

# $\label{eq:chemical} \begin{array}{l} \mbox{Chemical and structural stability of} \\ \mbox{La0.6Sr0.4Co0.2Fe0.8O3}{-}\delta\ \mbox{ceramic vs. medium/high} \\ \mbox{water vapor pressure} \end{array}$

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## Chemical and structural stability of $La_{0.6}Sr_{0.4}Co_{0.2}Fe_{0.8}O_{3-\delta}$ ceramic vs. medium/high water vapor pressure

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#### Abstract:

During the last decades, perovskite-type oxides have received large attention as potential electrolytes and electrodes for Solid Oxide Fuel Cells (SOFC), including Proton Ceramic Fuel Cells (PCFC), gas separation membranes and High Temperature Steam Electrolysers (HTSE). A thermal treatment in autoclave, at a temperature close to an operating temperature, was used to measure the chemical stability of  $La_{0.6}Sr_{0.4}Co_{0.2}Fe_{0.8}O_{3-\delta}$ (LSCF6428) ceramic under medium and high water pressure (~20 & 40 bar). This mixed ionic-electronic conductor (MIEC) exhibits interesting properties as cathode of fuel cell materials. The reactivity rate of the investigated LSCF6428 sample under the protonation process conditions (several weeks at 550°C using CO<sub>2</sub>-free and CO<sub>2</sub>-saturated water) was studied in order to evaluate a potential use of this compound. Bulk and surface tructural/chemical changes were characterized by optical microscopy, TGA, dilatometry, Raman and ATR-FTIR spectroscopy. The results revealed only minor surface modifications in the case of ceramic treated under medium vapor pressure (20 bar) using CO<sub>2</sub>-free water. On the contrary, under higher pressure (40 bar) and CO<sub>2</sub>-saturated water several second phases were detected, namely strontianite, cobalt oxides and hematite. The chemical/structural stability of LSCF6428 is compared with previously investigated Rare-Earth nickelate ceramics:  $La_2NiO_{4+\delta}/Pr_2NiO_{4+\delta}/Nd_2NiO_{4+\delta}$ .

Keywords: B: Spectroscopy; Surface; C: Corrosion; Lifetime; D: perovskite; E: Fuel cell

#### 1. Introduction

Solid Oxide Fuel Cells (SOFC) received a large attention because of their high efficiency and low pollution emissions [1]. The oxygen ion conducting SOFCs based on zirconia/ceria electrolyte operate at high temperature, typically above 800°C. Such a high working temperature reduces the material's lifetime and increases significantly operating costs. To overcome these drawbacks, recent studies have paid attention to proton conducting electrode/electrolyte materials able to operate at lower temperatures (500-700°C) [2-5]. Such lower operating temperatures allow the use of less expensive steels and keeping of low-cost catalyzers. Reversibly, the electrochemical cells can be adapted to be used in High Temperature Steam Electrolyzer (HTSE). One great advantage of proton conductors for HTSE is the production of dry pure hydrogen in the cathode side. Electrode materials should exhibit important mixed ionic and electronic conductivity (MIEC) as well as show high efficiency and high stability vs. protons and/or oxygen/hydrogen. Furthermore, they should be also stable vs. other gas components such as CO<sub>2</sub>, NO<sub>x</sub> which could reduce the stability and the efficiency of the MIEC-materials as well as accelerate its ageing through chemical/structural transitions [6-8].

Different perovskite-related oxides exhibiting a large range of properties regarding to the variety of structure and chemical compositions have been tested as electrodes successfully [9]. Among various electrodes,  $La_{0.6}Sr_{0.4}Co_{0.2}Fe_{0.8}O_{3-\delta}$  (LSCF6428) exhibits excellent ionic-electronic conducting properties over a wide temperature range [10]: for example at 900°C the electronic conductivity can reach 230 S.cm<sup>-1</sup> whereas oxygen conductivity is 0.2 S.cm<sup>-1</sup> [11, 12]. Proton conductivity is expected but not yet

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demonstrated. Only few results are reported in the literature concerning electrochemical characterizations of similar MIEC-materials as a function of  $p(H_2O)$  [3, 13, 14]. At room temperature, LSCF shows almost cubic structure, with a small orthorhombic or rhombohedral distortion in the case of low Co and Sr contents. At high temperatures and/or high Sr content, a cubic phase is observed [15-17]. Structural stability of LSCF7382 was analyzed under H<sub>2</sub> atmosphere in a wide temperature range; reversible transition from the perovskite ABO<sub>3</sub> phase to the brownmillerite  $A_2B_2O_5$  was detected at 250-300°C. Moreover, decomposition of the structure into Co,  $La_2O_3$ , SrO and  $LaSrFeO_4$  was observed above 750°C [18, 19].

In our previous studies, we demonstrated the potential of high pressure/middle temperature autoclave to study the chemical stability of various perovskite-type electrolytes and electrodes vs. high pressurized water [20, 21]. Such treatment (550-600°C, 10-40 bar  $p(H_2O)$ ) that allows also incorporation of protonic species into a host material structure is called "Protonation". Note, the use of medium pressure allows also to optimize the economic efficiency of Fuel Cells and Electrolysers. Operating conditions (typically 10 to 20 bar  $p(H_2O)$ ) [IND] are the must. The use of high water pressure by industrialists allows an increased efficiency and in consequence decreasing of Hydrogen production/conversion cost [22-27]. Our previous studies, performed in a strict collaboration with Industrial partner, showed however that high water pressure can significantly accelerate a material ageing [membrane]. Consequently, our protonation procedure performed under very high pressure (40 bar  $p(H_2O)$ ) can be considered as an accelerated stability test which allows very fast to choose the most stable [22, 23].

