The roles of constituting oxides in rare-earth cobaltite-based perovskites on their pseudocapacitive behavior

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ABSTRACT

The role and influence of strontium and its oxide on structure and capacitive response of materials containing mixed lanthanum cobalt oxides, LC, and lanthanum strontium cobalt oxides, LSC, as a capacitive materials were investigated in this study. The mixed oxides were synthesized by the single-step ultrasonic spray pyrolysis (USP) technique. The microstructures and electrochemical properties of the samples were characterized by X-ray diffraction, scanning electron microscopy, cyclic voltammetry, potentiostatic electrochemical impedance spectroscopy and galvanostatic charge/discharge cycling. It was found that strontium oxide induces the formation of the perovskite structure of promoted pseudocapacitive behavior over an enhancement of redox transitions of cobalt. The measurements showed that the capacitive stability and rate capability were lower for the samples of higher specific capacitance. Among the prepared materials, the LSC prepared at a USP temperature 600 °C showed the best capacitive characteristics in 0.10 M KOH due to having the most defined spherical perovskite structure leading to well-defined reversible charge-discharge performances.

1. Introduction

In order to reach highly reversible and stable materials for energy storage processes, investigation of novel materials, as suitable supports for noble ones, has been predominant in the past few years [1-9].

Perovskite oxides are promising candidates for electrochemical power sources due to their unique physical and electronic properties, of which its structural stability is one of the most important. Perovskites can accommodate mobile oxygen ions under electrical polarization. It was shown that oxygen ion mobility and vacancies play an important role in charge storage and in catalysis in general [10]. The composition and structural stability of Perovskite not only allow storage of large quantities of energy, but also ensure high rates of delivery [11].

Alkaline-stable perovskite materials with the general formula ABO3 (where A is a lanthanide and B is a transition metal) were proven to have metallic conductivity and some redox capacitive behavior [10-17], which makes them promising candidates as supports in supercapacitive materials support of reversible and stable response. Oxygen-vacancy mediated redox pseudocapacitance for a nanostructured lanthanum-based perovskite, LaMnO3 was demonstrated for the first time for fast energy storage [10]. Rare-earths in ruthenates are also known for improvement of the specific capacitance, especially La replacement in SrRuO3 [12]. ABO3 materials with high electrochemical stability and fast charge/discharge rates can be synthesized in different material combinations [6,7,15,18-20]. It was found recently that simple impregnation of LaSrCoO3 with RuO2 leads to considerable mutual enhancement of the capacitive behavior of these two structural components [6]. Similarly, the synergy of Co/Mo has good redox ability and it facilitates high oxygen mobility, fast kinetics of charge storage and excellent cycle life [21].

As a synthesis procedure, a unique ultrasonic spray pyrolysis. (USP), was applied that appeared to be a highly promising technique for the synthesis of perovskite materials [6,22]. It was found that certain USP-synthesized perovskites as solid electrolytes have the ability to improve the performance of solid oxide fuel cells (SOFC) [22]. Perovskite materials can have long cyclability compared to carbon-supported composites in alkaline to neutral media [3,23]. Lanthanum...
cobalites (LC) and strontium-impregnated lanthanum cobalites (La$_{1-x}$Sr$_x$CoO$_{3-δ}$, LSC) are supposed to have wider voltage window than carbonaceous materials [4,6].

Although the perovskites have been proven to be of pronounced capacitance with respect to the constituting oxides, the roles and influences of the separate components in the capacitive responses of enhancing perovskite structures are not clear. Zhang et al. [23] performed ex situ doping of LC with Sr oxide and found some improvements in the electrode capacitive response and also nickel foam supported asymmetric capacitor. However, the unique redox influence of strontium oxide doping into the structure of LC was not reported.

The aim of the reported investigation was to bring detailed insights concerning separated influences of constituting oxides to this issue, in order to reveal the redox electrochemistry behind perovskite structures as supports for supercapacitive applications. The synthesis approach involved in situ impregnation of the LC structure with Sr oxide.

LC, as the “Sr-free” counterpart of strontium-doped lanthanum cobaltite (LSC) nanopowders were also synthesized by the ultrasonic spray pyrolysis process and their structural and capacitive properties were analyzed and compared.

