ni nanoparticles, along the thickness of the electrode. The average Ni particle size infiltrated in the electrode is 10-20 nm. This size matches with ~10-20 nm Ni particles resulted from Ni nitrate calcination and subsequent reduction at 800 ºC in H₂, which are observed in the TEM image of Figure 9.17 (measured in powder). Electrode microstructural parameters are of great importance especially for infiltrated electrodes, since the pore should host the Ni nanoparticles and allow the mean free path of the gas molecules. Indeed, the SEM images of the anodes (Figure 9.16) after measurements in reducing conditions show that the highly-porous backbone is adequate for Ni hosting and proper gas transport.
Figure 9.16. Schematic diagram of the LSCN anode a) and its corresponding SEM image (b) and scheme of the LSCN+5Ni anode c) and its real SEM images d) and e). 5 M water solution of Ni Nitrate was dropped on the surface of the LSCN electrode and allowed to penetrate into the pores for 30 min. The infiltration step was repeated 4 times. Each infiltration step was followed by calcination at 300 °C to decompose the nitrates.
Mixed protonic-electronic conductors based on lanthanum chromites

Figure 9.17. TEM image of Ni nanoparticles obtained by firing nitrate precursor at 800 °C in H₂. Particle size is in between 10 and 20 nm.

The electrochemical properties of the three different tested anodes, LSC, LSCN and LSCN+5Ni, can be evaluated to determine the polarization resistance (Rp) values as a function of the temperature, plotted in Figure 9.18. At 700 °C in wet hydrogen LSC anode shows Rp of 347 Ω·cm². In the case of the LSCN anode, the segregation of the Ni nanoparticles results in two orders of magnitude Rp decrease, showing a Rp value of 3.54 Ω·cm² at 700 °C. This indicates that the presence of the Ni nanoparticles actively helps the anode performance, which may be attributed to the Ni catalytic activity over the as H₂ oxidation reaction. Ni infiltration on LSCN anode decreases the Rp almost another order of magnitude, reaching a value of 0.47 Ω·cm² at 700 °C. This improvement can be directly related to the higher amount of active Ni nanoparticles in the anode.
In order to discern the nature of the electrochemical improvements due to the Ni nanoparticles that are on one hand segregated from the Ni doped compound and, on the other, obtained by Ni infiltration, EIS spectra are analyzed for the different materials.

Figure 9.19 shows the EIS spectra at 700 °C of LSC anode (a and b, Nyquist and Bode plots, shown again for a direct comparison) and LSCN and LSCN+5Ni (c and d, Nyquist and Bode plots) after subtracting the electrolyte contribution. The spectra show two different contributions in both LSC and LSCN anodes, i.e., two different arcs at different frequencies. Otherwise, LSCN+5Ni anode only shows one main contribution, despite a small noise ascribed to the contacts appears at low frequencies and high temperatures. Figure 9.19b demonstrates that the LSC anode is limited by low frequency (LF) associated processes, although high frequency (HF) associated processes remain an important contribution as well. The structural Ni doping the LSCN is able to segregate in the surface of the material under operating conditions as
nanoclusters, so that the LF contribution is drastically decreased (almost two orders of magnitude), although it is still limiting the overall anode performance. LSCN anode also shows another contribution at higher frequencies. By Ni nanoparticles infiltration along the anode surface, the LF contribution disappears and high frequency (HF) associated processes limit the LSCN+5Ni anode performance.

Figure 9.19. Impedance spectra recorded at 700 °C in wet hydrogen of LSC a) Nyquist and b) Bode plots and LSCN and LSCN+5Ni c) Nyquist and d) Bode plots. Solid lines represent the depressed arcs fitting the different electrode resistance contributions.

These different HF and LF contributions observed for LSC and LSCN can be analyzed in terms of two depressed arcs by fitting EIS spectra to a double R∥CPE circuit (Figure 9.11). In the case of LSCN+5Ni anode only one arc is needed, fitted to a single R∥CPE circuit. Modeled resistances and capacitances (calculated from CPEs) are depicted in Figure 9.20a and b, and their characteristic frequencies are written in the graph. LSC anode shows both limiting LF (2-25 Hz) associated processes and also very high resistances
associated to HF (1-10 kHz) processes. Its associated capacitance values are \(10^3\) F/cm\(^2\) for LF processes and \(10^5-10^4\) F/cm\(^2\) for HF, which combined with their characteristic frequencies can be assigned to surface associated processes and LSC transport properties, respectively.

![Figure 9.20. Temperature dependency of modelled resistances a) and capacitances b) of LSC, LSCN and LSCN+5N samples.](image)

When the Ni is added as dopant, the LSCN anode shows LF (10-100 Hz) and medium frequency, MF (0.1-0.5 kHz), contributions. Associated capacitances are \(10^{-2}\) F/cm\(^2\) and \(10^{-3}-10^{-2}\) F/cm\(^2\) for LF and MF processes, respectively. These characteristic frequencies and their corresponding associated capacitances can be related to surface processes. In particular, the MF is associated to surface exchange coupled with bulk ionic transport in mixed conducting electrodes. [36] LF resistance decreases more than one order of magnitude compared to that of LSC, while the range of characteristic frequencies shift to higher values. This decrease of the LF resistance can only be attributed to the effect of the active Ni nanoparticles formed on the surface of the anode after reduction, which improves the surface dissociation of hydrogen. The disappearance of this LF contribution in the
LSCN+5Ni anode demonstrates the great importance of the catalytic activity of these Ni nanoparticles. In fact, LSCN+5Ni anode only shows HF (1-15 kHz) resistance with associated capacitances of $10^{-3}$-$10^{-4}$ F/cm$^2$. These HF processes may be attributed to the transport properties of the LSCN material or to the LSCN/LWO interfacial resistances, i.e., (i) the charge transfer between Ni nanoparticles and the oxide and (ii) the diffusion of charged species to the three phase boundary. [33,34] The pronounced resistance improvement confirms that infiltration is a suitable method to support Ni catalyst nanoparticles to increase the surface contact with fuel.

**Catalytic surface promotion of $La_{0.75}Ce_{0.1}Sr_{0.15}CrO_3-\delta$ (LSCCe) anodes**

The second anode development strategy consisted of improving the total conductivity of the material, which was achieved by Ce doping on the A-site. Symmetrical LSCCe electrodes were screen printed on the LWO dense electrolyte and were infiltrated afterwards by different Ni loads (0, 1 and 5 Ni loads) to boost the surface reaction.

Figure 9.21 presents the XRD analysis of the LSCCe powder and the Ni-infiltrated LSCCe anode after being submitted to the reduction and to the electrochemical characterization. Powder sintered at 1200 °C still present traces of SrCrO$_4$ and CeO$_2$, which disappear either at higher temperatures or in reducing atmospheres. [24, 30] In fact, LSCCe anode after testing only shows diffraction peaks corresponding to the chromite, the LWO from the electrolyte, Au from the current collector and Ni catalyst. The lack of any other diffraction
peaks that could be assigned to any reaction product denotes the compatibility of the LSCCe with the LWO electrolyte, even after the Ni infiltration, and the stability in anode working conditions.

![XRD pattern of LSCCe as prepared powder and anode after Ni infiltration and reduction in wet H₂ up to 800 ºC. XRD positions of LaCrO₃, LWO and CeO₂ are indicated as reference.]

**Figure 9.21.** XRD patterns of LSCCe as prepared powder and anode after Ni infiltration and reduction in wet H₂ up to 800 ºC. XRD positions of LaCrO₃, LWO and CeO₂ are indicated as reference.

Besides the material stability under working conditions, Figure 9.22 shows that the material is stable in CO₂ containing atmospheres. The thermogravimetric curve (Figure 9.22b) shows a slight weight loss at temperatures up to 400 ºC, attributed to water and oxygen desorption, while at higher temperatures the mass remains practically constant. The lack of weight gain indicates that carbonation processes/reactions (CO₂ uptake) at temperatures below 800 ºC are not happening.
In order to investigate the transport properties of the material, DC-conductivity measurements were carried out. Figure 9.23a plots the total conductivity of LSCCe in wet H₂ and D₂ as a function of temperature. For comparison the LSC conductivity in H₂+H₂O (anode operating conditions) is also depicted. LSCCe exhibits higher total conductivity under reducing atmospheres than LSC. Besides, the conductivity in H₂+H₂O is higher than in D₂+D₂O, which denotes an isotopic effect typical of protonic transport.

In the studied pO₂ range, the conductivity of chromites usually presents a predominant p-type electronic character. [18, 19] Moreover, ionic (O²⁻ and H⁺) transport represents an important contribution due to the oxygen vacancies created upon Cr and Ce reduction. [24, 30, 36, 37] This is confirmed for LSCCe by means of DC-conductivity analysis as a function of pO₂. Figure 9.23b and 9.23c display the conductivity power dependency on pO₂ (σ ∝ pO₂ⁿ) at 700 ºC in dry and wet conditions, respectively. Figure 9.23b confirms the conductivity dependence of pO₂¹⁄₄ expected for a predominant p-type electronic conductor in
dry atmospheres, as it is the case of the reported chromites. [20-22] In wet atmospheres there is a decrease of the \( pO_2 \) slope up to 1/6, which is likely a consequence of the introduction of protonic carriers.

Summarizing, the protonic conductivity is deduced from the observed isotopic effect and from the smaller \( pO_2 \) dependency (1/6), and combined with the dry dependency, the mixed ionic-protonic transport of the material under these conditions is pointed out. Due to this mixed electronic-protonic behaviour one may expect the length of triple phase boundary (TPB) within the electrode to increase up to the whole electrode surface, enhancing the anode performance.
In order to further improve this anode, different amounts of Ni were infiltrated into the sintered anode aiming to study the effect of the catalytic coating on the electrochemical activity. The three different LSCCe anodes discussed are shown schematically in Figure 9.24: (a) bare LSCCe anode, (b) LSCCe infiltrated by using 1 M water solution of Ni nitrate (LSCCe+1Ni) and (c) LSCCe infiltrated by using 5 M water solution of Ni nitrate (LSCCe+5Ni). The real SEM images of the cross section anodes after measurements are shown in Figures 9.24d, 9.24e and 9.24f, respectively. The study of these three
different anodes aims (1) to compare results of the new Ce-doped LSC material with previously studied LSC based anodes and (2) to analyse the effect of the surface catalytic promotion due to the Ni infiltration.

SEM images (Figure 9.24e and f) provide evidence that the quantity of Ni nanoparticles increases with the Ni concentration of aqueous solution employed in the infiltration. Further, the average Ni particle size is similar (∼10-20 nm) to that shown for the Ni powder reduced from precursor nitrates in Figure 9.17.

![Figure 9.25](image)

**Figure 9.25.** Rp as a function of the inverse temperature of the four anodes $\text{La}_{0.85}\text{Sr}_{0.15}\text{CrO}_3-\delta$ (LSC), $\text{La}_{0.75}\text{Ce}_{0.1}\text{Sr}_{0.15}\text{CrO}_3-\delta$ (LSCCe) and LSCCe infiltrated with 1Ni (LSCCe+1Ni) and with 5Ni (LSCCe+5Ni).