The protonation process leads to subtle modifications of a material structure whatever the small amount of incorporated protons [28-35]. The quantity of  $CO_2$  dissolved in the water used is one of the stringent parameters. With high water pressure, the water activity at the ceramic surface increases, and enhances many reactions (hydroxylation, carbonation, or protonation). Consequently, a significant amount of secondary phases can be formed in relatively small experimental time (days to weeks). Careful analysis of the secondary phases with usual analytical techniques permits the study of the corrosion mechanism. Following the procedure of the previous studies, the LSCF6428 ceramic sample was treated under medium/high water vapor pressure ( $\sim 20$  and  $\sim 40$  bar) at 550°C for several days using a home-designed autoclave device [20, 28-32]. Two kinds of water were used: i) CO<sub>2</sub>-free water ii) CO<sub>2</sub>-saturated water. Thermogravimetry (TGA), thermal expansion (TE), X-ray diffraction (XRD), Attenuated Transmitted Reflected Fourier Transform-infrared spectroscopy (ATR FT-IR), Raman Spectroscopy (RS) were employed in order to identify corrosion films as well as eventual structural changes associated to the proton/water insertion. The aim of this study is the comprehension of the reaction mechanisms in operating proton conductor-based device conditions. Structural/chemical modifications allowing to estimate ageing of LSCF6428 ceramic were compared with other alternative cathode materials, namely Ln-nickelates [21].

#### 2. Materials and Methods

*Sample preparation: A* ceramic of La<sub>0.6</sub>Sr<sub>0.4</sub>Co<sub>0.2</sub>Fe<sub>0.8</sub>O<sub>3- $\delta$ </sub> (LSCF6428) composition, called hereafter 'as-prepared' sample, was prepared in pellet form ( $\emptyset$  = ~16 mm, thickness = 1.48 mm, theoretical density of 95 % max.). The preparation method was described by Mauvy and Batocchi [3]. Commercial powders provided by Marion Technology (Verniolle, France)

were isostatically pressed into pellets at 3000 bar for 5 minutes and then sintered in air at 1350°C for 4 hours. The thermal treatment of some as-prepared sample was carried out from 40°C to 1000°C using a TG balance (heating rate = 10 °C/min, 3 cycles) under He atmosphere.(99.99%, H<sub>2</sub>O < 3 ppm vol., O<sub>2</sub> < 2 ppm vol., C<sub>n</sub>H<sub>m</sub> < 0.5 ppm vol., Alphagaz-1, Air Liquide, France).

*Autoclave treatment:* The as-prepared ceramic was treated under high/medium water vapor pressure (20/40 bar, 550°C, CO<sub>2</sub>-free and CO<sub>2</sub>-saturated water) for period of 8 to 47 days using the home designed autoclave device [20, 28-32]. This device is equipped with a temperature and pressure controller, which allows us to follow the protonation process/incorporation of water/protonic species from the pressure drop. Note, Figure S1 (Supplementary Material) illustrates temperature and pressure curves recorded during the protonation process. The curves consist of two regions - start-up and protonation. The pressure drop in the protonation region was adjusted to 30 bar by the mean of a small leak valve. The further pressure decreases with a time is related to the absorption of water by the ceramic. The sample weight was systematically analyzed by non-destructive techniques after each run of autoclave treatment in order to observe the kinetic of protonation/reactions.

*Optical microscope analysis: Micro*-structures were analyzed on the surface as well as on a fresh fracture section of protonated samples using an Optical Microscope Olympus BX51 instrument. The 50x and 100x objectives were selected in order to observe microstructure modifications in the 5-1000  $\mu$ m range. Grains and pores sizes were measured using a line-intercept method (ASTM E112) with high magnification (10  $\mu$ m) of the micro-feature. Image contrast was optimized using the Microsoft Office Picture software.

*Thermal analysis:* TGA was performed between 40°C and 1000°C using a Setaram Setsys Evolution thermobalance. The use of Pt-crucible and He- atmosphere (99.99%, H<sub>2</sub>O < 3 ppm vol.,  $O_2 < 2$  ppm vol.,  $C_nH_m < 0.5$  ppm vol., Alphagaz-1, Air Liquide, France) allows the significant enhancement of the measurement accuracy. Thermal expansion (TE) measurements were carried out between 40°C and 600°C under Ar- atmosphere (99.99%, H<sub>2</sub>O < 0.5 ppm vol.,  $O_2 < 0.1$  ppm vol., CO < 0.1 ppm vol.,  $CO_2 < 0.1$  ppm vol., H<sub>2</sub> < 0.1 ppm vol., CO < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., CO<sub>2</sub> < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., Co < 0.1 ppm vol., H<sub>2</sub> < 0.1 ppm vol., Co < 0.1 ppm vol.,

*Vibrational spectroscopy analysis:* Raman spectra were recorded at room temperature under 458 nm and 514 nm wavelengths using a HR800 spectrometer (Horiba, John Yvon) equipped with a Peltier effect cooled CCD detector. The measurements under 785 nm wavelength were also performed using the solid state source line of a Senterra spectrometer (Bruker Optics, Karlsruhe, Germany). Raman spectra were recorded between 50 cm<sup>-1</sup> and 4000 cm<sup>-1</sup> through the 50x long working distance objectives (N.A. = 0.50). It was previously pointed out that moderate power of illumination may significantly heat the black ceramic samples [21, 36]. Consequently, in order to select the convenient laser power and to avoid phase transformation, preliminarily power tests were performed by varying the output power from 0.1 to 20 mW. Correct signal/noise ratios were obtained with relatively high counting duration of 100-500s. The spectra were fitted using Origin Software. The Raman mapping, furthermore, was performed on fresh fracture section of the protonated samples. The spectra were recorded from fresh surface deep to the sample's core with interval line of a few microns (5 – 50 µm).