2. Experimental

2.1. Synthesis of LaCoO$_3$ and La$_{0.6}$Sr$_{0.4}$CoO$_3$ powders by ultrasonic spray pyrolysis

The synthesis of Co-based perovskites was realized by single-step USP procedures. The solutions for the synthesis of LC and LSC were prepared by mixing solutions of the starting precursors in stoichiometric mole ratios, i.e. La:Co = 1:1 for LC and La:Sr:Co = 3:2:5 for LSC.

An La: Sr ratio of 3:2 was chosen in order to prepare stoichiometric La$_{0.6}$Sr$_{0.4}$CoO$_3$ perovskite, according to a previously published work [6]. Aqueous 0.10 M solutions of La(NO$_3$)$_3$6H$_2$O (99.9% rare earth oxide), Sr(NO$_3$)$_2$ (99% ) and Co(NO$_3$)$_2$ (98%), all from Alfa Aesar, were used as precursors for the synthesis of LC and LSC. The USP conversion temperature was adjusted and controlled using a thermostated furnace. All powders were synthesized by ultrasonic spray pyrolysis in equipment with horizontal nebulia flow.

Fogging of the prepared precursors solutions occurred in an ultrasonic atomizer (Gapusol 9001, RBI/France) with an ultrasonic nebulizer (Prizma Kragujevac, Serbia) to create an aerosol [6,24]. The aerosol with droplets having diameter of around 2.3 µm was produced with an ultrasonic frequency of 2.5 MHz [6,25]. The atomization was performed in an O$_2$/N$_2$ atmosphere as carrier gas, having O$_2$ to N$_2$ volume ratio of 2:1 and continuous flow rate of 3 dm$^3$ min$^{-1}$. The synthesis temperature was adjusted to 600 °C or 800 °C for both LC and LSC, in order to investigate the influence of these USP synthesis conditions. The produced LC and LSC particles were collected in water bottle collectors.

3. Characterization techniques

3.1. Surface morphology and structural analysis

The morphology and elemental composition of LC and LSC powders were analyzed by scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDS). Scanning electron microscopes (Zeiss DSM 982 Gemini; Vega TS 5130 MM Tescan) were used to examine the appearance of the obtained particles. The elemental composition was determined by EDS with a Si(Bi) X-ray detector connected to the SEM and a multi-channel analyzer. The identification of elements was performed via the unique set of peaks from the reflected X-ray spectrum.

Structural and phase analysis of the composite samples were examined by X-ray diffraction (XRD) measurements on a Philips PW 1050 (Royal Philips, Amsterdam, the Netherlands) powder diffractometer at room temperature with Ni-filtered Cu Kα radiation (λ = 1.54178 Å) and scintillation detector within the 2θ range of 10°–82° in steps of 0.05°, at a scanning rate of 5 s per step. Phase analysis was performed using EVA v. 9.0 software.

3.2. Electrochemical measurements

Electrochemical properties of LC and LSC were studied by cyclic voltammetry (CV), potentiostatic electrochemical impedance spectroscopy (PEIS) and galvanostatic charge–discharge (GCD) tests. The electrochemical measurements were performed in a conventional three-electrode cell. A platinum wire and Ag/AgCl were used as counter and reference electrode, respectively. All potentials in the discussion are referred to Ag/AgCl. The working electrode was glassy carbon (GC, Sigraud – Sigri, Electrographite, GmbH, Germany) with a surface area of 0.196 cm$^2$ covered by a thin layer of LC or LSC according to the following procedure: 20 µL of LC or LSC water suspension (5 mg mL$^{-1}$) was added onto the GC using a micropipette. The mass loading of both LC and LSC electrode materials was 0.51 mg cm$^{-2}$. After 2 h of drying in air, the same volume of nafion solution (1:100 vs. water), achieved by dissolving 10 µL of nafion perfluorinated resin solution in 990 µL of water, was added on top of the layer and left to additionally dry at room temperature. This very thin nafion film was used for binding the investigated powders to the GC substrate. The cell was purged with N$_2$ for 30 min prior to CV electrochemical measurements. A potentiostat/galvanostat measuring station (BioLogic SAS, SP-240, Grenoble, France) provided with physical electrochemistry software was used. The CV responses in 0.10 M KOH were recorded at a scan rate of 50 mV s$^{-1}$. The specific capacitance, the most important indicator for the evaluation of electrode properties for supercapacitors [7,22,23], was calculated from the CV data according to Eq. (1):

$$C_{sp} = \frac{1}{m \ln |A|} \int_{Ea}^{Ec} I(E) dE$$

(1)

where $C_{sp}$ (F g$^{-1}$) is the specific capacitance, m (g) is the mass of the active material, $I_{sp}^{Ea} I(E) dE$ (A V$^{-1}$) is the integral area under the CV curve, which gives the charge stored during cathodic and anodic scans, $\nu$ (V s$^{-1}$) is the scan rate, and $E_a$ and $E_c$ (V) are the anodic and cathodic integration limits of the potential or the capacitive potential window.