Electrochemical properties of the different anodes were analyzed by means of EIS measurements. Polarization resistances ($R_p$) extracted at different temperatures (Figure 9.25) for all the studied anodes allow comparing (1) the improvement caused by the enhancement of transport properties of the backbone chromite material, and (2) the further improvement of the LSCCe
anode performance achieved through the infiltration and surface coating with Ni nanoparticles.

Figure 9.26. EIS spectra at 700 °C in H₂ of LSC and LSCCe anodes, a) Nyquist and b) Bode plots and EIS spectra at 700 °C in H₂ of LSCCe infiltrated with 1Ni and 5Ni, c) Nyquist and d) Bode plots.

The impedance spectra at 700 °C in wet H₂ are plotted in Figure 9.26a-d. They can be analysed by adjusting to a double serial R∥CPE elements (Figure 9.11). With the two arcs interpretation we are able to distinguish different processes occurring in different ranges of frequencies, low, medium and high.
Figure 9.27. Modeled R contributions as a function of the inverse temperature of the LSCCe anode and when infiltrated with different amounts of Ni: 1Ni (LSCCe+1Ni) and with 5Ni (LSCCe+5Ni).

Figure 9.28. Associated capacitances as a function of the inverse temperature of the LSCCe, LSCCe+1Ni and LSCCe+5Ni anodes.

Figure 9.27 presents the modeled resistances for the pristine LSCCe and the two infiltrated LSCCe electrodes (LSCCe+1Ni and LSCCe+5Ni) as a function of temperature (600–800 °C); the corresponding associated
Mixed protonic-electronic conductors based on lanthanum chromites - 301 -

capacitances are available in Figure 9.28. The LSCCe anode (Figure 9.24a and d) is principally limited by processes ascribed to low frequencies (LF, 0.3-25 Hz) and this becomes more evident with decreasing temperatures. LSCCe anode shows another important contribution at medium frequencies (MF, 33-200 Hz). Due to the characteristic frequencies and associated capacitances, both contributions can be related to surface processes, being the MF one typically associated to surface exchange coupled with bulk ionic transport in mixed conducting electrodes. [36, 37]

With the introduction of a limited amount of Ni nanoparticles (LSCCe+1Ni, Figure 9.24b and e) the LF contribution disappears, the MF associated processes shift to higher frequencies (MF, 200–500 Hz) and a new contribution becomes visible at high frequencies (HF, 1–100 kHz). This confirms the catalytic role of Ni particles, which reduce the surface associated resistances and confirm the surface nature of the abovementioned LF and MF processes. Additionally, the decrease of the LF and MF resistances allows the observation of other, less resistive processes, at higher frequencies. In summary, the reduction of the LF process can be directly linked with the improvement in surface kinetics due to the Ni dispersion on the LSCCe grains.

When the amount of infiltrated Ni is increased (LSCCe+5Ni, Figure 9.24c and 9.24f) the MF contribution disappears and exclusively the HF contribution (HF, 25-100 kHz) remains. The HF contribution is ascribed to the transport properties of the LSCCe material or to the LSCCe/LWO interfacial resistances, which remain unchanged upon Ni infiltration.

As a summary, a complete comparison between the different chromite based anodes on LWO electrolytes studied in this chapter can be seen in Figure
9.29, where the Rp at 700 °C is plotted for LSC, LSCCe, LSCN10, LSCN, LSCCe+1Ni, LSCCe+5Ni and LSCN+5Ni. The graph depicts the gradual decrease of the Rp from the 347 Ω·cm² obtained for LSC to 4.85 and 3.55 Ω·cm² when the LSC material is firstly substituted by 10 and 20% Ni, respectively and 37.81 Ω·cm² obtained for 10% Ce doped LSC. This anode improvement is increased with the Ni infiltrations on both LSCCe and LSCN anodes, being LSCN+5Ni anode the best electrode (Rp = 0.47 Ω·cm²) with slightly better performance than LSCCe+5Ni (Rp = 0.76 Ω·cm²).

Figure 9.29. Rp at 700 °C in H₂+H₂O of different chromite based anodes: LSC, LSCCe, LSCN10, LSCN, LSCCe+1Ni, LSCCe+5Ni and LSCN+5Ni.
9.3. Conclusions

In conclusion, this chapter presents the large potential of chromite-based anodes for LWO-based PC-SOFC. Aside from the chemical compatibility of LSC, LSCN10, LSCN and LSCCe with LWO, the protonic conductivity of reduced anodes combined with sufficient $p$-type electronic conductivity is illustrated. These properties are requisites for protonic anodes and make it possible to avoid the blending with other protonic or electronic materials, as it typically happens with protonic composite cathodes for PC-SOFC and Ni-based cermet anodes. Furthermore, the mixed conductivity allows enlarging the surface area, i.e., TPB, available for the $\text{H}_2$ oxidation reactions and proton incorporation.

LSCN10 anode on LWO, in symmetrical cell configuration, demonstrates that under typical anode reducing conditions, structural Ni is segregated on the surface. Thus, the Ni nanoparticles become catalytically active for the $\text{H}_2$ oxidation reaction improving the anode performance regarding to pure LSC, i.e., $2.68 \, \Omega \cdot \text{cm}^2$ at $750 \, ^\circ \text{C}$ being principally limited below $850 \, ^\circ \text{C}$ by LF processes, ascribed to surface reactions. The surface properties of the chromite backbone structure (mixed $\text{H}^+/\text{e}^-$ conducting) can be boosted either by higher Ni or Ce doping and specially by Ni infiltration. This results in the coating of the electrode surface with Ni nanoparticles and consequently in a strong improvement of the anode performance. Specifically, the infiltrated anode with the highest Ni loading (LSCN+5N) only presents HF associated resistance and Rp as low as $0.26 \, \Omega \cdot \text{cm}^2$ at $750 \, ^\circ \text{C}$ in wet $\text{H}_2$. 
9.4. References


Mixed protonic-electronic conductors based on lanthanum chromites


Chapter 10. Conclusions and remarks

With the aim of finding new MIEC materials stable in the harsh working conditions found in the applications as ion transport membranes, SOFC and PC-SOFC electrodes, four classes of materials have been studied. The results presented in this thesis led to the following conclusions, which are divided by blocks.

**Fluorite structured \( \text{Ce}_{0.9}\text{Ln}_{0.1}\text{O}_{2-\delta} \) and \( \text{Ce}_{0.9}\text{Ln}_{0.1}\text{O}_{2-\delta} + 2 \text{ mol}\% \text{ Co} \)**

- 10% of trivalent lanthanides are fully soluble in the fluorite lattice of ceria. Eu and Yb maintain their trivalent state in oxidising conditions. The trivalent lanthanides produce a linear distortion of the cell and the introduction of a fixed number of extrinsic oxygen vacancies.
- Pr and Tb dopants are in a mixed tri- and tetravalent state. The ratio \( \text{Ln}^{3+}/\text{Ln}^{4+} \) depends on temperature and oxygen partial pressure.
- The addition of 2 mol% of \( \text{CoO} \) succeed as sintering aid for all the doped cerias. It helps the grain growth and causes the densification improvement.
- Raman spectroscopy results demonstrate that the introduction of lanthanides dopants produces a disorder due to the creation of oxygen vacancies. This disorder increases with the cell parameter.
- All the lanthanides improve the total conductivity with respect to pure ceria. Except of Tb and Pr doped ceria, all the compounds are predominantly ionic conductors, as deduced from the conductivity behaviour, which is independent on \( p\text{O}_2 \).
- Conductivity of Tb and Pr doped ceria show a dependency on \( p\text{O}_2^{-1/6} \) at the most oxidizing conditions due to the formation of oxygen vacancies as \( p\text{O}_2 \) reduces. The conductivity is still mainly ionic.
The addition of Co particularly improves the conductivity of Tb and Pr doped ceria due to the effect in the reducibility of the dopant (which occurs at low temperatures) and to the electronic conductivity promotion by either the creation of electronic paths through the grain boundary or the dissolution of part of the Co in the structure.

- Differences in conductivity of materials with the same concentration of ionic charge carriers are due to differences in the mobility of ions over the intrinsic oxygen vacancies created in the fluorite lattice.
- The highest ionic conductivity is obtained for the less distorted cells, *i.e.*, Gd and Nd, which represent an optimum balance between Coulomb interactions, steric effects and oxygen vacancy distribution.
- The lowest $E_a$ was found for materials with long cell parameters, up to a maximum.

**Co addition on Ce$_{0.9}$Pr$_{0.1}$O$_{2.4}$**

- The CoO$_x$ causes the grain size enlargement by accelerating the densification rate and grain growth.
- Co does not dissolve in the ceria lattice.
- Co has a catalytic effect on the reduction of ceria, which shifts the bulk ceria reduction temperature in H$_2$.
- The addition of Co improves the total conductivity of Pr-doped ceria, attributed to electronic pathways along the grain boundaries.
- The transport properties depend on the microstructure of the specimen. Higher sintering temperature produces larger grain size and the agglomeration of CoO$_x$ in microdomains.
- Low temperature sintering conditions produce small particle size and the homogeneous spreading of the Co additive along the grain boundaries.
- The highest conductivity has been obtained for samples sintered at 1000 °C, which are a trade off between grain size, Co distribution and densification.
Materials based on Ce$_{1-x}$Tb$_x$O$_{2.δ}$ and Ce$_{1-x}$Tb$_x$O$_{2.δ}$ + 2 mol% Co

- The cell parameter depends on the concentration of dopant following the Vegard’s rule.
- A constant ratio between Tb$^{4+}$/Tb$^{3+}$ is observed for all the concentrations at room temperature. The Co addition affects the oxidation state of Tb increasing the ratio Tb$^{4+}$/Tb$^{3+}$.
- Part of the CoO$_x$ dissolves in the fluorite lattice. The sintering at very high temperatures may cause the segregation in the grain boundaries.
- The addition of Co increases the material reducibility, i.e., it lowers the reduction temperature.
- The addition of Co improves the densification of the specimens
- Ce$_{1-x}$Tb$_x$O$_{2.δ}$ shows predominantly ionic or mixed ionic-electronic conductivity depending on the Tb concentration, Co addition and temperature range.
- Tb doping introduces new electronic levels in the ceria band gap, which allow the small polaron hopping between Tb$^{4+}$/Tb$^{3+}$. Low Tb content (10%) seem not to be enough for the hopping, while 50% Tb-doped ceria shows p-type electronic conductivity.
- High doping levels and low oxygen partial pressures may induce the oxygen vacancy association with the cations.
- The addition of Co further creates oxygen vacancies, so that improves the conductivity. The Co not incorporated in the structure may create electronic pathways along the grain boundaries.
- The characterization by XPS reveals the presence of Tb$^{4+}$, Tb$^{3+}$ and Co$^{2+}$ in the surface and thus oxygen vacancies. Ce$^{3+}$ is not observed as expected for oxidizing conditions (air).
- PIE experiments demonstrate the importance of oxygen vacancies and electrons on the surface exchange rate, $\mathcal{R}_o$. It is found to increase profoundly with increasing Tb concentration.
- $9_{\alpha}$ is limited by the dissociative adsorption step rather than the oxygen incorporation in the ceria lattice, which is two orders of magnitude higher.
- CT10 does not show any oxygen permeation flux, which is attributed to the mainly ionic conductivity since the low doping level still not allows the electron hopping.
- CT10Co shows an oxygen permeation flux up to 0.02 ml cm$^{-2}$ min$^{-1}$ at 1000 °C. This is a consequence of the electronic pathways introduced by the Co with regard to CT10.
- CT20Co increases the oxygen permeation up to 0.08 ml cm$^{-2}$ min$^{-1}$ at 1000 °C. The enhancement is ascribed to the higher oxygen vacancy concentration as well as the higher density of new electronic levels in the ceria bandgap and percolation pathways.
- CT20, CT50 and CT50Co membranes are not tested due to the impossibility to obtain dense specimens by pressing and firing.
- CTx and CTxC families are stable in CO$_2$ containing atmospheres, contrarily to the most widely used membrane material, the BSCF.