ATR FT-IR spectra were recorded between 600 cm<sup>-1</sup> and 4000 cm<sup>-1</sup> at room temperature using the Bruker IR-scope FTIR spectrometer in ATR modes: a Diamond ATR (2x2 mm<sup>2</sup>) and a Germanium micro ATR ( $20x20 \ \mu m^2$ ) crystal.

*X-ray Diffraction:* XRD patterns of ground pellets were collected on a PANalitycal X'pert MPD-PRO Bragg-Brentano of  $\theta$ -2 $\theta$  geometry diffractometer equipped with a germanium primary monochromator over an angular range of  $2\theta = 10-130^{\circ}$ . Each acquisition lasted for 64 hours. The Cu-K $\alpha_1$  ( $\lambda = 1.54056$  Å) radiation was generated at 45 kV and 40 mA. Unit-cell parameters were calculated by Le Bail profile matching. The phase identification was carried out using the ICCD database and PANatycal X'pert highscore software. The relative quantities of the different phases were estimated according to the ratio of the relative intensities of the major peaks recorded in the XRD patterns.

#### 3. Results and Discussion

#### 3.1 Characterization of the pristine LSCF ceramic

#### 3.1.1 Mean symmetry

The XRD results presented in Fig.1 show that LSCF6428 exhibits a single rhombohedral phase with *R*-3c space group. The pristine symmetry of the material investigated here is then similar to previous literature studies [3, 37, 38]. Structural modifications involved by the autoclave treatment will be discussed in section 3.3.

#### 3.1.2 Raman scattering

Because of the black color of the as-prepared LSCF6428 sample that leads to high conversion of the laser beam to heat at the examined spot, Raman spectra shown in Fig. 2a

were recorded as a function of laser power (0.1 - 20 mW, 458 nm, 135 – 1250 cm<sup>-1</sup>). At the lowest power (~0.15 mW), the spectrum reveals the presence of typical LSCF perovskite phase consisting of a main peak at 662 cm<sup>-1</sup> and three broad bands at 569, 470, and 292 cm<sup>-1</sup>. A small broad band at 180 cm<sup>-1</sup> is also detected. The first two bands at 662 and 569 cm<sup>-1</sup> are similar to those observed in the literature [18, 39-42]. It should be noted that according to its complex composition, LSCF6428 can be considered as a solid solution between La<sub>1-x</sub>Sr<sub>x</sub>CoO<sub>3-δ</sub> (LSC) and La<sub>1-x</sub>Sr<sub>x</sub>FeO<sub>3-δ</sub> (LSF). The large bandwidth is related to the Co/Fe and La/Sr partial substitutions. The group theory analysis in the case of rhombohedral symmetry with the *R*-3c space groups predicts 18 optical phonons of irreducible representations [40]. Fives modes ( $A_{1g}+4E_g$ ) are Raman actives, and eight modes ( $3A_{2u}+5E_u$ ) are IR actives, while the remaining modes ( $2A_{1u}+3A_{2g}$ ) are inactive. According to the literature [39], the Raman bands at 650 and 590 cm<sup>-1</sup>, typical of the LSCF material, correspond to  $E_g$  modes.

With increase of the laser power, the temperature at the sample surface increases, which can lead to oxidation reaction. Simultaneously a wavenumber downshift is expected and observed under 2.5 mW laser beam. According to the study of Siebert et al. [18], the Raman spectrum of fully oxidized LSCF7382 exhibits an overlapping of two broad bands centered at 560 and 630 cm<sup>-1</sup>. Our spectra showing bands at 569 and 662 cm<sup>-1</sup>, Fig. 2a, are consistent with a low oxygen deficiency. Overall area of spectrum and relative intensity between the band at 569 and 662 cm<sup>-1</sup>, plotted as a function of the output laser power, are shown in Fig. 2b. The result shows a decrease of the sum of peak area with the increase of the laser power. This vanishing of Raman signature is consistent with the progressive transition toward the cubic structure [20, 43]. Moreover, a jump of the relative intensity

(I<sub>569</sub>/I<sub>662</sub>) at 2.5 mW of laser power was clearly observed and indicates a phase transition due to local heating under the laser beam. Consequently, only the Raman spectra recorded below this limit power of 2.5 mW, can be considered as showing the "true" signature of the pristine LSCF6428 ceramic at room temperature, as analyzed by XRD (Fig. 1).

Raman spectra, Fig. 3, were recorded under different laser lines: 458 nm, 514 nm and 785 nm with 50x objectives in several surface spots. The 458 nm spectrum under 0.15 mW laser power shows the signatures of a pure LSCF perovskite phase as discussed earlier. A rather similar spectrum is obtained under 3 mW with 785 nm exciting line indicating a lower light absorption - and heating - in near IR range. Additional narrow Raman peaks at about 685, 518, and 480 cm<sup>-1</sup> were observed. These peaks can be assigned to the contribution of Co<sub>3</sub>O<sub>4</sub> second phase. A reference spectrum given in Fig. 3 has been obtained by heating of CoO.OH heterogenite [44]. The small wavenumber shift and relative intensity change arise from the (pre)resonance Raman effect [45]. No "well-defined" spectrum of CoO phase seems to be available in the literature. Very low intense spectrum with broad bands at 465 and 675 cm<sup>-1</sup> is given by Tang et al. [46]. According to Tang et al.' assignment, the CoO contribution in the Raman spectrum is detected by the broadening of the 461 cm<sup>-1</sup> peak. Alternatively, the Raman spectrum of CoO.OH (Fig. 3) shows a narrow peak at 498 cm<sup>-</sup> <sup>1</sup> and a broad feature at  $\sim$ 620 cm<sup>-1</sup> that may explain the rather high intensity of the  $\sim$ 480 cm<sup>-1</sup> peak. Anyway, the intense light absorption of the black material promotes the transformation of (hydroxylated) CoO into  $Co_3O_4$  under the laser beam. The observation of a band at  $\sim$ 1140 cm<sup>-1</sup> is also consistent with hydroxylated CoO [44]. Traces of metal transition oxides are commonly observed in similar compounds [46].

