IR characterization of Ln₂-xSrxCoO₄ (x≥1; Ln=La, Nd) oxides

S. Castro-García ^a, M. Sánchez-Andújar^a, C. Rey-Cabezudo^a, M.A. Señarís-Rodríguez^a, C. Julien^b

^a Dpto. Química Fundamental, Universidad de A Coruña, 15071 A Coruña, Spain

^b LMDH, Université Pierre et Marie Curie, 4 place Jussieu, 75252 Paris, France

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Abstract

We have recorded the FTIR spectra of powder samples of Ln_2 -xSrxCoO $_4$ (Ln=La, Nd) at room temperature. We have identified the infrared active modes ($3A_{2u}$ + $4E_u$), and analyzed how they change as a function of Ln and the Sr doping. We correlate the obtained results with structural data obtained from powder X-ray diffraction studies and with the electronic properties displayed by these samples.

Keywords

Magnetically ordered materials; Chemical synthesis; Crystal structure and symmetry; Phonons; Light absorption and reflection

1. Introduction

Ln_{2-x}SrxCoO₄ oxides (Ln=La, Nd) are systems that crystallize in the K₂NiF₄-type structure (Fig. 1), that is the 2D analogue of the perovskite type [1]. Their structure has been described as a sequence of tetragonally distorted [CoO₆] octahedra, the Ln³⁺ and Sr²⁺ ions being located in 9-coordinated sites between the layers. These compounds closely related to $Ln_1-xSrxCoO_3-\delta$ perovskite are materials [2], [3], [4], [5], [6], [7], [8] and [9], that have attracted a lot of attention in view of how their magnetic and electrical properties can be varied by changing the rare earth [4], [5] and [7] and the degree of Sr-doping [3] and [6]. Nevertheless $Ln_2-xSrxCoO_4$ (with $x\ge 1$) compounds have been relatively less studied, probably due to the difficulty to synthesize them and to stabilize the Co³⁺, Co⁴⁺ formal states in the K₂NiF₄ structure. The magnetic and electrical properties of the Ln_{2-x}SrxCoO₄ also change with x [10] and [11]: their electrical resistivity decreases as x increases, even if the metallic regime is not achieved; also, a ferromagnetic contribution starts to develop for x>1.2 at T≤175 K even if to a lesser extend than in the corresponding Ln_2 -xSrxCoO $_3$ - δ materials.

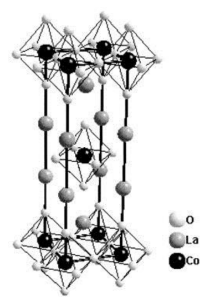


Fig. 1.
The K₂NiF₄-type structure.

To get more information about these relatively poorly characterized Ln_2 -xSrxCoO $_4$ materials, we have studied them by means of powder X-ray diffraction and IR spectroscopy. In this paper we present and discuss the results obtained for Ln=La and Nd with $1.0 \le x \le 1.4$ and $1.0 \le x \le 1.3$, respectively.

2. Experimental

Samples of La_{2-x}SrxCoO₄ (1.0 \le x \le 1.4) and of Nd_{2-x}SrxCoO₄ (1 \le x \le 1.3) were prepared by a nitrate decomposition method. Stoichiometric quantities of La₂O₃/Nd₂O₃, SrCO₃ were dissolved in HNO₃ (30%) followed by the addition of stoichiometric amounts of Co(NO₃)₂·6H₂O. The resulting solution was gently warmed up so as to slowly evaporate the solvent. To facilitate the formation of highly oxidized samples, a melting agent, KNO₃, was then added to the so-obtained mixture of nitrates, in a mole ratio 1 KNO₃:1 Co(NO₃)₂·6H₂O. This mixture was then treated in air at 400°C for 1 h, 600°C for 48 h, and 975°C for 24 h with intermediate grinding after each heating step. X-ray powder diffraction measurements were carried out with a Siemens D-5000 diffractometer and CuK_a=1.5418 Å radiation. Semiquantitative EDAX analysis was carried out in a scanning electron microscope (model JEOL 6400) at 40 kV. FTIR absorption spectra were recorded at room temperature using an interferometer (Brucker model IFS113v) equipped with a 3.5-µm-thick beamsplitter, a global source, and a DTGS/PE far-infrared detector. Samples were ground to fine powders into ICs pellets. Data were collected in transmission mode at a spectral resolution of 2 cm⁻¹ after 256 scans in vacuum atmosphere.