Materials based on Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ (X=Gd, Cr, Mg, Bi and Ce)

- Gd doped ceria has high ionic conductivity. The co-doping with other metals aims to promote the electronic conductivity. Gd, Cr, Mg, Bi and Ce as X dopant have been incorporated in the ceria structure.
- Electrical conductivity of LSM-Ce$_{0.8}$Gd$_{0.1}$X$_{0.1}$O$_{2-\delta}$ (X=Gd, Cr, Mg, Bi and Ce) composite has been proved to be sufficient for SOFC cathode applications. Doped ceria contributes the ionic conduction improving the triple phase boundary area of a LSM cathode.
- These materials have been tested as LSM-CGXO composite cathodes in symmetrical cells. The polarization resistance decreases with increasing the conductivity of the ceria based material.
- Among the dopants, Mg seems to boost both the oxygen ion transport and the surface process for oxygen reduction.
Conclusions

- However, the performance of these new co-doped ceria composite cathodes is not notably better than Ce_{0.8}Gd_{0.2}O_{1.9} material.
- EIS data may be fitted to an equivalent circuit comprising three ZARC elements in series with an inductance and a pure resistance.
- The use of Ce_{0.8}Gd_{0.1}Mg_{0.1}O_{2-δ} instead of Ce_{0.8}Gd_{0.2}O_{1.95} in the composite may be economically viable due to the lower price of the raw material.

Materials based on doped-CaTiO_3 perovskites

- The perovskite stability predictions made by Goldschmidt and octahedral factor are not experimentally accomplished for all the doped CaTiO_3.
- The stable perovskites have been electrochemically tested by DC-conductivity. All the dopants improve the total conductivity with respect to pure CaTiO_3.
- The best transport properties are observed for Fe and Fe-Mg doping in the B-site (named CTF and CTFM, respectively).
- DC conductivity up to 1000 °C demonstrated an increase of conductivity of CTFM over that of CTF upon 800 °C, which has been ascribed to the mobility of associated oxygen vacancies released at high temperature. Below 800 °C, the decrease in conductivity of CTFM is attributed to the oxygen vacancy association.
- CTF and CTFM present mainly ionic conductivity at high temperatures, while p-type electronic conductivity is ascertained in the low temperature range.
- XPS measurements demonstrate the presence of oxygen vacancies in the surface of CTFM, since Ti, Fe and Mg in tri- and divalent oxidation states of are found.
- PIE experiments indicate that CTF and CTFM show a similar oxygen exchange rate, $\tilde{\beta}_o$ up to 700 °C. At this temperature, the transport properties of both materials are similar, so that the possible effect of the oxygen vacancy increase in CTFM is not quantifiable.
CTF and CTFM present a comparatively high $\mathcal{R}_o$ with regard to YSZ and CGO10, known as solid ionic conductors, and LSC20 perovskite. $\mathcal{R}_o$ is comparable to that of CGO20 and CT50. On the other hand, CTF and CTFM $\mathcal{R}_o$ are lower than for BSCF.

Using a catalytic layer over the membrane surface may allow overcoming the limits imposed by the oxygen surface exchange rate.

CTFM membrane with a Pt catalytic layer delivers an oxygen permeation flux one order of magnitude higher than referenced CTF membranes, which is attributed to the temperature dependent increase of oxygen vacancies concentration and mobility.

The CTFM membrane stability is tested under CO$_2$ containing sweep gas, where it demonstrates to be stable.

CTFM is stable to harsh environment conditions involving 1000 ppm COS, 100 ppm HCN, 4% CO$_2$, 46% CO, 46% H$_2$ and 4% H$_2$O at 35 bar and up to 400 °C.

CTFM is not stable against a two weeks treatment on a continuous stream of CO$_2$ containing 5% O$_2$, 1% of SO$_2$ and 2.5% of H$_2$O at 800 °C.

**Composite materials based on doped-ceria/ spinels**

One-pot synthesis of fluorite/spinel composites has been demonstrated to be the most suitable method to obtain pure phases. One pot method produces narrow grain size distribution and homogeneous distribution of grains and phases.

60/40 spinel to fluorite phase volume ratio is the optimum for the creation of percolative ionic and electronic paths from side to side of the membrane.

From the tested spinels, only Fe$_2$NiO$_4$, Fe$_2$CrO$_4$ and Mn$_{1.8}$Cu$_{1.2}$O$_4$ are stable at high temperatures and to the contact with the fluorite. However, Fe$_2$CrO$_4$ and Mn$_{1.8}$Cu$_{1.2}$O$_4$ are not able to deliver dense specimens.

Fe$_2$NiO$_4$ (FNO) is the spinel of choice as it is capable to form dense composites without reacting with some of the fluorites. FNO only reacts with the Ce$_{0.8}$Gd$_{0.1}$Pr$_{0.1}$O$_{2.8}$.
Conclusions

- Stable composites are achieved by mixing FNO and Ce$_{0.8}$Gd$_{0.2}$O$_{1.9}$ (CGO), Ce$_{0.8}$Tb$_{0.2}$O$_{2.8}$ (CTO) and Ce$_{0.8}$Tb$_{0.2}$O$_{2.8}$ +2 mol% Co (CTOCo).
- At high temperatures, the FNO-CGO composite shows the highest conductivity, ascribed to the higher extrinsic ionic conductivity of CGO.
- At temperatures below 640 ºC, the conductivity of FNO-CTO overwhelms that of FNO-CGO, due to the improved electronic conductivity of the fluorite phase.
- The lowest conductivity is observed for FNO-CTOCo.
- Contrarily to single phase FNO, dense specimens are achieved by the combination with the fluorites, which are able to act also as sintering aid.
- A deeper study on FNO-CTO composite microstructure did not show evidence of impurities or reaction between phases up to the limits of the XRD and SEM devices.
- SEM analysis of FNO-CTO reveals a narrow grain size distribution and homogeneous allocation of grains. The existence of percolative paths from side to side of the dual-phase membranes necessary for the oxygen permeation is ascertained as well as a proper packing and bounding among the different grains in the membrane.
- The transport properties of FNO-CTO showed that the ambipolar transport involves (i) hopping of charge carriers, i.e., hopping of localized electrons between multivalent cations located in the octahedral Fe sublattice and (ii) the ionic transport in though the doped ceria lattice. Additionally, CTO presents significant $p$-type electronic conductivity although the ionic conductivity prevails.
- The oxygen permeation of FNO-CTO membrane has been measured and compared with the flux obtained by a FNO-CGO reference composite membrane, both using CO$_2$ as sweeping gas. Both membranes are stable against CO$_2$, but the oxygen permeation of FNO-CTO is twice the flux obtained by FNO-CGO.
- Higher $pO_2$ gradients across the membranes lead to substantial increases in the oxygen flux.
FNO-CTO composite is stable against an environment containing wet SO₂ and CO₂, at 800 ºC during 14 days. Contrarily, FNO-CGO and LSCF are not stable to the same treatment.

Mixed protonic-electronic conductors based on lanthanum chromites

- La₀.₈₇Sr₀.₁₃CrO₃ (LSC) has been doped both in A and B site in order to increase the mixed protonic-electronic conductivity.
- LSC, La₀.₈₅Sr₀.₁₅Cr₀.₉Ni₀.₁O₃ (LSCN10) and La₀.₈₅Sr₀.₁₅Cr₀.₈Ni₀.₂O₃ (LSCN20) show SrCrO₃ impurities that disappear in reducing conditions.
- Several dopants succeeded to form a pure perovskite at low temperatures, but they were not stable at the high temperatures required to obtain dense specimens, i.e., La₀.₈₅Sr₀.₁₅Cr₀.₉Fe₀.₁O₃ (LSCrF), La₀.₈₅Sr₀.₁₅Cr₀.₉Al₀.₁O₃ (LSCA), La₀.₈₅Sr₀.₁₅Cr₀.₉Ti₀.₁O₃ (LSCT) and La₀.₈₅Sr₀.₁₅Cr₀.₉Y₀.₁O₃ (LSCY).
- Other dopants only released a pure perovskite structure after sintering at high temperatures or in reducing conditions, i.e., La₀.₇₅Sr₀.₁₅Ce₀.₁CrO₃ (LSCCe).
- The highest conductivities in wet H₂ atmospheres were obtained for LSCCe and LSCY. LSCCe was chosen for further investigations.
- Densification of these materials is very difficult, so they have been focus on the application as anodes for PC-SOFC based on LWO electrolyte. Thus, the first stage of the study considered LSC, LSCN10, LSCN20 and LSCCe materials and the improvement of surface reactions by Ni infiltration.
- These materials do not react with the LWO electrolyte, contrarily to the widely used anode of NiO.
- Structural Ni in LSCN10 and LSCN20 is able to segregate from the structure to the surface in reducing conditions. Thus, Ni nanoparticles become catalytically active for the H₂ oxidation reaction. The more Ni in the structure the better performance of the anode. TEM images demonstrated the presence of the nanoparticles in the electrode surface.
- The improvement of LSCN10 and LSCN20 anodes with respect to the LSC is ascribed to the surface promotion by the Ni nanoparticles. They reduce the
overall resistance, but they are still limited by low frequency resistance (LF), especially at low temperatures.

- The improvement of LSCCe anode with respect to the LSC is ascribed to the improvement in the mixed protonic electronic conductivity. It reduces the overall resistance, although the anode operation is still limited by the LF, \textit{i.e.}, surface processes.

- The infiltration of Ni nanoparticles allows decreasing the LF of the corresponding anodes without infiltration, which suggests that the surface reaction of \( \text{H}_2 \) oxidation is catalyzed.