#### 3.2 Protonation treatment under medium/high pressure

#### 3.2.1 Reactivity vs. water

The investigated LSCF6428 ceramic was treated, similarly to our previously investigated nickelate-based samples [21, 36], using an autoclave device under ~20 bar of CO<sub>2</sub>-free water pressure at 550°C. The mass variation between pristine and protonated ceramics was recorded versus time. Consequently, the mass difference per sample surface area (mg/mm<sup>2</sup>) exposed to water pressure was calculated using equation 1. The graph of the calculated mass change vs. protonation time is plotted in semi-logarithmic scale as shown in Fig. 4.

Mass uptake 
$$(mg / mm^2) = \frac{m_i - m_0}{S}$$
, i = 1, 2, 3, ... ... (1)

where  $m_0$  is the as-prepared sample weight (mg),  $m_i$  is the sample weight of i = 1, 2, 3...protonation runs up to the complete studied time range and *S* is sample surface area,  $2\pi r^2 + h(2\pi r)$ .

The exponential fitting curves, Fig. 4, reveal very low corrosion of the LSCF6428 ceramic in comparison with our previously investigated nickelates [21, 36]. The reactivity rate of the LSCF6428 sample is about 1:42, 1:22 and 1:8 of the reactivity rate in the case of LNO, PNO and NNO sample, respectively. Hence, the LSCF6428 ceramic exhibits the highest stability under 20 bar of CO<sub>2</sub>-free water pressure at 550°C.

#### 3.2.2 Microstructures

The samples were categorized in different states; as-prepared/pristine (a), as-prepared sample thermally treated at 1000°C (a'), protonated (b, c, d), and deprotonated by thermal

treatment at 1000°C (b', c', d'). Figure 5 illustrates the surface microstructure features of each sample states. The surface of as-prepared sample (Fig. 5a) as well as that of the thermally treated sample (Fig. 5a') show grains of ~1-2  $\mu$ m with pores of ~2 to 10  $\mu$ m. A white contour is observed at the grain periphery and assigned to 2<sup>nd</sup> phase traces already identified by XRD and Raman analysis. In the case of the ceramic protonated under 40 bar of CO<sub>2</sub>-free water for a short period of 8 days (Fig. 5b), the observed surface habits are similar to those detected for the as-prepared sample. However, we observe an increase of the number of grains affected by the white contour. This may indicate an increase of the secondary phase content. In the case of thermally treated protonated LSCF6428 using CO<sub>2</sub>-free water (Fig. 5b' and 5c'), the surface habit remains rather similar to that of the as-prepared samples, in agreement with good stability of the ceramic.

In the case of the ceramic protonated under  $CO_2$ -saturated water, the surface features were more affected - small aggregates (~4 µm) were covered by another phase. After deprotonation performed at 1000°C (Fig. 5c'), the microstructure remains rather stable. By comparison with previously analysed protonated nickelate homologues [21, 36], no crack was observed neither for the protonated nor the deprotonated LSCF6428 ceramics, treated either under  $CO_2$ -free or  $CO_2$  saturated water.

#### 3.3 Chemical and structural modifications

#### 3.3.1 Thermogravimetry and thermal expansion

Fig. 6a compares the TGA curves of non-protonated and protonated LSCF6428 ceramic. According to the oxygen mobility [11, 15, 16], the weight loss observed above 600°C (He atmosphere) in the case of non-protonated pellets can be related to the equilibration of the oxygen stoichiometry with the new environment, different from the sintering atmosphere [47]. The variation is very weak according previous measurement of the oxygen stoichiometry [2]. Furthermore the low delta value limits the number of vacancies. In order to highlight the contribution solely due to the autoclave treatment, the oxygen stoichiometry variation was suppressed by subtracting the measure made on the non-protonated, as-prepared sample: the relative mass-loss difference between non-protonated and protonated samples is plotted in Fig. 6b. The difference curve clearly shows the presence of one weak peak at ~400°C in the case of medium-pressure (~20 bar) protonated samples. For the high-pressure (40 bar) protonation, a second stronger peak is observed at ~800°C. According to previous studies [20, 28, 29], the phenomenon at 300-400°C can be assigned to the decomposition of hydroxide(s), and that at ~800-900°C to the carbonate decomposition.

According to the above TG data and hypothesis that the ceramic exhibits a single phase and that the initial  $\delta$  oxygen stoichiometry is not modified at 550°C in the autoclave, the following formula can be proposed for protonated LSCF6428 ceramic under 40 bar of CO<sub>2</sub>absorped water pressure:  $La_{0.4}Sr_{0.6}Fe_{0.2}Co_{0.8}O_{3-\delta}(OH)_{0.01}.(CO_3)_{0.04}$ . In the case of ceramics treated under free-CO<sub>2</sub> water pressure in both medium and high pressure, the content of hydroxides moieties was much lower (< 0.001 mole/mole pH<sub>2</sub>O).