PEIS studies were performed in the frequency range of 10$^{-2}$–10$^6$ Hz using a 10 mV root mean square sinusoidal potential amplitude around the potentials 0.1 and 0.3 V for all the samples. These potentials are recognized as characteristic values according to the CV responses. The impedance spectra were analyzed and fitted by ZView® software [28-30].

GCD tests were performed for cycle-life assessments of all the samples. These tests were also performed on potentiostat/galvanostat station (BioLogic SAS, SP-240, Grenoble, France). Galvanostatic charging and discharging curves were measured at a 50 µA current density until the 600 mV voltage window is spent (adopted from the CV measurements). The specific capacitances were calculated by graphical differentiation according to the following equation [18,31]:

$$C_{sp} = \frac{I dt}{m \Delta E}$$

(2)

where $C_{sp}$ (F g$^{-1}$) is the specific capacitance, $I$ (A) represent the charge/discharge current, dt (s) is the time required to spend the infinitesimal potential window, $\Delta E$ (V), and m (g) represents the mass of the active material.
4. Results and discussion

4.1. SEM and EDS analysis

The porosity of a material governs the maximal availability of the active sites \[6,25,26\]. The morphologies of the LC and LSC powders produced at 600 and 800 °C are presented in Fig. 1, which shows the typical appearance of the produced materials under SEM examination. The EDS elemental compositions of produced powders is presented in Table 1.

The lanthanum cobalt oxides powder synthesized at 600 °C (LC 600, Fig. 1a) is highly agglomerated and aggregated. The tight agglomerates consist of almost spherical particles in the size range of 50 nm – 200 nm. The temperature of the synthesis plays an important role. The influence of temperature could be seen by comparing powders from Fig. 1a and Fig. 1b. The later shows the appearance of lanthanum cobalt oxides powder synthesized at 800 °C (LC 800). The particles appear considerably less agglomerated, and the formed spherulites are in the size range of 1 µm. The initial particle sizes are also in the range of 50 nm – 200 nm, but the aggregation is not as pronounced as for the LC 600. The above mentioned features are more clearly seen in Fig. 2, which shows LC 600 and LC 800 at higher magnification.

The LSC powder synthesized at 600 °C (LSC 600) is shown in Fig. 1c. Comparing to Fig. 1a and Fig. 1b, the influence of Sr is evident. The powder is more spherical, more rounded and more compact, with clearly defined pores between the uneven spheres, which is in correlation with the results of previous research \[6\]. The sphere size of the powder is in the range of 1 µm – 2 µm, with the majority of particles having a diameter of 1 µm. The \(\text{La}_{0.6}\text{Sr}_{0.4}\text{CoO}_3\) powder synthesized at 800 °C (LSC 800) is shown in Fig. 1d. The influence of synthesis temperature on the powder morphology is evident once again, along with the fine sphere-generating influence of strontium oxide. The powder particles are well defined fluffy spheres with the powder sphere size in the range of 0.5 µm – 1.5 µm. It could be noticed from Fig. 1 that both the synthesis temperature and presence of strontium ions influence the morphology of the powder in the sense that spherical particles become well defined, more rounded and larger. The unique finding is that strontium forms fluffy structure at the surface of the spherical particles at higher temperatures. This could indicate that some structural composition occurs upon insertion of strontium oxide at higher temperatures.

EDS composition mapping was performed on the powder particles at different spots and different samples and the average value of these elemental compositions are listed in Table 1.

As it could be seen in Table 1, the detected content ratios of Co to La for LC powders are considerably higher with respect to stoichiometric 1:1. As the material becomes less compact and more porous at higher temperature (Figs. 1 and 2), the ratio indicates slightly higher presence of La. This enrichment in Co also holds for the LSC sample. However, the contents of La and Sr are lower at higher temperature, which also generates the fluffy surface structure of the spheres. It appears that the Co oxide tends to grow into surface needles, which mask the EDS response of the La and Sr oxides. The stoichiometric ratio of La to Sr of 3:2 appears well preserved in the LSC samples. It could be hypothesized that besides perovskite LC and LSC powders, some separate phases of Co-based oxides are formed. This hypothesis was further investigated by XRD measurements.