- The most concentrated infiltration (5Ni) completely eliminates the LF resistance in both anodes LSCN20 and LSCCe, and only high frequency (HF) processes remain.

- If the infiltration concentration decreases, as is the case of LSCCe+1Ni, the LF resistance disappears and the medium frequency (MF) resistance shifts to higher frequencies, combined with a new contribution at HF.

- The HF contribution is ascribed to the transport properties of the anode backbone and/or to the anode/electrolyte interfacial resistances, which remain unchanged upon Ni infiltration.

- The best anode performance is obtained for LSCNi20+5Ni. Polarization resistance is as low as 0.26 \( \Omega \cdot \text{cm}^2 \) at 750 °C in wet \( \text{H}_2 \).

\textbf{10.1. Final remarks}

The key objectives of the research described in this thesis were to find new materials capable to transport oxygen ions or protons and electronic carriers that are stable in the working conditions to which they are submitted. Several materials emerged from this work have achieved their purpose. \( \text{Ce}_{1-x}\text{Tb}_x\text{O}_{2-\delta} \) and \( \text{Ce}_{1-x}\text{Tb}_x\text{O}_{2-\delta} +2 \text{ mol\% Co} \) are proposed as a family of materials.
that may provide competent oxygen permeation fluxes and be stable against CO₂ containing atmospheres. Besides, the inclusion of this material as a component of a dual-phase membrane has increased the oxygen flux possibilities. The combination with a stable, Co and alkali-earth free spinel has delivered a promising membrane material for the integration in oxyfuel power plants. Furthermore, the studies about the behaviour of the materials and sintering properties may contribute to the next optimization of the microstructure and membrane architecture.

On the other hand, after a wide screening of doped CaTiO₃ perovskites, CTFM has been shown as a promising material offering good oxygen permeation fluxes and keeping stable against CO₂.

Finally, the study on lanthanum chromites as mixed protonic-electronic conductors allowed developing a new generation of anodes for LWO based PC-SOFC. Although deeper study on the transport properties of the doped materials and the optimization of the synthesis and sintering conditions is still required, milestone polarization resistance values have been reached with these anodes.
Scientific Contribution

Peer reviewed articles

C. Solís, M. Balaguer, F. Bozza, N. Bonanos and J. M. Serra, Catalytic surface promotion of highly performance La_{0.85}Sr_{0.15}Cr_{0.8}Ni_{0.2}O_{3-δ} anodes for La_{5.6}WO_{11.4-δ} based proton conducting fuel cells, *Applied Catalysis B* (2013), to be submitted.

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**Patent**


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Selected poster presentations

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List of Figures and Tables

List of figures

Figure 1.1. a) Frenkel defect: cation in interstitial hole, and b) Schottky disorder: missing pair of ions.............................................................................................................. 18

Figure 1.2. a) Polaron particle: a charge inside its self-created lattice distortion [12] and b) ionic diffusion mechanisms (1) via vacancy, (2) via interstitial site............................... 22

Figure 1.3. Electron (e), and hole (h), mobilities in oxides. The lower limit for large polarons corresponds to a mean-free path equal to the interatomic distance. The upper limit for small polarons corresponds to one hop every lattice vibration period. Note the opposite trends with temperature: for small polarons and ions, thermal activation is needed, so higher temperatures raise the rate; for large polarons, phonon scattering reduces the mobility at higher temperatures. [13]................................................ 23

Figure 1.4. Kroger-Vink diagram for acceptor doped material. The acceptor concentration represents the ionized acceptor concentration........................................... 31

Figure 1.5. Kröger-Vink diagram for donor doped material. The donor concentration represents the ionized donor concentration......................................................... 31

Figure 1.6. Oxygen permeation scheme for a dense membrane with catalytic layers for oxygen dissociation and recombination. ................................................................. 35

Figure 1.7. Schematic of an individual SOFC based on oxygen-ion conductors fed by H₂......................................................................................................................... 38

Figure 2.1. Scheme of the fabrication of the lanthanide doped ceria. The addition of Co by impregnation is indicated. The calcination temperature ensures the decomposition of the nitrates and carbonates from the precursors...................................................... 62

Figure 2.2. Main steps for rectangular probes fabrication. From left to right: initial powder, ball milling step, rinsing of the powder, pressing, high temperature calcination and final configuration of sintered samples................................................. 64
Figure 2.3. Main steps for membrane fabrication. From left to right: initial powder, ball milling step, rinsing and drying of the powder, uniaxial pressing, high temperature calcination and final configuration of sintered samples. Afterwards, a catalytic layer may be screen printed on the membrane surface according to the membrane surface.

Figure 2.4. Main steps for electrode fabrication. From left to right: initial powder, ball milling step, rinsing and drying of the powder, ceramic ink fabrication, screen printing of the electrode on the electrolyte surface and final electrode configuration after high temperature calcination. In addition, the electrode on the right is screen printed with a gold mesh for current collection.

Figure 2.5. Incident beam on a crystalline material and diffracted angle ($2\theta$), which is recorded.

Figure 2.6. Schematic diagram for grain size and crystallite of a crystalline powder sample. A particle or grain of a real crystalline powder generally consists of many crystallites, which can be considered as single crystals.

Figure 2.7. Four point configuration for DC conductivity measurements. The contacts are made with silver paste and wires.

Figure 2.8. a) Combinatorial DC conductivity setup, b) detail of the sample chamber containing the ten samples and c) from top to bottom, furnace controller, programmable current source (Keithley 2601) that and sixteen channel multimeter (Keithley 3706).

Figure 2.9. Experimental setup permeation scheme. The gas inlet and outlet may be reversed. The membrane is pressed by a spring sited in the lid.

Figure 2.10. Equivalent circuits a) simple parallel resistance-capacitor (RC) circuit and its corresponding perfect semicircle response and b) double RC circuit, represented as a double arc.

Figure 3.1. Fluorite cubic structure of CeO$_2$. Doping occurs by the Ln substitution of a Ce cation.

Figure 3.2. Thermogravimetry analysis of Ce$_{0.9}$Tb$_{0.1}$O$_{2.8}$ and Ce$_{0.9}$Er$_{0.1}$O$_{2.8}$, in air. The TG was performed up to 800 ºC, the calcination temperature of the as prepared materials.

Figure 3.3. XRD pattern of Ce$_{0.8}$Ln$_{0.2}$O$_{2.8}$ and Ce$_{0.8}$Ln$_{0.2}$O$_{2.8}$ + Co 2mol% calcined at 800 ºC.
Figure 3.4. Lattice parameter of doped ceria as a function of the trivalent ionic radius of the dopant cation (and tetravalent for Pr and Tb) in eight-fold coordination. 98

Figure 3.5. SEM images of Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ and Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ + Co 2 mol% calcined at 1300 °C (Ln = Tb, Pr, Eu and La). 100

Figure 3.6. HT-XRD heating up in air (left side) and cooling down in N<sub>2</sub> (right side) of Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ. In the insets, the calculated cell parameters (a 700 ºC) at 700 °C in air (left side) and in N<sub>2</sub> (right side) are plotted as a function of the trivalent ionic radii (and tetravalent for Pr and Tb). 102

Figure 3.7. a) Raman spectra of the Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ (Ln=La, Gd, Eu, Pr, Tb, Yb, Er) (sintered at 1300 °C) from 400-750 cm<sup>-1</sup> and b) F2g Raman shift as a function of the cell parameter Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ and Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ + Co calcined at 1300 °C and 800 °C (top) and after subtracting Grüneisen shifts (bottom). The inset shows a magnification of the band related to the oxygen vacancy formation. 104

Figure 3.8. Correlation length obtained from Raman line-shape fitting as a function of the cell parameter. 105

Figure 3.9. a) Arrhenius plot of the total conductivity in air of CeO<sub>2</sub> and Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ (Ln=Tb, Pr, Eu and Er), and b) log-log plot of total conductivity at 700 °C of the Co-free and Co-containing CeO<sub>2</sub> and Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ (Ln=Tb, Pr, Eu and Er) as a function of pO<sub>2</sub>. 108

Figure 3.10. a) Arrhenius plot of the total conductivity in air of Co-free Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ (Ln=Yb, Gd, La and Nd) and b) log-log plot of the total conductivity versus pO<sub>2</sub> at 700 °C of the Co-free and Co-containing CeO<sub>2</sub> and Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ Ce<sub>0.9</sub>Ln<sub>0.1</sub>O<sub>2</sub>-δ (Ln=Yb, Gd, La and Nd). 108

Figure 3.11: E<sub>a</sub> (top) and total conductivity at 700 °C in air (bottom) of the different doped materials versus cell parameter measured at 700 °C. Cell parameters at 700 °C in air have been calculated from HT-XRD or by using TEC values for similar dopants. E<sub>a</sub> solid lines have been fitted to Eq. 4 only taken into account samples with trivalent dopants while dashed fits include Pr and Tb. 111

Figure 4.1. a) XRD patterns and SEM images and b) temperature dependence of cell parameter of CP and CPCo sintered at 1300 °C. 125

Figure 4.2. a) TPR profile of CeO<sub>2</sub> and CP and b) TPR profile of CeO<sub>2</sub> + 2 mol% Co and CPCo. All the samples are samples sintered at 1300 °C. The flow gas is 10% H<sub>2</sub> in Ar; heating ramp was 5 °C/min. 127
Figure 4.3 SEM pictures of CPCo for different sintering temperature: 900, 1000, 1200, 1300 and 1450 °C. The grain size grows with the temperature from 60 nm at 900 °C to 7 µm at 1450 °C. ................................................................. 129

Figure 4.4. Back scattering SEM of CPCo sintered at a) 1000 °C and b) 1450 °C. c) Less magnified image of CPCo at 1450 °C and d) EDS Co mapping indicating Co rich zones. Arrows indicate the presence of the CoOx particles. .............................................. 130

Figure 4.5. FIB-lamella TEM of CPCo sintered at 1000 °C (left) and 1450 °C (right). 131

Figure 4.6. TEM-EDS on a 1000 °C CPCo grain. Co appears dispersed along the grain surface, while fine CP-CP grain interface is shown. ......................................................... 131

Figure 4.7. TEM-EDS on a 1450 °C CPCo grain boundary. A fine CP-CP grain interface is shown. Co was not detected either in the grain boundaries or in fringes and surfaces. ................................................................. 132

Figure 4.8. a) XRD patterns and b) TPD of oxygen in Ar for CPCo sintered at 1000, 1200 and 1450 °C. ........................................................................................................ 133

Figure 4.9. a) Total conductivity in air at 800 °C and 400 °C as a function of the sintering temperature and b) total conductivity of CPCo sintered at five different temperatures in air. For comparison, conductivity at 5x10⁻⁵ atm for CPCo sintered at 1000 °C is plotted. ........................................................................................................ 134

Figure 4.10. a) Arrhenius plot of the total conductivity of CPCo sintered at 1000 and 1300 °C and CP sintered at 1300 °C, in air. b) pO₂ dependence of the total conductivity of CPCo sintered at different temperatures, 400 °C (open symbols) and 800 °C (solid symbols). ................................................................. 136