Comparison of thermal expansion (TE) curves characteristic of non-protonated and protonated LSCF6428 samples is presented in Fig. 6c. The area located in-between heating and cooling curves of pristine, non-protonated sample is rather small, revealing stability of the material (measured temperature range is far below the sintering temperature of the ceramic pellet). Only the first heating TE curve of protonated pellet (~40 bar of CO<sub>2</sub>-free vapor pressure) shows clearly a contraction jump at ~350°C. A similar phenomenon was observed in our previous works devoted to stability studies of nickelate-based oxides [21, 36]. Furthermore, a study on phase transition of LSCF7382 under H<sub>2</sub> atmosphere by Siebert et al. [18], points out the transformation from rhombohedral perovskite structure to orthorhombic  $A_2B_2O_5$  brownmillerite phase at 350°C. Formation of a hydroxide film at the ceramic surface could also explain the presence of such a jump since dehydration of hydroxides is observed in this temperature range. The lack of cracks in the ceramic pellet is more consistent with the assignment of this jump modification to the presence of  $2^{nd}$  phase surface film than to the effect of the phase transition involving a contraction of the unit-cell in the bulk.

#### 3.3.2 Unit-cell symmetry

Figures 1a and 1b compare XRD patterns of as-prepared and 20 bar protonated/ 40 bar protonated using  $CO_2$ -free water LSCF6428, respectively. Extra Bragg peaks observed after protonation treatment confirm the presence of cobalt oxide(s), namely CoO. According to the refinement of XRD patterns and relative ratio analysis, the amount of cobalt oxide(s) traces can be estimated to 1-2% depending on the protonation pressure - the higher pressure value, the higher percentage of CoO. The detection of  $Co_3O_4$  by Raman spectroscopy could be due to the presence of both CoO and  $Co_3O_4$ . Note, the higher Raman intensity of  $Co_3O_4$  hinders the observation of CoO signature and/or the oxidation of CoO can appear under the laser beam heating. Taking into account the unit-cell parameters and space group given in Table 1, it is clear that the protonated LSCF6428 keeps the rhombohedral symmetry, no matter the protonation pressure value is. The protonation treatment under medium and high pressure causes a small expansion of unit-cell volume: 0.57% for 20 bar protonation and 1.10% for 40 bar protonation. This volume expansion could be consistent with the contraction observed in Fig. 6c, because of the departure of protons inserted in the perovskite structure.

	phase		Unit-cell parameters				
Sample	symmetry of perovskite structure	Space group	a (Å)	b (Å)	<b>c</b> (Å)	<b>V</b> (Å <sup>3</sup> )	ΔV (%)
LSCF6428	rhombohedral	<i>R</i> -3c	5.4878(5)	5.4878(5)	13.374(4)	348.8(1)	_
$LSCF6428.H^+$	rhombohedral	<i>R</i> -3c	5.4936(8)	5.4936(8)	13.423(8)	350.8(2)	+ 0.57
$(20 bar CO_2$ -free water)	+ CoO						
LSCF6428.H+	rhombohedral	<i>R</i> -3c	5.5107(1)	5.5107(1)	13.4088(3)	352.64(1)	+ 1.10
(40 bar CO <sub>2</sub> -free water)	+ CoO						

 Table 1: XRD analyses of as-prepared and protonated LSCF6428 samples.

#### 3.3.3 Vibrational spectroscopy analysis

Raman spectra recorded at the surface of protonated LSCF6428 samples, Fig. 7a, clearly reveal the presence of secondary phases i.e. oxide(s), hydroxide(s) and carbonate(s) depending on protonated/thermal treatment conditions.

i) In the case of protonation performed under 20 bar using CO<sub>2</sub> free- water (20 bar, *P20*), Raman signatures at 688, 611, 493, 407, 289, and 221 cm<sup>-1</sup> were observed. These Raman peaks are assigned to hematite,  $\alpha$ –Fe<sub>2</sub>O<sub>3</sub> [48].

ii) For higher pressure of  $CO_2$  free-water (40 bar, P40), the Raman spectrum shows traces of lanthanum oxycarbonate (La<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>). Our reference spectra of LaOOH.(CO<sub>3</sub>) and

La(OH)<sub>3</sub> obtained by *in*-situ thermal treatment of hydroscopic La<sub>2</sub>O<sub>3</sub> powder are given in Fig. S2 (Supplementary Material). The Raman signature detected at 1066 cm<sup>-1</sup> is assigned to internal stretching mode (A<sub>1g</sub>) of C-O bonds [49].

iii) For the ceramic treated under 40 bar of  $CO_2$ -saturated water pressure (40 bar, *P40/CO<sub>2</sub>*), Raman spectra recorded at the surface show carbonate signature: the peaks at 1073, 702, 247, 182, and 150 cm<sup>-1</sup> were assigned to strontium carbonate, SrCO<sub>3</sub>, with its characteristic 1073 cm<sup>-1</sup> symmetric stretching mode ( $v_1$ ) of C-O bond [50]. A peak at ~865 cm<sup>-1</sup> and a weak band at ~567 cm<sup>-1</sup> can be assigned to a strontium oxide/peroxide. This result is similar to that presented in the study of Waal [51], where a peak at 863 cm<sup>-1</sup> is attributed to the O-O stretching mode of SrO<sub>2</sub> composition, and three very weak bands at 563, 537, 522 cm<sup>-1</sup> are assigned to translation modes of Sr-O vibrational unit. The presence of Sr-based oxides could be related to the thermal decomposition of the Sr-carbonate under the laser spot because of a black matrice.

 Table 2: Raman wavenumbers of the different second phases found on the bulk

 and surface of protonated LSCF6428 ceramics.

