

Substitution site and effects on magnetism in Sr-for-Ca substituted CaBaCo₄O₇

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Cationic substitutions in the novel magnetoelectric compound CaBaCo₄O₇ lead to profound changes in its magnetic and electric behaviors. In this work, we present a structural study of the isovalent substitution Sr-for-Ca in CaBaCo₄O₇. X-ray diffraction, as well as neutron powder diffraction experiments, are reported for a series of samples $Ca_{1-x}Sr_xBaCo_4O_7$ with $0 \le x \le 0.10$. Special emphasis is given to the identification of the substitution site, as Sr has also been reported to substitute for Ba in this crystal structure. The solubility limit for Sr at the Ca site is shown to be at $x \simeq 0.08$. The variation of lattice constants with Sr-doping firmly supports the Sr-for-Ca substitution. Rietveld refinements of the Sr-substituted samples are presented, and used as starting point to analyse the local structure around Sr by means of X-ray absorption spectroscopy at the Sr K-edge. Both the near-edge absorption and the extended absorption fine-structure confirm the substitution of Sr for Ca, giving definite support to the proposed nominal formula. In addition, macroscopic magnetization measurements are presented which reveal the striking effects of Sr-substitution over the magnetic landscape of this puzzling compound. © 2015 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4931615]

I. INTRODUCTION

During the last decade, a novel member of the cobaltrich oxides family has gained increasing attention. The so-called "114"-cobaltate $RBaCo_4O_7$ was discovered in 2002 for R=Y (Ref. 1) and, since then, synthesized with other lanthanides. This compound displays at least two very attractive features. First, it offers the possibility to study frustrated magnetism in the unique topology of a three-dimensional framework of Co tetrahedra in interconnected triangular and Kagomé lattices. Geometric frustration is related to intensively studied phenomena in spin ices and multiferroic materials; and this compound opens a new possible scenario for its study. Second, it shows an outstanding capacity of reversible absorption and desorption of gases, which turns these cobaltates promising candidates for oxygen-storage and gassensors applications. 4,5

In 2009, Caignaert *et al.*⁶ synthesized this compound for R = Ca which triggered even more exciting features, as the Ca-member, by changing the mixed valence state of Co, the structure, and the charge order, dramatically varies the magnetic and electric properties. The CaBaCo₄O₇ compound was shown to become

In this work, we aim at studying the effect of an isovalent substitution at the Ca site by replacing Ca²⁺ with Sr²⁺. Other cationic substitutions at the Co and Ba sites have shown to affect the magnetic properties. This paper focuses in the room-temperature structural aspects of the Sr-substitution, given the importance of correctly identifying the substitution sites in the structure. To this end, we have performed not only diffraction experiments using X-rays and neutrons but also studied the local environment of Sr atoms by means of X-Ray absorption spectroscopy (XAS). Finally, we analyse the striking effects that this substitution has on the magnetic properties, reporting macroscopic dc and acmagnetization measurements as a function of temperature and frequency.

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ferrimagnetic below 60 K (Refs. 7 and 8) with a complex non-collinear arrangement of spins, and it was later discovered that it concomitantly displays a dramatic increase in electric polarization of $17\,000\,\mu\text{C/cm}^2$, driven by exchange-striction. 9,10 Whereas it was initially thought that the compound was ferroelectric, 11 it is rather defined as a ferromagnetic pyroelectric, as the polarization cannot be switched by an external electric field. 10,12 These very recent findings open up the route to new magnetoelectric and multiferroic materials in which the structural properties play a major role.

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II. EXPERIMENTAL METHODS

A. Synthesis

Polycrystalline samples of nominal CaBaCo₄O₇ (hereafter called Sr0%) and Ca_{1-x}Sr_xBaCo₄O₇ with x=0.02 (Sr2%), x=0.05 (Sr5%), x=0.07 (Sr7%), and x=0.10 (Sr10%) were prepared by conventional solid-state reaction. High-purity powders of previously dried Y₂O₃, BaCO₃, SrCO₃, and Co₃O₄ were thoroughly mixed in an agate mortar at stoichiometric weights. After a de-carbonation process at 1173 K for 12 h, the mixtures were pressed into pellets. Pellets were annealed during 12 h–15 h at 1373 K and cooled at 3 K/min in the furnace. After a regrinding, the compression and annealing at 1373 K in air were repeated and as a final step, samples were quenched from 1373 K to L-N₂ to obtain the stoichiometric compound. ^{6,8}