Figure 5.1. XRD patterns of CTx and CTxCo sintered at 1450 °C recorded at room temperature. ........................................................................................................ 145

Figure 5.2. Scanning microscope pictures taken by backscattered electrons of CTx and CTxCo membrane cross sections, at 1200 °C................................................................. 145

Figure 5.3 Cell parameter dependence on the dopant amount for CTx and CTxCo and theoretical Vegard’s slope for Tb⁺³ and Tb⁺⁴ total substitution. .............................................. 146

Figure 5.4. a) H₂ TPR signal for CT10 and b) H₂ TPR for CTxCo (x=10 and 20). ..... 149
Figure 5.5. Crystallite size dependence on the cell parameter for CT10, CT20, CT10Co and CT20Co after sintering at 1300 ºC.............................................................. 149

Figure 5.6. Normalized TPD measured in Ar of CTx and CTxCo. Inset: CeO$_2$ and CeO$_2$+Co. The intensity (ion current) corresponds to $m/z = 32$. ........................................ 150

Figure 5.7. (a) Oxygen non-stoichiometry ($\delta$) determined by TG in air and (b) $\delta$ determined by TG in N$_2$, for CTx and CTxCo. Temperature range include from 430 to 1000 ºC. ................................................................. 152

Figure 5.8. Kröger-Vink diagram for acceptor doped fluorite. Four $pO_2$ regions delimit the main key defect reactions for a general fluorite. .................................................. 153

Figure 5.9. Arrhenius plot of the total conductivity in air and in Ar (0.21 and 5·10$^{-5}$ atm, respectively) for a) CTx and b) CTxCo................................................................. 157

Figure 5.10. Total conductivity dependency on Tb dopant concentration in (a) air, $pO_2 = 0.21$ atm and (b) Ar, $pO_2 = 2·10^{-5}$ atm. ......................................................... 159

Figure 5.11. Conductivity of CTx and CTxCo systems as a function of $pO_2$ measured at (a) 400 ºC and (b) 800 ºC................................................................. 160

Figure 5.12. UV-vis absorption spectra $h\nu$F(R)$^2$ versus photon energy of (a) CTx and (b) CTxCo................................................................. 163

Figure 5.13. a) Ce 3d XPS spectra of CT20 and CT20Co quenched in air from 800 ºC and b) Ce3d XPS separation of peaks of CT20 and CT20Co spectra. Pass energy was 30 eV................................................................. 164

Figure 5.14. XPS of the Tb 3d region of a) CT20 and b) CT20Co, quenched in air from 800 ºC, acquired with a pass energy of 30 eV................................................................. 165

Figure 5.15. XPS of the Co 2d region of CT20 (a) and CT20Co (b), quenched in air from 800 ºC, acquired with a pass energy of 30 eV. The Co 2d peak is shifted to higher binding energy than found in CoO (778 eV), suggesting Co lattice substitution........ 167

Figure 5.16. Arrhenius plot of the exchange rates of CT10, CT10Co, CT20, CT20Co, CT50, CT50Co................................................................. 168

Figure 5.17. Surface exchange rate as a function of the oxygen nonstoichiometry at 700 ºC; ........................................................................................................ 169
Figure 5.18. Inverse temperature dependence of the rates of dissociative adsorption (\(R_a\)), oxygen incorporation (\(R_i\)) of the terbium doped studied series. ......................................... 171

Figure 5.19. Oxygen permeation flux as a function of inverse temperature for CT10Co and CT20Co. The oxygen flux for similarly configuration BSCF and LSCF membranes is shown for comparison. ......................................................... 173

Figure 5.20. a) Ionic transport number and b) total, ionic and electronic conductivity of CT20Co measured across a 1.2 mm thick membrane under a nominal gradient of \(pO_2\) 0.21 \(\parallel\) \(5 \times 10^{-5}\) atm (air/Ar). \(J(O_2)\) measured using chromatography analytical data. .... 174

Figure 5.21. Thermogravimetric analysis under CO\(_2\) flow of CT10 and CT10Co in CO\(_2\), and BSCF. ................................................................................................................................. 175

Figure 6.1. Room temperature XRD patterns of the as-sintered doped cerias CG82, CGMg, CGCr, CGBi and CG91) at 800 ºC................................................................. 187

Figure 6.2. Comparison of the initial particle size of a) CG82 (as-sintered) and b) LSM (as-received) and the resulting powders after calcination at 1150 ºC: c) CG82 and d) LSM. ................................................................. 189

Figure 6.3. SEM images of several composite cathodes sintered at 1150 ºC: a) CG82-LSM, b) CG91-LSM and c) CGBi-LSM................................................................. 190

Figure 6.4. a) Arrhenius plot in air and b) \(pO_2\) dependence, of the total conductivity of ceria-based materials ................................................................. 192

Figure 6.5. Arrhenius plot of the electrode polarization resistance of LSM-ceria-based composites and the single LSM electrodes. LSFC and LSM-YSZ are shown for comparison................................................................. 194

Figure 6.6. Polarization resistance as a function of conductivity at three different temperatures (700, 750 and 800 ºC) ................................................................. 195

Figure 6.7. Cross section SEM images of a) CGCr-LSM, b) CG82-LSM and c) CG91-LSM electrodes on a dense CG82-Co electrolyte. The upper layer seen in a) corresponds to the gold current collecting mesh................................. 196

Figure 6.8. Electrochemical impedance spectra of symmetrical cells with different LSM-doped-ceria electrodes at 800 ºC under air: a) Nyquist and b) imaginary
impedance Bode plots. Logarithm of selected frequencies is provided on the Nyquist plot.

Figure 6.9. Resistances corresponding to the different processes obtained by fitting the spectra of symmetrical cells at 800 °C under air according to the equivalent circuit \(LR(R_AQ_A)(R_BQ_B)(R_CQ_C)\) shown in the inset. The corresponding logarithm of summit frequency is indicated on top of each bar.

Figure 7.1. CaTiO\(_3\) orthorhombic perovskite from the family A\(^{II}\)B\(IV\)O\(_3\).

Figure 7.2. Comparative \(pO_2\) dependence of total electrical conductivity for pure and doped CaTiO\(_3\) perovskites.

Figure 7.3. a) Room temperature XRD pattern of orthorhombic CTFM calcined at 1200 °C and b) SEM picture of the formed membrane.

Figure 7.4. TG analysis under 5 % CO\(_2\)/air of a) CTFM and CTF and b) BSCF. Heating rate was 10 °C·min\(^{-1}\).

Figure 7.5. a) DTG in air, b) TPD of oxygen, m/z=32, in He \((pO_2=10^{-5}\) atm\), c) cell parameters extracted from HT-XRD in air and d) cell parameters extracted from HT-XRD in N\(_2\), for CTFM. Heating rate was 10 °C·min\(^{-1}\).

Figure 7.6. a) Arrhenius plot of the conductivity of CTFM, CTF and CaTiO\(_3\) and b) logarithm of the total conductivity versus \(pO_2\) of CTFM and CTF at 1000, 800 and 400 °C.

Figure 7.7. Ca 2p spectrum of CTFM quenched from 800 °C in Ar. Pass energy was 30 eV.

Figure 7.8. Ti 2p spectrum of CTFM quenched from 800 °C in Ar and deconvolution of the spectrum. Ti\(^{4+}\) and Ti\(^{3+}\) components are highlighted. Pass energy was 30 eV.

Figure 7.9. a) Fe 2p XPS spectrum and b) Mg 2s and Fe3s XPS spectra of CTFM quenched from 800 °C in Ar. Pass energy was 30 eV.

Figure 7.10. O1s XPS spectra of CTFM quenched from 800 °C. Pass energy was 30 eV.

Figure 7.11. Arrhenius plot of CTF and CTFM (a) the overall surface exchange rate \(\mathcal{R}_0\) and (b) partial \(\mathcal{R}_a, \mathcal{R}_i\) and \(\mathcal{R}_b\).
Figure 7.12. Comparative Arrhenius plot of the CTFM and CTF \( \Re \), with literature data of widely used oxides measured by PIE as YSZ, CGO10, CGO20, CT50 or IEDP as LSC20 and BSCF. [30-32] .......................................................... 224

Figure 7.13. Oxygen permeation flow for CTFM membrane as a function of the temperature, with a sweep flow of 400 mL min\(^{-1}\) of Ar. Synthetic air and pure oxygen at a flow rate of 100 mL min\(^{-1}\) are used as feed gas. .................................................... 226

Figure 7.14. a) Oxygen permeation flux for CTFM membrane as a function of the Ar sweep flow at three different temperatures using synthetic air as feed and b) using air and \( \text{O}_2 \) as feed gas at 1000 \( ^\circ \text{C} \). ........................................................... 227

Figure 7.15. a) Oxygen permeation flow after 24 hours in \( \text{CO}_2/\text{Ar} \) atmosphere. Oxygen flux using Ar as sweep gas is represented using open symbols, b) Oxygen permeation flux in Ar before and after of \( \text{CO}_2 \) treatment. \( Q_{\text{Ar}}=400 \text{ mL\cdot min}^{-1} \) ........................................... 228

Figure 7.16. XRD patterns of as sintered CTFM powder and membranes after the stability treatments. Reference pattern and XRD of sample holder has been plotted for comparison. .......................................................... 229

Figure 8.1. Oxygen and electron transport scheme in a dual phase composite membrane composed of a) pure electronic and mixed ionic electronic conducting phases and b) pure electronic and ionic conducting phases. ............................................. 239

Figure 8.2. XRD patterns of tested spinels: \( \text{Fe}_2\text{NiO}_4 \), \( \text{Cr}_2\text{FeO}_4 \), \( \text{Co}_2\text{FeO}_4 \), \( \text{Mn}_2\text{CuO}_4 \). Grey circles indicate impurity peaks. ............................................................ 242

Figure 8.3. a) XRD patterns of the composites prepared by SS and OP, and the \( \text{Fe}_2\text{NiO}_4 \) and \( \text{Ce}_{0.8}\text{Gd}_{0.2}\text{O}_{1.9} \) single phases. b) Arrhenius plot of total electrical conductivity for the 50/50 phase ratio composites prepared by SS and OP and for 60/40 phase ratio OP composite. c) SEM picture of 50/50 FNO-CGO prepared by PM and d) SEM picture of 50/50 FNO-CGO prepared by OP. ............................................................. 243

Figure 8.4. XRD patterns of FNO-CLnO synthesized by OP and sintered in air at 1400 °C. Reference fluorite and spinel structure peaks are also indicated (ICDD 00-043-1002 and 00-044-1485, respectively). ................................................................. 244

Figure 8.5. Arrhenius plot of the total electrical conductivity in air of FNO-CLnO composite series. ........................................................................................................ 246

Figure 8.6. a) BS-SEM picture of the fractured cross-section of a FNO-CTO composite membrane sintered at 1400 °C in air; b) detail of the grain boundary between spinel
(dark) and fluorite (light) phases; c) detail of two fluorite grains fringe; d) detail of two spinel grains fringe and; e) CGO-LSM-Pd catalytic layer on the composite.