Phase	Symbol*	wavenumber (cm <sup>-1</sup> ) <sup>+</sup>	References
SrO	ф	<u>560</u> , 540, 520	[51]
SrO <sub>2</sub>	Δ	<u>863</u>	[31]
$Co_3O_4$	σ	<u>686, 614, 520, 479, 191</u>	[44, 52-54]
γ-Fe <sub>2</sub> O <sub>3</sub>	γ	<u>700, 600, 495, 406, 292, 247, 226</u>	[48]
I aO OH		424, <u>383</u> , <u>345</u> , 312, 216, 202, 165, 135, 123,	[49]
Lao.on	μ	116	[עד]
SrCO <sub>3</sub>	*	1446, <u>1071</u> , <u>710</u> , 511, 258, <u>245</u> , 237, <u>180</u> ,	[50, 55]

	<u>148,</u> 100				
I-La <sub>2</sub> O <sub>2</sub> CO <sub>3</sub>	**	<u>1064, 868, 721, 670, 315</u>	[49]		

<sup>\*:</sup> symbols used in Fig. 7

+: underline Raman wavenumbers are observed in this study

Figure 7b shows representative Raman spectra recorded in the bulk of LSFC6428 ceramic treated under different protonation conditions. Measurements are made on the fresh fracture of the ceramic pellet, from surface to the bulk centre.

- i) Raman spectra characteristic of the sample treated under 20 bar/CO<sub>2</sub>-free (at 510  $\mu$ m bulk) and 40 bar/CO<sub>2</sub>-saturateded water (at 50  $\mu$ m bulk), reveal the presence of traces of Cobalt spinel (Co<sub>3</sub>O<sub>4</sub>: 689, 618, 523, 485, and 195 cm<sup>-1</sup>). However, some slight peak downshifts are detected between pure Co<sub>3</sub>O<sub>4</sub> and that present in protonated LSCF6428.
- ii) Raman spectrum recorded on bulk, 50  $\mu$ m depth (protonated under 20bar, Fig. 7b), shows the bands at 606, 501, 404, 288 and 221 cm<sup>-1</sup> which are characteristic of hematite ( $\alpha$ -Fe<sub>2</sub>O<sub>3</sub>) [56].

ATR-FTIR technique specially probes the surface, typically a few (tenths) micron indepth penetration: the exact penetration depth varies with the wavenumber depending on the optical index of the crystal probe (diamond or germanium, Fig. 8). Depending on the band intensity and optical characteristic of the investigated solid (the higher optical index , the higher reflection), the band shape is complex and can be described as a mixture of absorption and reflection signal [57, 58]. Thus, it is not possible to discus the crystallinity of the second phases. The ATR-FTIR tests were performed on the as-prepared (a, b), protonated in different conditions (c, d, and e) and thermally treated as-prepared (f) samples. A sharp peak observed at ~3604 cm<sup>-1</sup> can be assigned to the stretching mode of OH<sup>-</sup> in Sr(OH)<sub>2</sub>. This contribution confirms the presence of protonic species absorbed on the ceramic surface. The absorption bands at 600-1500 cm<sup>-1</sup> detected in the case of protonated samples correspond to traces of SrCO<sub>3</sub> [30]. The strong absorption peak centered at 1422 cm<sup>-1</sup> and the band at 701 cm<sup>-1</sup> are assigned to the asymmetric stretching ( $\upsilon_3$ ) and bending ( $\upsilon_4$ ) vibrations of C-O bonds, respectively. A strong narrow peak at ~859 cm<sup>-1</sup> can be assigned to the out-of-the-plane bending ( $\upsilon_2$ ) mode, and that observed at ~1071 cm<sup>-1</sup> to the fully symmetric stretching ( $\upsilon_1$ ) mode of  $CO_3^{2^-}$  ions. The absorption bands characteristic of Sr(OH)<sub>2</sub>, and SrCO<sub>3</sub> observed here are similar to those presented in the literature [59, 60].

#### 3.4 LSCF6428 chemical stability

A Raman mapping was performed on the cross-section of a fresh fractured sample – from the top surface (directly exposed to the vapor pressure) down to the bottom one (in contact with the autoclave chamber) in step of few micron to 50 µm. This procedure is the so-called Raman profilometry method discussed in details in our previous studies [20, 21, 35, 36]. Figure 9 compares the Raman mapping spectra recorded for ceramics protonated using three distinct conditions: 20 bar of free-CO<sub>2</sub> water pressure (47 days), 40 bar of free-CO<sub>2</sub> water pressure (8 days) and 40 bar of CO<sub>2</sub>-saturated water pressure (8 days). The relative volume section of phase modification is summarized in Table 3.

#### Table 3: comparison of corrosion in LSCF6428 dense ceramic.

Section (%)

Protonation conditions

	20 bar/47days free-	40 bar/8days	40 bar/8days	
	CO <sub>2</sub> water	free-CO2 water	saturated-CO <sub>2</sub> water	
Top layer	<0.1	~0.4	~0.7	
core	99.9	99.2	98.9	
Bottom layer	<0.1	~0.4	~0.4	

Due to the possibility of focusing at the microscale on a sample surface, the Raman technique has higher ability than the usual XRD technique to detect even small structural changes as well as secondary phases. However, the combination of the black color of the matrix/substrate and the laser spot can lead to uncontrolled heating and then structural/chemical transformations (partial dehydration/oxidation) under the laser light as discussed in section 3.12. The dehydration of those hydrated/hydroxylated substrate derivatives may induce phase transitions/transformations and hence a higher number of phases that explains the higher number of phases detected by vibrational spectroscopy. Furthermore, the vibrational spectroscopy techniques are able to reveal the presence of amorphous oxyhydroxides, not detectable by XRD [20, 30]. Since the Raman intensity depends on the chemical bond polarisability, minor phases may have a stronger signature than the major matrix.