4.2. X-ray diffraction analysis

The XRD patterns of LC 600, LC 800, LSC 600 and LSC 800 powders are presented in Fig. 3. Sharp peaks appear in the 2θ range of 10–80° for all the samples. These XRD peaks could be assigned to LC and LSC compounds with the sharpness indicating the highly crystalline sin-
gle-phase nature of the samples with a rhombohedral crystal structure. The synthesized LC and LSC powders as perovskites, with variation in the crystal structure that can lead to distinct electrocatalytic activities \[6,27\]. The diffraction peaks of Co$_3$O$_4$ (JCPDS PDF No.01-078-1969), La$_2$O$_3$ (JCPDS PDF No.01-074-2430), CoO (JCPDS PDF No.01-089-2803) and SrO (JCPDS PDF No.00-048-1477) could be distinguished in the XRD patterns of LC and LSC. It can be derived from these XRD findings that there is formation of nanoparticles of above mentioned compounds.

Besides the good correlation to the development of perovskite structure (mixed oxide) for all the samples it could be seen that there is some significant quantity of a separate Co$_3$O$_4$ crystal phase in the LC 600, LC 800 and LSC 600 samples, and to very small extent of CoO phase in the LSC 800 sample. This confirms that the generation of separate Co oxide phase, beside that incorporated into perovskite structure, could aggregate separately within the powder particles, as indicated by the EDS measurements.

Apart from separate phases of Co oxides, the XRD patterns also indicated the appearance of separate La$_2$O$_3$ (for LC) and SrO (for LSC) phases. The peaks of La$_2$O$_3$ are more pronounced in the high temperature sample, which agrees with higher La/Co ratio found by EDS.

### 4.3. Electrochemical analysis and performances

Stable CVs of LC 600, LC 800, LSC 600 and LSC 800 thin powder layers on GC substrate in 0.10 M KOH (nitrogen atmosphere) were recorded at the scan rate of 50 mVs$^{-1}$, and the results are shown in Fig. 4.

The cycling started from the open circuit potentials (OCP) for the prepared electrodes: $-0.165$ V for LC 600, $-0.062$ V for LC 800, $0.35$ V for LSC 600 and $0.265$ V for LSC 800. Although the OCPs were more positive when Sr was present, phenomenologically there is no great difference in the material response whether strontium was present or not. On the other hand, Sr influences development of the anodic branch in the materials synthesized at the lower temperature. Comparing the voltammograms of the obtained material to the ones for the Co$_3$O$_4$ in the same potential range of $-0.1$ V to $0.6$ V in KOH electrolyte it could be concluded that some redox peaks appear

<table>
<thead>
<tr>
<th>Element/ at.%</th>
<th>LC 600</th>
<th>LC 800</th>
<th>LSC 600</th>
<th>LSC 800</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>61.8</td>
<td>69.0</td>
<td>64.1</td>
<td>64.2</td>
</tr>
<tr>
<td>Sr</td>
<td>–</td>
<td>–</td>
<td>4.2</td>
<td>2.8</td>
</tr>
<tr>
<td>La</td>
<td>1.9</td>
<td>2.3</td>
<td>6.5</td>
<td>4.5</td>
</tr>
<tr>
<td>Co</td>
<td>36.3</td>
<td>28.7</td>
<td>25.2</td>
<td>28.5</td>
</tr>
</tbody>
</table>

**Table 1**

EDS comparative analysis of LC 600, LC 800, LSC 600 and LSC 800 as-prepared powders (at.%).

Fig. 2. SEM microphotographs of lanthanum cobalt oxides powders synthesized at: a) 600 °C and b) 800 °C recorded at magnification of 50 k.

Fig. 3. XRD patterns of a) LC and b) LSC powders.

Fig. 4.
in the potential range of 0.4–0.6 V and are usually assigned to redox transitions of CoOx to CoOOH and CoOOH to CoO2 [32]. The other weak broader peaks that appear outside this range could belong to the redox reactions of the rare earth metal. These curves are not close to the symmetric shape presented by an electrochemical double layer capacitor which has a closed rectangular shape and stores energy through non-Faradaic processes [32]. The voltammograms show forms that are typical of materials that are strongly governed by Faradaic processes with defined anodic and cathodic peaks that represent pseudo-capacitive behavior. An effective symmetry might be associated with good reversibility of oxidation and reduction processes [33]. Poor linear behavior is affirmation of the pseudo-capacitive behavior of the electrode [34]. It was also shown that displacement of the oxidation peaks towards positive potentials and reduction peaks towards negative ones may be related to conductivity and to the polarization-induced capacitive ability of the electrode [35].