Figure 8.7. a) Total electrical conductivity recorded in air as a function of temperature for the single phase CTO, FNO and for the composite, b) Arrhenius plot of the total electrical conductivity of the composite in air and Ar atmospheres and c) total electrical conductivity at 800 °C as a function of the $pO_2$.

Figure 8.8. SEM image of Fe$_2$NiO$_4$ sample sintered at 1400 °C in air dwelling 10 h. ~90% relative density was achieved at these conditions.

Figure 8.9. a) Oxygen permeation flux as a function of temperature and sweep gas (Ar or CO$_2$) for FNO-CTO membrane and in Ar for single phase CTOCo; and b) flux stability with time on stream for FNO-CTO membrane in CO$_2$ sweeping at 900 °C.

Figure 8.10. Oxygen flux as a function of reciprocal temperature for different $pO_2$ gradients across the membrane, i.e., air-Ar, $O_2$-Ar and air-10% CH$_4$/Ar, on a 0.68 mm thick FNO-CTO membrane sintered at 1400 °C.

Figure 8.11. XRD pattern of both sides of the FNO-CTO membranes after 7 days of permeation experiments in Ar, CO$_2$, CH$_4$ sweeping.

Figure 8.12. Temperature dependency of oxygen permeation of FNO-CTO and FNO-CGO composites. The membranes were fed by air and the sweep gas was 100% CO$_2$.

Figure 8.13. XRD pattern of FNO-CTO and FNO-CGO composites after 7 days of treatment under a continuous flux consisted of 1% SO$_2$-5% O$_2$ in CO$_2$, at 800 °C.

Figure 9.1. XRD patterns of LSC calcined at 900 °C, LSCN10 and LSCN at 1050 °C and LSCrF at 1000 °C. The dotted lines represent the LSC reference pattern (ICSD 00-032-1240).

Figure 9.2. XRD patterns of A-site doped LSC calcined at 1600 °C and corresponding SEM microstructure images of (top to bottom) SmSC, NSC and LSCCe.

Figure 9.3. XRD patterns and corresponding SEM images of B-site doped LSC pressed and calcined at 1600 °C: LSCY, LSCT, LSCA and LSCrF. The peak positions of LaCrO$_3$ (LC, ICSD 00-024-1016) and LSC (ICSD 00-032-1240) perovskites are represented for comparison.
Figure 9.4: XRD patterns of (from bottom to top): LSC, LWO and 50 vol.% mixture at RT, treated at 1400 ºC in H₂ for 48 h and treated at 1150 ºC in air for 5 h. ................................ 275

Figure 9.5: XRD patterns of (from bottom to top): LWO and NiO 50 vol.% mixture at RT, treated at 900 ºC in H₂ for 48 h and treated at 1150 ºC in air for 5 h; and LSCN10 and LWO 50 vol.% mixture after calcination in air at 1150 ºC for 5 h. ......................... 276

Figure 9.6: a) SEM image and b) TEM image of a FIB-lamella of a LSCN10 anode on LWO electrolyte sintered at 1150 ºC. ........................................................................ 277

Figure 9.7: TEM images of as sintered LSCN10 on LWO electrolyte reveal the good bounding between electrolyte and porous electrode particles a) and LSCN10 powder after reduction at 900 ºC b). ........................................................................ 278

Figure 9.8: a) TPR analysis of LaCrO₃₋ₓ, LaCr₀.₉Ni₀.₁O₃₋ₓ and LSCN10 materials and b) Rp of LSCN10 as a function of time at 900 ºC in 100% H₂. ........................................... 279

Figure 9.9: Total conductivity of La₀.₈Sr₀.₁Cr₀.₉Ni₀.₁O₃₋ₓ in 5% H₂, 5% D₂, 5% H₂+H₂O and 5% D₂+D₂O. .................................................................................. 280

Figure 9.10: a) Nyquist and b) Bode plots of impedance spectra recorded in wet (2.5% vol. H₂O) 100% H₂ for LSC at 700 ºC and c) Nyquist and d) Bode plots of impedance spectra recorded in wet (2.5% vol. H₂O) 100% H₂ at different temperatures for LSCN10. .............................................. 281

Figure 9.11: Equivalent circuit used for EIS impedance modeling. CPS are used instead of perfect capacitors due to the porous nature of the anode. ................................................................. 282

Figure 9.12: a) Modeled LF and MF resistances as a function of the temperature and b) as a function of the pH₂. .............................................................................................. 283

Figure 9.13: XRD pattern in logarithm scale of LSCN10 anode after electrochemical test. The materials maintains its pure perovskite structure. .................................................. 284

Figure 9.14: a) XRD pattern and b) SEM image of LSCN powder sintered at 900 ºC. 285

Figure 9.15: XRD patterns of LSC, LSCN and LSCN+5Ni symmetrical cells. The peaks corresponding to metallic Au from the current collector of the symmetrical cell and metallic Ni (infiltrated and structural) are indicated. .................................................. 286
5 M water solution of Ni Nitrate was dropped on the surface of the LSCN electrode and allowed to penetrate into the pores for 30 min. Each infiltration step was followed by calcination at 300 °C to decompose the nitrates.

5 M water solution of Ni Nitrate was dropped on the surface of the LSCN electrode and allowed to penetrate into the pores for 30 min. Each infiltration step was followed by calcination at 300 °C to decompose the nitrates.

Figure 9.16. Schematic diagram of the LSCN anode a) and its corresponding SEM image b) and scheme of the LSCN+5Ni anode c) and its real SEM images d) and e). 5 M water solution of Ni Nitrate was dropped on the surface of the LSCN electrode and allowed to penetrate into the pores for 30 min. The infiltration step was repeated 4 times. Each infiltration step was followed by calcination at 300 °C to decompose the nitrates.

Figure 9.17. TEM image of Ni nanoparticles obtained by firing nitrate precursor at 800 °C in H2.

Figure 9.18. Rp as a function of inverse temperature for LSC, LSCN and LSCN+5Ni electrodes.

Figure 9.19. Impedance spectra recorded at 700 °C in wet hydrogen of LSC a) Nyquist and b) Bode plots and LSCN and LSCN+5Ni c) Nyquist and d) Bode plots.

Figure 9.20. Temperature dependency of modelled resistances a) and capacitances b) of LSC, LSCN and LSCN+5Ni samples.

Figure 9.21. XRD patterns of LSCCe as prepared powder and anode after Ni infiltration and reduction in wet H2 up to 800 °C. XRD positions of LaCrO3, LWO and CeO2 are indicated as reference.

Figure 9.22. TG analysis of the LSCCe in Ar (85%)-CO2 (15%). The temperature was increased at a rate of 5 °C/min.

Figure 9.23. a) LSCCe total conductivity in H2+H2O and D2 +D2O and LSC in H2+H2O, b) pO2 dependence of the total conductivity at 700 °C of LSCCe in dry atmosphere and c) pO2 dependence of the total conductivity at 700 °C of LSCCe wet atmospheres.

Figure 9.24. Schematic diagrams of the LSCCe a), LSCCe+1Ni b) and LSCCe+5Ni c) anodes and the corresponding SEM images d), e) and f). 1 M and 5 M water solution of Ni Nitrate (LSCCe+1Ni and LSCe+5Ni, respectively) were dropped on the surface of the electrodes and allowed to penetrate into the electrodes pores for 30 min. The infiltration step with the 5 M solution was repeated 4 times. Each infiltration step was followed by calcination at 300 °C to decompose the nitrates.

Figure 9.25. Rp as a function of the inverse temperature of the four anodes La0.65Sr0.15CrO3-δ (LSC), La0.75Ce0.15Sr0.15Cr O3-δ (LSCCe) and LSCCe infiltrated with 1Ni (LSCCe+1Ni) and with 5Ni (LSCCe+5Ni).

Figure 9.26. EIS spectra at 700 °C in H2 of LSC and LSCCe anodes, a) Nyquist and b) Bode plots and EIS spectra at 700 °C in H2 of LSCCe infiltrated with 1Ni and 5Ni, c) Nyquist and d) Bode plots.
Figure 9.27. Modeled R contributions of the LSCCe anode and when infiltrated with different amounts of Ni: 1Ni (LSCCe+1Ni) and with 5Ni (LSCCe+5Ni).

Figure 9.28. Associated capacitances and relaxation times of the LSCCe, LSCCe+1Ni and LSCCe+5Ni anodes.

Figure 9.29. Rp at 700 ºC of different chromite based anodes: LSC, LSCCe, LSCN10, LSCN, LSCCe+1Ni, LSCCe+5Ni and LSCN+5Ni.

List of tables

Table 1.1. Summary of electroneutrality equations in different regions.
Table 1.2. Region dependent equation for defect species concentration in acceptor doped material.
Table 1.3. Region dependent equation for defect species concentration in donor doped material.
Table 2.1. Summary of the seven crystal systems and Bravais lattices.
Table 3.1. Theoretical and experimental density of doped cerias.
Table 3.2. Thermal expansion coefficients calculated in Air and in N₂ for different La, Eu Gd, Pr and Tb doped ceria.
Table 4.1 Thermal expansion coefficient of Ce₀.₉Pr₀.₁O₂₋δ and CPCo 2% sintered at 1300 ºC.
Table 5.1. Apparent activation energies in the temperature ranges of 800-480 ºC (high T) and 480-400 ºC (low T) for CT10, CT10Co, CT20 and CT20Co and in the temperature range of 800-700 ºC (high T) and 700-400 ºC (low T) for CT50 and CT50Co. The conductivities at 800 ºC under air and 5x10⁻⁷ atm is indicated.
Table 5.2: Oxygen non-stoichiometry and surface exchange rate, at 700 °C, and apparent activation energies for the surface exchange rate for the different compounds (Fig 14).

Table 6.1 Cerium-based compositions synthesized as cubic fluorite structure with the nomenclature used along the text, and the corresponding lattice parameter and crystallite size extracted from XRD patterns.

Table 6.2. Fitting parameters obtained for each LSM-ceria-based composite at 800 °C under air. R is the resistance, $Q_0$ and $n$ are the constant phase element parameters and $\omega$ the summit frequency of each ZARC element, respectively. Capital letters (A, B, C) indicate the process according to the proposed electrical equivalent circuit.

Table 7.1. Ionic radius of constituent ions A and B, tolerance factor, octahedral factor ($r_B/r_O$) and symmetry of different compounds (O=orthorrombic, X= not single phase).

Table 7.2. Cell volume and cell parameters of formed perovskite compounds (fitting errors in brackets).

Table 8.1. Spinel and fluorite structured compounds selected for dual-phase membranes.

Table 9.1. Summary of doped lanthanum strontium chromites composition and short nomenclature. The main structural phase formation is indicated as well as the total conductivity in wet H$_2$ atmosphere at 700 °C, and the corresponding activation energy.
Appendix

Appendix 1

Summary of the Kröger-Vink notation of a compound MOₓ.