No transformation on the top layer of protonated LSCF6428 is observed after protonation under 20 bar. Traces of hematite ( $\alpha$ -Fe<sub>2</sub>O<sub>3</sub>) and cobalt oxide are observed in some spots in the bulk due to defects formed in the powder processing and/or sintering. On the contrary, the Raman spectra characteristic of the ceramics protonated under 40 bar of both CO<sub>2</sub>-free and -saturated water, show the presence of carbonate compounds (nearly surface, <1  $\mu$ m) as well as other hydroxide(s)/oxide(s) phases. The carbonate phases are identified as strontianite (SrCO<sub>3</sub>) and lanthanum oxycarbonate (I-La<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>). Hydroxide(s)/oxide(s) phase are identified as lanthanide oxyhydroxide (LaO.OH) and cobalt oxides (Co<sub>3</sub>O<sub>4</sub>). Note, comparison of 2<sup>nd</sup> phase volumes between medium and high vapor pressure protonation treatments indicates that they are not significantly different.

#### **4** Conclusions

According to the protonation treatment (~20 and ~40 bar of CO<sub>2</sub>-free or CO<sub>2</sub>-saturated water, 550°C) and several advanced analysis techniques, we can summarize that the LSCF6428 ceramic exhibits very good stability under medium and high water pressure, 5 to 30 times better than nickelates homologues (Table S1). XRD results show that the structure of protonated pellets remains rhombohedral (*R*-3c) with a small unit-cell expansion related to proton incorporation. Only traces of secondary phases were detected in the ceramic bulk. XRD reveals the formation of CoO. Fe<sub>2</sub>O<sub>3</sub> traces are also observed within the bulk as processing defects. Raman spectroscopy shows the presence of oxyhydroxide(s). According to TG results, the content of OH<sup>-</sup> (< 0.01 mole/mole.H<sub>2</sub>O) is very low. Raman mapping allows an identification of surface corrosion products: the hematite ( $\alpha$ -Fe<sub>2</sub>O<sub>3</sub>), cobalt oxides (CoO/Co<sub>3</sub>O<sub>4</sub>), tetragonal lanthanum oxycarbonate (I-La<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>), and strontianite (SrCO<sub>3</sub>). The type and content of these secondary phases depend on the protonation conditions (i.e. CO<sub>2</sub> contents, pressure, duration times). As expected, the highest content of carbonates is found after the treatment using CO<sub>2</sub>-saturated water.

Fig. 10 compares the rate of the formation of the corrosion film thickness formed at the surface of 'dense' ceramic pellets used as electrolytes (BCZ:YZ,  $BaCe_{0.5}Zr_{0.3}Y_{0.16}Zn_{0.04}O_{3-\delta}$ 

*BZ:Yb,*  $BaZr_{0.9}Yb_{0.1}O_{3-\delta}$ ; *SZ:Yb,*  $SrZr_{0.9}Yb_{0.1}O_{3-\delta}$ ) [20] and electrodes ( $Ln_2NiO_{4+\delta}Ln = La$ , LaNO, Ln = Pr, PrNO and Ln = Nd, NdNO; LSCF6428) [21]. Under medium water pressure the stability of electrode materials and of strontium zirconate electrolyte is good. Under high water pressure barium-base perovskite electrolytes are no more stable and if the water has been saturated with  $CO_2$  the superior stability of LSCF compound is demonstrated whatever the lower carbonation of Pr and Nd-based compounds vs. La, Ba or even Sr homologues.

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#### **Figure Captions**

- **Figure 1:** Comparison of X-ray patterns a) as-prepared vs. protonated LSCF6428 under 20 bar CO<sub>2</sub>-free water b) as-prepared vs. protonated LSCF6428 under 40 bar CO<sub>2</sub>-free water.
- **Figure 2:** a) Raman spectra (458 nm) of as-prepared LSCF6428 ceramic as a function of laser power b) Oxidation relative intensity peak (I<sub>569</sub>/I<sub>662</sub>) versus laser power.
- Figure 3: Raman signatures of as-prepared LSCF6428 ceramic at different surface area/laser power/wavelengths and of Co<sub>3</sub>O<sub>4</sub> obtained by thermal treatment of CoO.OH [44]. (\* corresponds to Co<sub>3</sub>O<sub>4</sub> traces).
- **Figure 4:** mass uptake evolution of LSCF6428 ceramic pellet during 47 days of protonation treatment (20 bar of CO<sub>2</sub>-free water and 550°C).
- Figure 5: Micro features (10 μm) of LSCF6428's surface sample in different states (a) asprepare sample (b) 40 bar of CO<sub>2</sub>-free water pressure for 8 days (c) 20 bar of CO<sub>2</sub>free water pressure for 47 days (d) 40 bar of CO<sub>2</sub>-saturated water pressure for 8 days.
- Figure 6: a) Thermal analysis of as-prepared and protonated LSCF6428 b) corresponding TG weight loss difference, %wt<sub>p</sub> - %wt<sub>np</sub> c) Two consecutive thermal expansion (length variation/initial length) curves of as-prepared and protonated LSCF6428 under free-CO<sub>2</sub> water pressure.
- **Figure 7:** Raman spectra of LSCF6428 ceramic in different states, protonated and deprotonated, recorded a) on surface b) in bulk. (Note: the used symbols are described in Table 2 and sample labels are indicated in the Text)

- **Figure 8:** ATR and micro ATR-FTIR spectra recorded at the surface of as-prepared (a, b), protonated under 20 bar free-CO<sub>2</sub> (c), under 40 bar free-CO<sub>2</sub> (d), 40 bar absorbed-CO<sub>2</sub> (e) and thermal treatment (f). (Note: \* indicates the spectra collected with the μ-ATR Ge crystal)
- Figure 9: Raman mapping along the fresh cross-section fracture of LSCF6428 ceramic pellets protonated under a) 20 bar of CO<sub>2</sub>-free water b) 40 bar of CO<sub>2</sub>-free water c) 40 bar of CO<sub>2</sub> saturated water.
- Figure 10: Comparison of corrosion rate ( $\mu$ m/day) of various Electrolytes [20] and Electrodes treated in Autoclave under different conditions (see text for label).