On comparing the LC and LSC CV curves, some differences could be noted. One of the major differences is the fact that the lowest CV currents are registered in the CV response of LC 600 and then in LC 800, which is an indication of the smallest active surface area of these electrode materials. Another difference reflects the fact that incorporation of Sr into the LC lattice leads to a considerable increase in the currents for Sr-containing samples. The temperature of the synthesis of anodic peaks towards positive potentials and reduction peaks towards negative potentials is not clearly evident at lower potentials. This promotes Sr, besides incorporation of Sr into the LC lattice leads to a considerable increase in the currents for the Sr-containing samples. The temperature of the synthesis of anodic peaks towards positive potentials and reduction peaks towards negative potentials is not clearly evident at lower potentials. This promotes Sr, besides Sr, higher temperature produces particles with higher specific surface area (Fig. 1), and hence having higher values of the specific capacitance, the inverse is applicable for the LSC samples. This could be associated to the Co oxide enrichment (opposite to LC, Table 1), which was found to cause the formation of the needles of the fluffy structure of the sphere surfaces. However, LSC 600 has higher specific capacitance compared to LSC 800, which does not present the benefits of needles formation. For better capacitive response of LSC, it seems that the not fully separated spheres of smooth surface (LSC 600, Fig. 1c) are beneficial. In addition, there is pronounced effect of multiple oxide structures present in the sample [6], especially separate CoO4 phase (Fig. 3). Besides presence of separate cobalt oxide, the presence of strontium in perovskite structure leads to high electrode/electrolyte contact areas and high rates of electrode reaction [6], since an enhancement due to separate Co3O4 was not registered in the LC samples.

These findings are in good agreement with previously published results [6] for ruthenized LSC. They all confirm the statements that strontium influences the development of the anodic branch, and that the processing temperature plays an important role in electrode material production. It appears that not only Sr promotes the capacitive response of LC, but also LSC favors and enhances redox transition of material in higher oxidation states.

Potentiostatic electrochemical impedance spectroscopy (PEIS) tests were performed at potential of 100 (poor conductive region, Fig. 4) and 300 mV (fully developed redox transitions) to evaluate the details of capacitive response and electrochemical behavior for all of the samples, including hydrothermally prepared Co3O4 for the sake of comparison. PEIS of LC 600, LC 800, LSC 600 and LSC 800 thin powder layers on GC substrate were recorded in 0.1 M KOH at a perturbing potential of 10 mV in the frequency range between 10 mHz and 1 MHz and the results are shown in Fig. 5. Based on the PEIS data, Fig. 6 shows equivalent circuit (ECC) models used to fit the data from Fig. 5 (ECC data are given by lines in Fig. 5).

The complex plane plots for LC and LSC all show two distinct regions. The first region consists of semicircle-like response at the high frequencies which corresponds to redox transitions and it reflects the charge transfer resistance-like (R3, Fig. 6) behavior of the electrode. The second region, represented by straight line in the low-frequency range, corresponds to ionic exchange in the electrolyte/material interphase – external (CPE1) and internal (CPE2–CPE4) [36,37]. The high-frequency intersection of the curve with the real axis represents the resistance of the aqueous KOH solution (R1). The internal capacitive response (CPE2–CPE4) required series ionic (to CPE2 and CPE4) or material bulk (to CPE3) resistance represented by R2 in Fig. 6.

The results obtained for the examined materials from fitting results using ECCs are given in Tables 2 and 3. The constant phase element, CPE, is used in EEC instead of a capacitor, C, in order to better embrace the non-ideal behavior of the C element, namely to address the surface heterogeneities, surface roughness, as well as defects on the surface. In general, Eq. 3 is used as the definition of the impedance of the CPE:

\[ Z_{\text{CPE}} = |Y(j\omega)|^{-1} \]

where \( Y \) is the frequency-independent admittance of the CPE, \( \omega \) being the angular frequency (\( \omega = 2\pi f \)) in rad s\(^{-1} \), \( f \) is the frequency, \( n \) is the value of the exponent of CPE between −1 for an ideal inductor and 1 for an ideal capacitor.