<table>
<thead>
<tr>
<th>Sites</th>
<th>Notation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interstitial oxygen ion</td>
<td>Oᵢ⁻</td>
</tr>
<tr>
<td>Foreign acceptor metal, A on M site</td>
<td>Aₘ⁺</td>
</tr>
<tr>
<td>Free electrons</td>
<td>e⁻</td>
</tr>
<tr>
<td>Occupied oxygen site</td>
<td>Oₐ⁺</td>
</tr>
<tr>
<td>Oxygen vacancy</td>
<td>Vₒ⁻</td>
</tr>
<tr>
<td>Foreign donor metal, D on M site</td>
<td>Dₘ⁻</td>
</tr>
<tr>
<td>Free positive holes</td>
<td>h⁺</td>
</tr>
<tr>
<td>Vacancy in the metal</td>
<td>Vₘᵢ⁻</td>
</tr>
</tbody>
</table>
Appendix 2

Information of plane spacing for the seven Bravais crystal systems.

Cubic \[ \frac{1}{d^2} = \frac{h^2 + k^2 + l^2}{a^2} \]

Tetragonal \[ \frac{1}{d^2} = \frac{h^2 + k^2}{a^2} + \frac{l^2}{c^2} \]

Hexagonal \[ \frac{1}{d^2} = \frac{1}{4} \left( \frac{h^2 + h k + k^2}{a^2} + \frac{l^2}{c^2} \right) \]

Trigonal \[ \frac{1}{d^2} = \frac{(h^2 + k^2 + l^2) \sin^2 \alpha + 2(hk + kl + hl)\cos^2 \alpha - \cos \alpha}{a^2(1 + 3 \cos \delta + 2 \cos \beta)} \]

Orthorhombic \[ \frac{1}{d^2} = \frac{h^2}{a^2} + \frac{k^2}{b^2} + \frac{l^2}{c^2} \]

Monoclinic \[ \frac{1}{d^2} = \frac{1}{\sin \beta} \left( \frac{h^2}{a^2} + \frac{k^2 \sin^2 \beta}{b^2} + \frac{l^2}{c^2} - \frac{2hl \cos \beta}{ac} \right) \]

Triclinic \[ \frac{1}{d^2} = \frac{1}{V} \left( S_{11} h^2 + S_{22} k^2 + S_{33} l^2 + 2S_{12} hk + 2S_{13} kl + 2S_{23} hl \right) \]

On the triclinic system, \( V \) is the volume of a unit cell and the coefficients are given below.

\[
S_{11} = b^2 c^2 \sin^2 \alpha, \quad S_{12} = ab^2 (\cos \alpha \cos \beta - \cos \gamma), \\
S_{22} = a^2 c^2 \sin^2 \beta, \quad S_{23} = a^2 bc (\cos \beta \cos \gamma - \cos \alpha), \\
S_{33} = a^2 b^2 \sin^2 \gamma, \quad S_{13} = ab^2 c (\cos \gamma \cos \alpha - \cos \beta)
\]
## Appendix 3

<table>
<thead>
<tr>
<th>Symbol</th>
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<th>Units</th>
</tr>
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<tr>
<td>( A_{\text{effective}} )</td>
<td>Effective area of a membrane</td>
<td>cm²</td>
</tr>
<tr>
<td>( \Delta \mu_{O_2} )</td>
<td>Total oxygen chemical potential difference</td>
<td></td>
</tr>
<tr>
<td>( \nabla \mu_{O_2} )</td>
<td>Oxygen chemical potential gradient</td>
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<tr>
<td>( a )</td>
<td>Cell parameter of a crystal</td>
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<td>( a )</td>
<td>Conductivity probe width</td>
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<td>( b )</td>
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<tr>
<td>( C )</td>
<td>Capacitor</td>
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<td>( d )</td>
<td>((h k l)) interplanar space</td>
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<tr>
<td>( E_a )</td>
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</tr>
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<td>( E_{\text{ass}} )</td>
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<tr>
<td>( E_b )</td>
<td>Binding energy (characteristic of each element)</td>
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<td>( E_f )</td>
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<td>( E_g )</td>
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<tr>
<td>( E_m )</td>
<td>Migration barrier</td>
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<td>( E_w )</td>
<td>Work function of the spectrometer</td>
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<tr>
<td>( h, k, l )</td>
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<tr>
<td>( h\nu )</td>
<td>Energy of the photons</td>
<td>eV</td>
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<tr>
<td>( J_{O_2} )</td>
<td>Permeation flux</td>
<td>mL min⁻¹ cm⁻²</td>
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<tr>
<td>( l )</td>
<td>Distance between electrodes in the conductivity probe</td>
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<tr>
<td>( L )</td>
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<td>( N )</td>
<td>Defect concentration</td>
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<tr>
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<tr>
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<td>Flow</td>
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<td>( q )</td>
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<tr>
<td>( R )</td>
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<td>Ohm</td>
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<tr>
<td>( t )</td>
<td>Crystallite diameter</td>
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<tr>
<td>( z )</td>
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<td>( Z )</td>
<td>System impedance</td>
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<tr>
<td>( \beta )</td>
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<td>( \Gamma )</td>
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<td>Unit</td>
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<tr>
<td>--------</td>
<td>-------------</td>
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</tr>
<tr>
<td>δ</td>
<td>Oxygen nonstoichiometry</td>
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<tr>
<td>Δω</td>
<td>Raman frequency shift</td>
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</tr>
<tr>
<td>θ</td>
<td>Incidence angle of X-ray</td>
<td>°</td>
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<tr>
<td>λ</td>
<td>Wavelength of X-rays</td>
<td>nm</td>
</tr>
<tr>
<td>μ</td>
<td>Carrier mobility</td>
<td>cm² V⁻¹ s⁻¹</td>
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<tr>
<td>σₐₘᵇ</td>
<td>Ambipolar conductivity</td>
<td>S cm⁻¹</td>
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<tr>
<td>σₑ</td>
<td>Electronic conductivity</td>
<td>S cm⁻¹</td>
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Appendix 4

<table>
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<td>Auger Electrons</td>
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<tr>
<td>ASR</td>
<td>Area Specific Resistivity</td>
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<td>Air Separation Unit</td>
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<td>Backscattered electrons</td>
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<td>CEC</td>
<td>Chemical Expansion Coefficient</td>
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<td>Catalytic Membrane Reactor</td>
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Resumen / Resum / Summary
Resumen

Los materiales conductores mixtos de electrones e iones (oxígeno o protones) son capaces de separar oxígeno o hidrógeno de los gases de combustión o de corrientes de reformado a alta temperatura. La selectividad de este proceso es del 100%. Estos materiales, óxidos sólidos densos, pueden usarse en la producción de electricidad a partir de combustibles fósiles, así como formar parte de los procesos que forman parte del sistema de captura y almacenamiento de CO$_2$. Las membranas de transporte de oxígeno (MTO) se pueden utilizar en las plantas energéticas con procesos de oxicombustión, así como en reactores catalíticos de membrana (RCM), mientras que las membranas de transporte de hidrógeno (MTH) se aplican en procesos de precombustión. Además, estos materiales encuentran aplicación en componentes de sistemas energéticos, como electrodos o electrolitos de pilas de combustible de óxido sólido, de ambas clases iónicas y protónicas (SOFC y PC-SOFC).

Los procesos mencionados implican condiciones de operación muy severas, como altas temperaturas y grandes gradientes de presión parcial de oxígeno ($pO_2$), probablemente combinadas con la presencia de CO$_2$ y SO$_2$. Los materiales más que mayor rendimiento de separación presentan y más ampliamente investigados en este campo son inestables en estas condiciones. Por tanto, existe la necesidad de encontrar nuevos materiales inorgánicos estables que proporcione alta conductividad electrónica e iónica.

La presente tesis propone una búsqueda sistemática de nuevos conductores iónicos-electrónicos mixtos (MIEC, del inglés) con diferente estructura cristalina y/o diferente composición, variando la naturaleza de los elementos y la estequiometría del cristal. La investigación ha dado lugar a
materiales capaces de transportar iones oxígeno, protones o cargas electrónicas y que son estables en las condiciones de operación.

La caracterización de una amplia serie de cerias (CeO$_2$) dopadas con lantánidos proporciona una comprensión general de las propiedades estructurales y de transporte, así como la relación entre ellas. Además, se estudia el efecto de la adición de cobalto a dicho sistema. Se ha completado el análisis con la optimización de las propiedades de transporte a partir de la microestructura. Todo esto permite hacer una clasificación inicial de los materiales basada en el comportamiento de transporte principal y permite adecuar la estructura y las condiciones de operación para obtener las propiedades deseadas para cada aplicación.

Algunos de los materiales extraídos de este estudio alcanzaron las expectativas. Las familias de materiales basadas en Ce$_{1-x}$Tb$_x$O$_{2-δ}$ y Ce$_{1-x}$Tb$_x$O$_{2-δ}$ +2 mol% Co proporcionan flujos de oxígeno bajos pero competitivos, ya que son estables en atmósferas con CO$_2$. Además, la inclusión de estos materiales en membranas de dos fases aumenta el flujo de oxígeno. La combinación con una espinela libre de cobalto y de metales alcalinotérreos como es el Fe$_2$NiO$_4$, ha dado lugar a un material prometedor en cuanto a flujo de oxígeno y estabilidad en CO$_2$ y en SO$_2$, que podría ser integrado en el proceso de oxicombustión.

Por otra parte, se ha añadido metales como codopantes en el sistema Ce$_{0.9-x}$M$_x$Gd$_{0.1}$O$_{1.95}$. Estos materiales, en combinación con la perovskita La$_{1-x}$Sr$_x$MnO$_3$ usada comúnmente como cátodo de SOFC, han sido capaces de disminuir la resistencia de polarización del cátodo. La mejora es consecuencia de la introducción de conductividad iónica por parte de la ceria.

Las perovskitas dopadas basadas en CaTiO$_3$ forman el segundo grupo de materiales investigados. La dificultad de obtener perovskitas estables y que
presenten conducción mixta iónica y electrónica se ha hecho evidente. De entre los dopantes utilizados, el hierro y la combinación hierro-magnesio han sido los mejores candidatos. Ambos materiales presentan conductividad principalmente iónica a alta temperatura, mientras que a baja predomina la conductividad electrónica tipo p. \( \text{CaTi}_{0.73}\text{Fe}_{0.18}\text{Mg}_{0.09}\text{O}_{3.8} \) se ha mostrado como un material competente en la fabricación de membranas de oxígeno, que proporciona flujos adecuados a la par que estabilidad en CO₂.