B. X-ray diffraction

Room-temperature X-ray powder diffraction (XRPD) data were collected using a PANalytical Empyrean diffractometer with Cu- K_{α} radiation to check the phase purity. A small amount of a secondary phase was detected for sample Sr10%, which was identified as a perovskite belonging to the (Sr, Ba)CoO_{3- δ} family, most likely Sr_{0.2}Ba_{0.8}CoO_{2.3} (reference code ICSD 260871). We relate this to the solubility limit for Sr, as we discuss later in the manuscript.

C. Neutron diffraction

High-resolution neutron powder diffraction (NPD) experiments were performed at the Institut Laue-Langevin (ILL) in Grenoble, France. NPD data at room temperature were collected in the D2B diffractometer for samples Sr2% and Sr5%. A wavelength of ~ 1.59 Å was used, with an angular span of 150° and a step of 0.05° . The collimated beam on the sample was of $32 \, \text{mm} \times 12 \, \text{mm}$. A NaCaAlF standard was used to calibrate the neutrons wavelength, yielding the value $\lambda = 1.5943 \pm 0.0001$ Å. Measurements were performed in vanadium cylinders of 6 mm diameter and 8 cm height filled with sample, with a collection time of 4 h. Data were processed with the LAMP software 17 and the diffractograms thus obtained were refined with the full-pattern analysis Rietveld method, using the program FULLPROF. 18

D. XAS

XAS is particularly powerful to determine the substitution site of dopants, because it is an element-selective tool, allowing to study the local environment of selected atomic species. Extended X-ray absorption fine structure (EXAFS) spectra at the Sr K-edge (16 105 eV) were collected in fluorescence mode at the DIFFABS beamline of the Soleil synchrotron (France). EXAFS spectra were acquired with a step of 1 eV in the near edge region (16–16.4 keV), and 2 eV in the EXAFS region (up to 16.8 keV). The acquisition was done using a four-element silicon drift detector. A double-crystal monochromator equipped with two silicon (111) crystals was used. Energy was calibrated using a 25 μ m Y foil (K edge: 17 038 eV). Three scans were collected at room

temperature and ambient pressure for a total acquisition time of 3 h.

X-ray Absorption Near-Edge Structure (XANES) spectra at the Sr K-edge were simulated using the *ab-initio* FEFF8.4 code. ¹⁹ The full multiple scattering (FMS) and self-consistent-field (SCF) atomic potential calculation routines were found to be crucial to obtain good simulations. The Hedin–Lundqvist exchange potential ²⁰ was used. For the calculation of the SCF muffin-tin atomic potential, a cluster of size 7.2 Å (113 atoms) was used. A cluster of 300 atoms was used for the FMS calculation. The theoretical XANES spectra were shifted in order to match the experimental edge energy.

Standard procedures were followed to extract the EXAFS signal, ²¹ i.e., subtraction of the pre-edge background, normalization, alignment and energy calibration, and spline modeling of the post-edge atomic background. The EXAFS signal was Fourier filtered to exclude contributions of noise and coordination shells above the third one. The extraction of the EXAFS signal and the structural refinements were performed using the ESTRA-FITEXA software package.²² Minimizations were conducted in the wave vector space (k) to avoid possible artifacts introduced by Fourier filtering. The number of free parameters in the fits never exceeded the number of independent points $(N_{ind} = (2\Delta k \cdot \Delta R)/\pi + 1)$. The amplitude and phase photoelectron functions were calculated ab-initio using the FEFF8 code. 19 The crystallographic clusters centered on the absorption specie (Sr) used for FEFF calculations were obtained from the structure refined from neutron diffraction data, as described in the following.

E. Magnetization measurements

DC magnetization measurements were performed using two superconducting quantum interference devices (SQUID, Quantum Design, San Diego, USA) magnetometers at low temperature (5 K < T < 350 K) under different applied magnetic fields between 1 kOe and 140 kOe, at warming/cooling rates of 2 K/min. The ac-susceptibility was measured using a Physical Property Measurement System (Quantum Design, San Diego, USA) with frequencies ranging from 1 Hz to 10 kHz with H_{ac} = 10 Oe and H_{dc} = 0 Oe. For all the magnetization experiments, small fragments of the sintered ceramic pellets were used.