3. Results and discussion

3.1. XRD characterization

According to their XRD patterns, all compounds were obtained as single-phase crystalline materials and were K^+ -free, as shown by EDAX analysis. This means that a complete solid solution between Ln and Sr can be formed in $Ln_{2-}xSrxCoO_4$ in the range $1 \le x \le 1.4$ for Ln=La and $1 \le x \le 1.3$ for Ln=Nd. These compounds crystallize in the K_2NiF_4 structure, that is body-centered tetragonal, space group I4/mmm with Z=2. This structure can be thought to consist of a perovskite layer separated by a rock-salt type $(Ln,Sr)_2O_2$ layer along c-axis. The cobalt atoms are located at the 2a site, the (La or Nd)/Sr randomly occupy the 4e site. There are two types of oxygen atoms, O(4c) located in the ab planes and O(4e) located in the $(Ln,Sr)_2O_2$ block above and below the cobalt atoms. Consequently, the cobalt atoms are in a distorted octahedral environment, where there are two different Co-O bonds: an axial Co-O(I) bond (in the ab plane) shorter than the apical Co-O(II) bond (along the c direction).

Fig. 2a and b show the variation of the lattice parameters and the cell volume as a function of x in Ln_{2-x}SrxCoO₄ compounds for Ln=La and Nd. As it can be observed in these figures, in both series the cparameter clearly increases with doping, whereas the a parameter changes very slightly. The cell volume in Nd₂-xSrxCoO₄ series increases with Sr content. In the La_{2-x}SrxCoO₄ system the same trend is observed when x is small, but for x>1.2 it remains almost constant. To explain these variations we need to take into account that when Ln is substituted by Sr, a larger size Sr⁺² ion $(^{IX}r_{Sr+2}=1.30 \text{ Å})$ is incorporated in place of smaller size Ln^{3+} ion $(^{IX}r_{Nd3+}=1.16)$ \mathring{A} , IX r_{La+3}=1.216 \mathring{A}) [12]. On the other hand, upon doping Co^{3+} (VI r_{Co3+(l.s.)}=0.55 Å, $^{\text{VI}}$ r_{CollI+(h,s.)}=0.61 Å) oxidates to the smaller Co⁴⁺ ions ($^{\text{VI}}$ r_{Co4+(h,s.)}=0.53 Å) [12]. In the case of the Nd-cobaltites, where there is a bigger size difference between Ln3+ and Sr^{2+} , the former factor predominates, so that the cell volume and the c parameter are seeing to increase with x. Nevertheless, in the case of the La-cobaltites where the size difference between La3+ and Sr2+ is relatively smaller the former factor dominates only for small x (x<1.2). For higher x, the two factors — that work in opposite directions are balanced, their effect gets cancelled and neither the volume nor the c parameter change upon further doping.

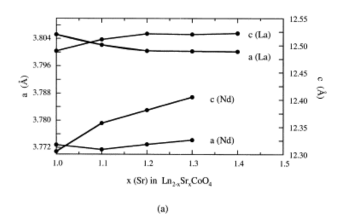
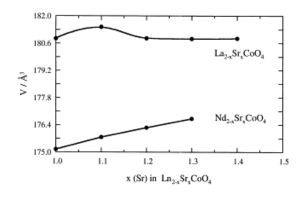


Fig. 2. Evolution of (a) the lattice parameters and (b) the cell volume as a function of x in $Ln_2-xSrxCoO_4$ compounds with Ln=La and Nd



It should be noted that small differences in structural parameters such as Co–O distances and the degree of distortion of the [CoO₆] octahedra are observed among the La and Nd series of compounds. In this context, Table 1 gives the Co–O(I) and Co–O(II) distances corresponding to LnSrCoO₄ compounds (Ln=La and Nd) calculated by Rietveld analysis.

Table 1.

Co–O distances in LaSrCoO₄ and NdSrCoO₄ compounds as obtained from XRD Rietveld analysis. d(Co-O(I)) corresponds to the axial bond in the *ab* plane while the d(Co-O(II)) corresponds to the apical bond along the *c* direction

Compound	d(Co–O(I)) (Å)	d(Co–O(II)) (Å)		
LaSrCoO ₄	1.9026	2.0374		
NdSrCoO ₄	1.8864	2.0135		

3.2. FTIR characterization

The factor group analysis of the K_2NiF_4 structure with I4/mmm (D_{4h}^{17}) space group indicates that there are seven infrared-active modes ($3A_{2u}+4E_u$) and three Raman-active modes; we can describe them following the interpretation of Daturi et al. [13] for Nd_2CuO_4 compounds that distinguishes 'internal' vibrations of the $[CuO_2]_n$ sheets, from internal vibrations of the $[Nd_2O_2]_n$ blocks and the lattice vibrations of these two units.