The CPE1 element has the capacitive response that resembles the capacitive response of pure Co3O4. The C1 values of capacitance at 300 mV after fitting the results (Tables 2 and 3) are close to the value of pure Co3O4 (3.5 \( \mu F \)) which was fitted by simple RC EEC in series (Fig. 6c). This finding is in accordance with the XRD results of the prepared powders, which show the presence of separate Co3O4 phase besides the perovskite structure. The rest of the EEC circuit show interaction of Co3O4 with other oxide structures in the electrode material.

**Fig. 4.** Cyclic voltammograms of LC 600, LC 800, LSC 600 and LSC 800 at a scan rate of 50 mV s\(^{-1} \) in 0.10 M KOH in N\(_2\) atmosphere at room temperature.
Fig. 5. Complex plane and phase shift (ϕ) plots of LC 600, LC 800, LSC 600 and LSC 800 powders, with insets showing enlarged high frequency region PEIS analysis at a) 100 and b) 300 mV.

Fig. 6. The equivalent electrical circuits used to fit the impedance spectra of: a) LC 600, LC 800 and LSC 800 recorded at 100 mV and 300 mV and LSC 600 recorded at 100 mV, b) LSC 600 recorded at 300 mV and c) Co$_3$O$_4$ recorded at 300 mV.
Comparing the results from Tables 2 and 3 it could be noticed that the resistance that precedes the capacitive response of the material is larger by up to one order of magnitude for the measurements performed at 100 mV. The modest capacitive response of the material could be seen when compared to the response at 300 mV. The values of the calculated total capacitance, $C_{\text{tot}}$, are considerably lower than the values obtained by CV. This indicates that material needs a much wider perturbing measuring potential range than 10 mV (PEIS) in order to develop a full capacitive response (700 mV in CV). This is best seen in the PEIS results and EEC of the LSC 600 sample recorded at 300 mV (Table 3) where there is the considerably highest capacitance value of C3 being 12,963 $\mu$F. However, it is in series with the rest of the EEC, which gives the total capacitance of 589 × 10^{-2} $\mu$F g^{-1}. This series capacitance is also characteristic for all other samples, and has the highest value in EEC. This apparently indicates that full development of the capacitive response requires some preceding reactions to occur. Since CV gives higher values, these preceding reactions could take place out of the potential range spent in PEIS perturbation. Hence, in order for full capacitive response to be evidenced, there is need for consecutive electrochemical transformations to occur.

As it was indicated by CV measurements, PEIS measurements proved that strontium promotes the structure and pseudocapacitive behavior of the synthesized perovskite material. The former statement is in good agreement with the findings that porous metal oxide structure with high specific surface area is responsible for the high $C_{\text{sp}}$ values since it enables and promotes the Faradaic reactions. Sr promotes the capacitive response of LC and it favors and enhances redox transitions of the material in higher oxidation states.

The charging-discharging stability and rate capability towards potential applicability of the LC and LSC powders as supercapacitor electrode materials was further tested and examined by galvanostatic charge–discharge (GCD) tests. The GCD measurements were performed in 0.10 M KOH at currents of 50 and 200 mA. The results of GCD measurements of LC 600, LC 800, LSC 600 and LSC 800 powders are shown in Fig. 7.

As was discussed and explained in relation to Fig. 4, typical pseudocapacitive behavior is reflected in some deviation of the curves from full linearity [24,38]. GCD curves lack mirror symmetry in the investigated potential range due to redox reactions that occur on the electrode surface [39]. The deviation appears not dependent on the GCD rate, which indicates that related redox transitions are uniformly distributed throughout the material. At a glance, Fig. 7A–C shows the higher stability at higher currents, due to indicated distribution of the components and phases as discussed in relation to Table 1 and Fig. 3. Specific capacitances for the first and 500th cycle, gained as the mean by graphical differentiation of G-C/DGCDC curves (since a rather nonlinear response was registered (Fig. 7) [40]), according to Eq. (2). All of the calculated values for the specific capacitance obtain from GCD curves follow the order found by CV (Fig. 4) and PEIS measurements (Table 3).