III. RESULTS

A. Room-temperature structure

Rietveld refinements of the NPD room-temperature data of samples Sr2% and Sr5% are presented in Fig. 1. Structural data obtained from the refinement are summarized in Table I. The refinements return a structural model in the $Pbn2_1$ space group, in line with that reported for the parent compound.⁷ Additional trials, explicitly including Sr at the nominal Ca site and/or at the Ba site, did not improve the results significantly. No unexplained reflections are observed in these samples, indicating that Sr has not been segregated in a secondary phase within the limits of detection of NPD for these samples. The occupation of oxygen sites refined to full occupancy within the uncertainty limits, in agreement

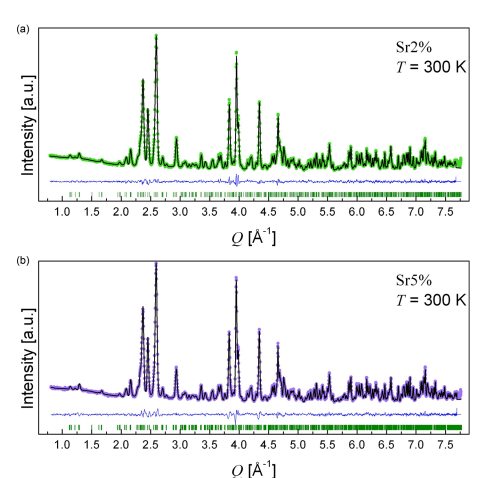


FIG. 1. Rietveld refinement for samples Sr2% and Sr5% from NPD data collected at room temperature in the high-resolution diffractometer D2B at ILL. Vertical bars at the bottom indicate Bragg reflections from the orthombic *Pbn*2₁-phase. The line at the bottom corresponds to the difference between the experimental and calculated patterns.

with the stoichiometric formula expected from the synthesis protocol used in preparing the samples.

The most relevant bond-lengths are summarized in Tables II and III for samples Sr2% and Sr5%, together with the results of bond-valence-sum (BVS) calculations

performed using the KDist software.²³ Negligible differences in the interatomic distances are observed between samples Sr2% and Sr5%. It is interesting to compare these data with the interatomic distances in the parent compound CaBaCo₄O₇. The room-temperature structure for

TABLE I. Structural parameters for samples $Ca_{0.98}Sr_{0.02}BaCo_4O_7$ (x=0.02) and $Ca_{0.95}Sr_{0.05}BaCo_4O_7$ (x=0.05) at room temperature. Data collected in the D2B diffractometer at ILL. Fractional coordinates correspond to the space group $Pbn2_1$ with all atoms in the Wyckoff position 4a. Isotropic thermal parameters (B_{150}), lattice constants, and reliability factors for the refinements are also indicated.