Figure 1: Comparison of X-ray patterns a) as-prepared vs. protonated LSCF6428 under 20 bar

*CO*<sub>2</sub>-free water *b*) as-prepared vs. protonated *LSCF*6428 under 40 bar *CO*<sub>2</sub>-free water.



**Figure 2:** a) Raman spectra (458 nm) of as-prepared LSCF6428 ceramic as a function of laser power b) Oxidation relative intensity peak ( $I_{569}/I_{662}$ ) versus laser power.



**Figure 3:** Raman signatures of as-prepared LSCF6428 ceramic at different surface area/laser power/wavelengths and of  $Co_3O_4$  obtained by thermal treatment of CoO.OH [44]. (\* corresponds to  $Co_3O_4$  traces).



*Figure 4:* mass uptake evolution of LSCF6428 ceramic pellet during 47 days of protonation treatment (20 bar of  $CO_2$ -free water and 550°C).



**Figure 5:** Micro features (10  $\mu$ m) of LSCF6428's surface sample in different states (a) asprepare sample (b) 40 bar of CO<sub>2</sub>-free water pressure for 8 days (c) 20 bar of CO<sub>2</sub>-free water pressure for 47 days (d) 40 bar of CO<sub>2</sub>-saturated water pressure for 8 days.



weight loss difference,  $\%wt_p$  -  $\%wt_{np}$  c) Two consecutive thermal expansion (length variation/initial length) curves of as-prepared and protonated LSCF6428 under free-CO<sub>2</sub> water pressure.



**Figure 7:** Raman spectra of LSCF6428 ceramic in different states, protonated and deprotonated, recorded a) on surface b) in bulk. (Note: the used symbols are described in Table 2 and sample labels are indicated in the Text)



**Figure 8:** ATR and micro ATR-FTIR spectra recorded at the surface of as-prepared (a, b), protonated under 20 bar free- $CO_2$  (c), under 40 bar free- $CO_2$  (d), 40 bar absorbed- $CO_2$  (e) and thermal treatment (f). (Note: \* indicates the spectra collected with the  $\mu$ -ATR Ge crystal)



**Figure 9:** Raman mapping along the fresh cross-section fracture of LSCF6428 ceramic pellets protonated under a) 20 bar of  $CO_2$ -free water b) 40 bar of  $CO_2$ -free water c) 40 bar of  $CO_2$  saturated water.



Figure 10: Comparison of corrosion rate ( $\mu$ m/day) of various Electrolytes [20] and Electrodes treated in Autoclave under different conditions (see text for label).

# Chemical and structural stability of $La_{0.6}Sr_{0.4}Co_{0.2}Fe_{0.8}O_{3\cdot\delta}$ ceramic vs. medium/high water vapor pressure

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Figure S1 : Example of a plot presenting pressure drop and temperature evolution v.s. protonation time.



Figure S2: Raman spectra of La(OH)<sub>3</sub>, LaOOH.(CO<sub>3</sub>) and A-La<sub>2</sub>O<sub>3</sub> phase obtained by in-situ thermal treatment of hydroscopic La<sub>2</sub>O<sub>3</sub> powder sample, and a Raman spectrum of II-La<sub>2</sub>O<sub>2</sub>CO<sub>3</sub> power sample at room temperature.

Sample Types	pristine state 🗕		Corrosion rate		
		time	T/°C	P / bar	(µm/day)
		Ele	ctrolyte group [19	)]	
BaCe <sub>0.5</sub> Zr <sub>0.3</sub> Y <sub>0.16</sub> Zn <sub>0.04</sub> O <sub>3</sub> - δ	high density (97%), ceramic pellet, middle brown	30 h	500	10	~30
BaZr <sub>0.25</sub> In <sub>0.75</sub> O <sub>3-δ</sub>	dense (90%), ceramic pellet, light brown	5 d	300	80	>150
$BaZr_{0.9}Yb_{0.1}O_{3-\delta}$	high density (97%), ceramic pellet, middle	23 d	200	15	~10
SrZr <sub>0.9</sub> Yb <sub>0.1</sub> O <sub>3-δ</sub>	high density (97%),	5-23 d	200	15	<0,25
	ceramic pellet, middle	- 1	or	0.0	
	brown	5 d	500	80	
		Ele	ectrode group [20	]	
La2NiO4+8	High density (95%), ceramic pellet, black	47 d	550	20	<0.1
		8	550	40	0.1
		8	550	40 (CO2-sturated)	~5
Pr <sub>2</sub> NiO <sub>4+δ</sub>	High density (95%), ceramic pellet, black	47 d	550	20	<0.1
		8	550	40	~0.1
		8	550	40 (CO <sub>2</sub> -sturated)	~1.25
Nd2NiO4+8	High density (95%), ceramic pellet, black	47 d	550	20	<0.1
		8	550	40	~0.1
		8	550	40 (CO2-sturated)	~0.6
La0.6Sr0.4Co0.2Fe0.8O3-8	High density (95%), ceramic pellet, black	47 d	550	20	< 0.01
		8	550	40	~0.1
		8	550	40 (CO2-sturated)	~0.1

## Table S1: Summary of Autoclave Treatment of Electrolyte and Electrode Samples