Fig. 3a and b show the FTIR spectra that we obtain for Ln_2 -xSrxCoO $_4$ compounds (Ln=La and Nd, respectively) that are quite similar to those shown by other oxides with K_2 NiF $_4$ structure [14] and [15], andTable 2 and Table 3 list the IR bands observed together with their assignment. Taking into account that powder morphology can affect these spectra [16], the position of the observed bands do not correspond to the transverse optical (TO) modes but are more or less shifted towards the longitudinal optical (LO) modes. Despite the fact that the LO-TO splitting can be large in oxide compounds, the phonon values have been evaluated from absorption experiments on powdered samples, for which FTIR reflectivity can not be accurately measured.

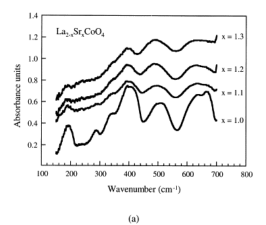


Fig. 3. FTIR spectra of $Ln_2-xSrxCoO_4$ compounds (Ln=La, Nd). The IR spectra have been vertically shifted according to their x content so that they can be easily seen.

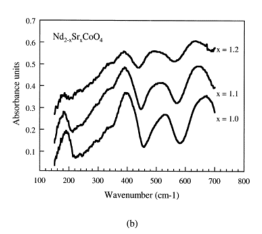


Table 2. Wavenumber (cm^{-1}) and assignment of the IR bands of the LaSrCoO $_4$ and NdSrCoO $_4$ compounds

Symmetry	v(LaSrCoO ₄)	v(NdSrCoO ₄)	Assignment
Eu	670,640	666	Asymmetric Co-O(I)-Co stretching in
			the ab plane
A_{2u}	510	528	Out of plane Co-O(I)-Co deformation
Eu	400	397	In plane Co-O(I)-Co deformation
A_{2u}	350	344	Deformations of (La or Nd)/Sr-O(II)
Eu	290	293	Stretching of (La or Nd)/Sr-O(II)
A_{2u}	240	237	Lattice mode:
			Motion of Co-O(I)-Co layers against the
			(La or Nd)/Sr-O(II) blocks
Eu	190	190	Lattice mode:
			Sliding of the Co-O layers with respect to
			(La or Nd)/Sr-O(II)

Table 3. Wavenumber (cm⁻¹) and assignment of the IR bands of the Ln₂₋xSrxCoO₄ compounds

Symmetry	La ₂₋ xSrxCoO ₄				Nd ₂ -xSrxCoO ₄		
<i>-</i> ,	<i>x</i> =1.0	<i>x</i> =1.1	<i>x</i> =1.2	<i>x</i> =1.3	<i>x</i> =1.0	<i>x</i> =1.1	<i>x</i> =1.2
Eu	670, 640	631	631	631	666	640	627
A_{2u}	510	492	483	483	528	510	492
Eu	400	391	391	391	397	396	389
Eu	189	185	185	185	189	183	180

The first interesting point to note about the FTIR spectra is that in both series of compounds the bands that appear in the high frequency region ($v>400 \text{ cm}^{-1}$) correspond to modes related to Co–O stretching and deformations; the bands that appear in the range $250 < v < 400 \text{ cm}^{-1}$ correspond to modes related to (Ln,Sr)–O vibrations and those appearing at $v < 250 \text{ cm}^{-1}$ correspond to modes related to lattice modes.

If we compare the spectra of the LaSrCoO₄ with those of the corresponding NdSrCoO₄ compound, we can give a mere estimation on the frequency shift due to the mass and Co–O distances changes, taking into account that the vibration energy is proportional to the square root of the bond strength and inversely to the square root of the atomic mass. We observe a slight frequency shift of the Co–O deformation modes due to the shorter bonding distances in NdSrCoO₄ (Table 1). As LaSrCoO₄ are lighter than NdSrCoO₄ we observe a frequency shift of the A_{2u} mode (corresponding to the deformations of the Ln, Sr–O bonds) from 344 to 350 cm⁻¹ which is in good accordance with the mass change.

If we compare how the spectra change as a function of x in a given $Ln_2-xSrxCoO_4$ series, the most remarkable feature is an increasing screening effect of the in-plane ab modes (see Fig. 3). This effect can be explained on the basis that upon doping charge carriers are created, resulting in an increase of the bidimensional conductivity of the $[CoO_2]_n$ sheets, as corroborated in fact by means of electrical conductivity measurements [17]. Nevertheless, none of these systems achieves the metallic regime in the compositional studied range.

For Nd-cobaltites, the Sr doping induces an increasing unit cell volume with higher Co–O distances. This results in the lowering of FTIR frequencies. For La-cobaltites we observe two different behaviors: (1) for $1.0 \le x \le 1.1$, a larger unit-cell volume with higher

Co–O distances is observed, resulting in a lowering FTIR frequencies and (2) for $1.1 \le x \le 1.3$, we do not observe any variations in both the lattice parameters and FTIR mode frequencies.

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Corresponding author. Tel.: +34-981-167-000; fax: +34-981-167-065 suquie@udc.es