The LC 600, LC 800 and LSC 600 GCD curves are almost identical in shape, representing a bell-like shape variation of the potential with time, which is a typical supercapacitor behavior, while the charge curve for LSC 800 has a half-cycle shape if compared to a more linear discharge curve as the corresponding counterpart, Fig. 7A–C. These findings are in accordance with previous research [6]. The difference between the investigated samples is that the LC 600, LC 800 and LSC 800 samples have an almost similar discharge time, while LSC 600 discharge time is almost twice of that for high-temperature samples. Although LSC 800 is of highest stability in charge/discharge cycles, it is of poor capacitive ability at the level of the LC samples.

Although the CD stability of the synthesized samples are more (high rates) or less (low rates) similar, the rate capabilities are quite different, Fig. 7D and E. It could be noted that the shape of the CD curves does not change much with the CD rate. However, the decrease in CD abilities upon increasing in CD rate was registered for all samples. The full-cycle charge exchanged at low (LR) and high (HR) CD rates, as well as that separated to half-cycles at low (LC) and high (HC) charging rates are presented in Table 4. The relative decrease of the charge exchanged with respect to low CD rates are given as corresponding $\delta$ values.

The LC 600 sample, being of lowest capacitance, also shows the lowest sensitivity to the CD rate. The decrease in charge exchanged is around 24 %, with the charging rate suffering more (27 %) than the discharging values (21%). This indicates that the interior of densely packed material (Fig. 2A) are hardly accessible at high CD rates. Once charged, however, the outer layers are relatively available more for discharge in comparison to the other samples.

Considerable larger decreases of CD rate capability are found for the other samples, with the highest one being for LSC 600 sample during charging. The mutual characteristic of LC 800, LSC 600 and LSC 800 is the initiation of (LC 800) towards the full generation of spherical structures (LSC samples), Fig. 1. Apparently, the CD dynamics “penetrate” harder into the dense internal structure of the sphere (Fig. 3). The highest decrease during the charging cycle is found for the sample of stuck spheres with a smooth surface (Fig. 1C). Consequently, LSC 600 sample suffers the most from charging rate capability decrease (59 %), but also shows the largest difference between the $\delta_{\text{HC}}$ and $\delta_{\text{LR}}$ values. This consideration is in accordance with that explained in relation to rate capability of the LC 600 sample.

Considering all previous investigations of the microstructure (SEM, EDS and XRD) to the electrochemical performances (CV, EIS and GCD) LSC 600 shows superior characteristics with respect to LC and its low temperature twin. All of these findings show the promising influence of strontium on the pseudocapacitive properties, as well as the influence of the processing temperature that plays an important role in the pro-

<table>
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<th>Sample</th>
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<th>R2(Ω)</th>
<th>R3(Ω)</th>
<th>C1(μF)</th>
<th>C2(μF)</th>
<th>C3(μF)</th>
<th>$C_{\text{sp}}$ ($\times 10^{-2}$ g^{-1})</th>
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duction of electrode material. It could be concluded that LSC 600 is promising supercapacitor electroactive material with fine reversible charge–discharge performances.

5. Conclusions

The influence of strontium oxide doping into LC shows that the synthesis temperature plays an important role in the characteristics of the final material, with particles becoming more spherical and less aggregated with increasing temperature. The addition of strontium oxide further defines the formation of more spherical powders (LSC). EDS and XRD analyses of the powders suggest that that besides LC and LSC perovskite powders, some other Co-based oxides are formed. The voltammograms show that the materials are strongly governed by Faradaic processes. Incorporation of Sr into the LC lattice leads to a considerable increase in the currents at positive potentials. The specific capacitances of the LSC samples are higher (2–4-fold) compared to the LC samples, with LSC 600 having the highest $C_{sp}$ value.

Calculated PEIS capacitance values are lower than CV obtained values because the material needs a much wider CD potential range in order to develop full capacitive performance. PEIS shows that strontium promotes the structure and pseudocapacitive behavior of the synthesized material, favoring and enhancing the redox transition of materials in higher oxidation states. Specific capacitance results of GCD measurements are in accordance and comparable with the values obtained from CV and PEIS measurements. The LC samples showed the lowest sensitivity to the CD rate, with the charging values suffering more than the discharging values, indicating that interior of densely packed materials are hardly accessible at high CD rates. Once charged, the outer layers are relatively more available for discharge in comparison to other samples. LSC 600 sample suffers the most from charging rate capability decrease, but also shows the largest difference between charging and discharging rates.

However, it is important to know that PEIS measurements indicated that full potential window cycling within electrolyte stability is required to gain full interactive capacitive performances of the obtained oxides as supports for supercapacitor materials.

CRediT authorship contribution statement

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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