Finalmente, la perovskita \( \text{La}_{0.87}\text{Sr}_{0.13}\text{CrO}_3 \) (LSC) ha sido dopada con el objetivo de aumentar la conductividad mixta protónica electrónica. Este estudio ha llevado al desarrollo de una nueva generación de ánodos para PC-SOFC basadas en electrolitos de LWO. Las perovskitas dopadas con Ce en el sitio del La (LSCCe) y con Ni en el sitio del Cr (LSCN) son estables en condiciones de operación reductoras, así como en contacto con el electrolito. El uso de ambos materiales como ánodo disminuye la resistencia de polarización con respecto al LSC. El LSCCe está limitado por los procesos que ocurren a baja frecuencia (BF), relacionados con los procesos superficiales, y que son atenuados en el caso del LSCN debido a la formación de nanopartículas de Ni metálico en la superficie. La infiltración posterior con nanopartículas de Ni permite disminuir la resistencia a BF lo que sugiere que la reacción superficial de oxidación del H₂ está siendo catalizada. La infiltración más concentrada en Ni (5Ni) elimina completamente la resistencia a BF en ambos ánodos, de forma que los procesos que ocurren a altas frecuencias son ahora limitantes. El ánodo constituido por \( \text{LSCNi}_{20+5}\text{Ni} \) dio una resistencia de polarización de 0.26 Ω·cm² at 750 ºC en H₂ húmedo.
Resum

Els materials conductors mixts d'electrons i ions (oxigen o protons) són capaços de separat oxigen o hidrogen dels gasos de combustió o de corrents de reformat a alta temperatura. La selectivitat d'aquest procés és del 100%. Aquests materials, òxids sòlids densos, poden usar-se en la producció d'electricitat a partir de combustibles fòssils, així com formar part dels processos que formen part del sistema de captura i emmagatzematge de CO₂. Les membranes de transport d'oxigen (MTO) es poden utilitzar en les plantes energètiques amb processos de oxicombustió, així com en reactors catalítics de membrana (RCM), mentre que les membranes de transport d'hidrogen (MTH) s'apliquen en processos de precombustió. A més, aquests materials troben aplicació en components de sistemes energètictes, com a elèctrodes o electròlits de piles de combustible d'òxid sòlid, d'ambdues classes iòniques i protòniques (SOFC i PC-SOFC).

Els processos esmentats implicuen condicions d'operació molt severes, com ara altes temperatures i grans gràdents de pressió parcial d'oxigen (\(pO_2\)), probablement combinades amb la presència de CO₂ i SO₂. Els materials que major rendiment de separació presenten i són més àmpliament investigats en aquest camp són instables en aquestes condicions. Per tant, existeix la necessitat de trobar nous materials inorgànics estables que proporcionen alta conductivitat electrònica i iònica.

La present tesi proposa una cerca sistemàtica de nous conductors iònics-electrònics mixts (MIEC, de l'anglès) amb diferent estructura cristal·lina i/o diferent composició, variant la naturalesa dels elements i l'estequiometria del cristall. La recerca ha donat lloc a materials capaços de transportar ions oxigen,
protons o càrregues electròniques i que són estables en les condicions d'operació.

La caracterització d'una àmplia sèrie de cerias (CeO₂) dopades amb lantànids proporciona una comprensió general de les propietats estructurals i de transport, així com la relació entre elles. A més, s'estudia l'efecte de l'addició de cobalt a aquest sistema. S'ha completat l'anàlisi amb l'optimització de les propietats de transport a partir de la microestructura. Tot açò permet fer una classificació inicial dels materials basada en el comportament de transport principal i permet adequar l'estrUCTura i les condicions d'operació per a obtenir les propietats desitjades per a cada aplicació.

Alguns dels materials extrets d'aquest estudi van aconseguir les expectatives. Les famílies de materials basades en Ce₁₋ₓTbₓO₂₋δ i Ce₁₋ₓTbₓO₂₋δ₊2 mol% Co proporcionen fluxos d'oxigen baixos però competitius, ja que són estables en atmosferes amb CO₂. A més, la inclusió d'aquests materials en membranes de dues fases augmenta el flux d'oxigen. La combinació amb una espinela lliure de cobalt i de metalls alcalinoterris com és el Fe₂NiO₄, ha donat lloc a un material prometedor quant a flux d'oxigen i estabilitat en CO₂ i en SO₂, que podria ser integrat en el procés de oxicombustió.

D'altra banda, s'ha afegit metalls com codopants en el sistema Ce₀.₉ₓMₓGd₀.₁O₂₋δ. Aquests materials, en combinació amb la perovskita La₁₋ₓSrₓMnO₃ usada comunament com a càtode de SOFC, han sigut capaços de disminuir la resistència de polarització del càtode. La millora és conseqüència de la introducció de conductivitat iònica per part de la ceria.
Les perovskites dopades basades en CaTiO$_3$ formen el segon grup de materials investigats. La dificultat d'obtenir perovskites estables i que presenten conducció mixta iònica i electrònica s'ha fet evident. D'entre els dopants utilitzats, el ferro i la combinació ferro-magnesi han sigut els millors candidats. Tots dos materials presenten conductivitat principalment iònica a alta temperatura, mentre que a baixa predomina la conductivitat electrònica tipus $p$. CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_{3.8}$ s'ha mostrat com un material competent en la fabricació de membranes d'oxigen, que proporciona fluxos adequats a l'una que estabilitat en CO2.

Finalment, la perovskita La$_{0.87}$Sr$_{0.13}$CrO$_3$ (LSC) ha sigut dopada amb l'objectiu d'augmentar la conductivitat mixta protònica electrònica. Aquest estudi ha portat al desenvolupament d'una nova generació d'ànodes per a PC-SOFC basades en electròlits de LWO. Les perovskites dopades amb Ce en el lloc del La (LSCCe) i amb Ni en el lloc del Cr (LSCN) són estables en condicions d'operació reductores, així com en contacte amb l'electròlit. L'ús de tots dos materials com a ànode disminueix la resistència de polarització pel que fa al LSC. El LSCCe està limitat pels processos que succeixen a baixa freqüència (BF), relacionats amb els processos superficiais, i que són atenuats en el cas del LSCN a causa de la formació de nanopartícules de Ni metàl·lic en la superfície. La infiltració posterior amb nanopartícules de Ni permet disminuir la resistència a BF, el que suggereix que la reacció superficial d'oxidació de l'H$_2$ està sent catalitzada. La infiltració més concentrada en Ni (5Ni) elimina completament la resistència a BF en tots dos ànodes, de manera que els processos que ocorren a altes freqüències són ara limitants. L'ànode constituït per LSCNi20+5Ni va donar una resistència de polarització de 0.26 Ω·cm$^2$ a 750 ºC en H$_2$ humit.
Summary

Mixed ionic (oxygen ions or protons) and electronic conducting materials (MIEC) separate oxygen or hydrogen from flue gas or reforming streams at high temperature in a process 100% selective to the ion. These solid oxide materials may be used in the production of electricity from fossil fuels (coal or natural gas), taking part of the CO₂ separation and storage system. Dense oxygen transport membranes (OTM) can be used in oxyfuel combustion plants or in catalytic membrane reactors (CMR), while hydrogen transport membranes (HTM) would be applied in precombustion plants. Furthermore, these materials may also be used in components for energy systems, as advanced electrodes or electrolytes for solid oxide fuel cells (SOFC) and proton conducting solid oxide fuel cells (PC-SOFC) working at high and moderate temperature.

The harsh working conditions established by the targeted processes include high temperatures and low O₂ partial pressures (pO₂), probably combined with CO₂ and SO₂ containing gases. The instability disadvantages presented by the most widely studied materials for these purposes make them impractical for application to gas separation. Thus, the need to discover new stable inorganic materials providing high electronic and ionic conductivity is still present.

This thesis presents a systematic search for new mixed ionic-electronic conductors. It includes different crystalline structures and/or composition of the crystal lattice, varying the nature of the elements and the stoichiometry of the crystal. The research has yielded new materials capable to transport oxygen ions or protons and electronic carriers that are stable in the working condition to which they are submitted.
The characterization of a large series of lanthanide-doped ceria (CeO$_2$) provides an overall understanding of the structural and transport properties of these materials. Besides, the addition of cobalt oxide on this system is similarly studied. The analysis is completed by the optimization of the transport properties dependence on the microstructure. This allows a preliminary classification of the materials based on the main transport behavior and shows the possibility of tailoring both the structure and operating conditions to achieve desired properties for a number of applications, i.e., solid oxide fuel cells (SOFC) electrodes or oxygen separation membranes.

Several materials emerged from this work have achieved their purpose. Ce$_{1-x}$Tb$_x$O$_{2.6}$ and Ce$_{1-x}$Tb$_x$O$_{2.6}$ +2 mol% Co are proposed as a family of materials that may provide competent oxygen permeation fluxes and stability against CO$_2$ containing atmospheres. Besides, the inclusion of this material as a component of a dual-phase membrane increases the oxygen flux possibilities. The combination with a stable, Co and alkali-earth free spinel has delivered a promising membrane material for the integration in oxyfuel power plants.

On the other hand, metal dopants on the Ce$_{0.9-x}$M$_x$Gd$_{0.1}$O$_{2.6}$ structure have been added to lanthanide-doped ceria. They have been studied as SOFC cathode in combination with La$_{1-x}$Sr$_x$MnO$_3$-perovskite. The results show that the ionic conductivity invested by the ceria allows decreasing the polarization resistance regarding to the single phase cathode.

The study of a second group of materials involves a wide screening of doped CaTiO$_3$ perovskites. The difficulty of obtaining stable perovskites with high mixed ionic-electronic conductivity is evidenced. Iron and iron-magnesium doped CaTiO$_3$ are the best candidates as they present mainly ionic conductivity at high temperatures, while $p$-type electronic conductivity is predominant in the
low temperature range. CaTi$_{0.73}$Fe$_{0.18}$Mg$_{0.09}$O$_3$-$\delta$ is shown as a promising material offering good oxygen permeation fluxes and stability against CO$_2$, although it reacts with SO$_2$.

Finally, La$_{0.87}$Sr$_{0.13}$CrO$_3$ (LSC) perovskites have been doped in order to increase the mixed protonic-electronic conductivity. The study has lead to the development of a new generation of anodes for LWO based PC-SOFC. Ce in the A-site (LSCCe) and Ni in the B-site (LSCN) are stable perovskites in the reducing anode working conditions and in contact to the LWO electrolyte. Both of them decrease the polarization resistance regarding to pure LSC. LSCCe is limited by low frequency processes (LF), which are alleviated in the case of LSCN due to the formation of metallic nanoparticles on the surface. The further infiltration of Ni nanoparticles allows decreasing the LF resistance, which suggests that the surface reaction of H$_2$ oxidation is catalyzed. The most concentrated infiltration (5Ni) completely eliminates the LF resistance in both anodes and only high frequency (HF) processes remain. The best anode performance was obtained for LSCNi20+5Ni. Polarization resistance was as low as 0.26 $\Omega$·cm$^2$ at 750 °C in wet H$_2$. 
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“What makes the desert beautiful,' said the little prince, 'is that somewhere it hides a well...”

Antoine de Saint-Exupéry, The Little Prince