x = 0.02					x = 0.05			
Atom	х	У	Z	$B_{iso} [\mathring{\mathrm{A}}^2]$	х	у	Z	B_{iso} [Å ²]
Ca1/Sr1	0.0005(10)	0.6700(5)	0.8737(5)	0.51(6)	0.0029(11)	0.6700(5)	0.8730(5)	0.84(9)
Ba1	0.0009(6)	0.6667	0.5000	0.60(5)	0.0014(7)	0.6667	0.5000	0.46(7)
Co1	0.0021(16)	-0.0046(13)	0.9371(8)	0.52(12)	0.0011(17)	-0.0046(13)	0.9371(9)	0.37(11)
Co2	-0.0005(18)	0.1714(8)	0.6882(12)	0.66(6)	-0.0047(19)	0.1714(8)	0.6898(12)	0.68(6)
Co3	0.2453(17)	0.0850(7)	0.1896(10)	0.66(6)	0.2454(19)	0.0850(7)	0.1885(11)	0.68(6)
Co4	0.2645(15)	0.9181(7)	0.6827(10)	0.66(6)	0.2631(17)	0.9181(7)	0.6798(10)	0.68(6)
O1	-0.0068(14)	-0.0010(6)	0.2530(6)	0.52(5)	-0.0075(14)	-0.0010(6)	0.2538(7)	0.57(5)
O2	-0.0007(9)	0.4935(4)	0.2303(6)	0.73(6)	-0.0005(10)	0.4935(4)	0.2294(6)	0.68(6)
O3	0.7842(11)	0.2589(5)	0.7796(8)	1.59(10)	0.7854(11)	0.2589(5)	0.7795(8)	1.50(11)
O4	0.7327(9)	0.7456(4)	0.2141(7)	0.70(9)	0.7333(9)	0.7456(4)	0.2133(7)	0.73(9)
O5	-0.0555(5)	0.1528(4)	0.5000(7)	1.06(6)	-0.0558(6)	0.1528(4)	0.4991(7)	1.24(7)
O6	0.2127(6)	0.1016(3)	0.0000(9)	1.03(6)	0.2123(7)	0.1016(3)	0.0000(10)	1.25(6)
O7	0.2609(6)	0.9465(3)	0.4997(11)	1.00(5)	0.2605(6)	0.9465(3)	0.5001(11)	0.97(5)
a [Å]	6.2877(1)				6.2893(1)			
b [Å]	11.0078(3)				11.0136(2)			
c [Å]	10.1914(2)				10.1917(2)			
χ^2	2.39				3.18			
R_B	3.02			3.49				

TABLE II. Cobalt—oxygen bond lengths determined at room-temperature from the Rietveld refinement of D2B data, for samples $Ca_{0.98}Sr_{0.02}BaCo_4O_7$ and $Ca_{0.95}Sr_{0.05}BaCo_4O_7$. The bond-valence-sums (BVS), indicative of the cobalt oxidation states, are also shown.

	x = 0.02			x = 0.05			
Со-О	Bond-length [Å]	Average value [Å]	BVS	Bond-length [Å]	Average value [Å]	BVS	
Co1–O5	1.785(14)	1.856(12)	2.66	1.798(14)	1.856(13)	2.63	
Co1-O1	1.877(10)			1.869(12)			
Co1-O6	1.878(13)			1.882(13)			
Co1-O7	1.884(12)			1.874(12)			
Co2-O3	1.904(13)	1.947(13)	2.02	1.867(13)	1.952(13)	1.99	
Co2-O4	1.934(12)			1.952(13)			
Co2-O5	1.960(14)			1.981(14)			
Co2-O1	1.989(11)			2.008(11)			
Co3-O2	1.941(11)	1.953(12)	1.99	1.937(13)	1.954(13)	1.97	
Co3-O6	1.952(14)			1.943(15)			
Co3-O1	1.957(13)			1.957(14)			
Co3-O3	1.962(10)			1.980(12)			
Co4-O2	1.834(11)	1.890(12)	2.39	1.859(11)	1.884(12)	2.41	
Co4-O4	1.841(9)			1.817(9)			
Co4-O7	1.891(15)			1.861(15)			
Co4-O1	1.993(12)			2.000(13)			

CaBaCo₄O₇ was only reported by Caignaert *et al.*⁶ Using their atomic positions, the calculated bond-lengths are slightly different to ours. Other reports of the low-temperature structure of CaBaCo₄O₇ (Ref. 7) and the room-temperature structure of Zn-doped CaBaCo_{4-x}Zn_xO₇ (Ref. 24) are, on the other hand, in full agreement with our

TABLE III. Calcium and barium nearest neighbor's bond-lengths determined at room-temperature from the Rietveld refinement of D2B data, for samples $Ca_{0.98}Sr_{0.02}BaCo_4O_7$ and $Ca_{0.95}Sr_{0.05}BaCo_4O_7$. The bond-valence-sums (BVS), indicative of the cation's oxidation state, are also shown.

	X =	= 0.02		x = 0.05			
М-О	Bond-length [Å]	Average value [Å]	BVS	Bond-length [Å]	Average value [Å]	BVS	
Ca1-O3	2.254(9)	2.335(8)	2.07	2.263(10)	2.338(9)	2.04	
Ca1-O2	2.318(7)			2.328(8)			
Ca1-O6	2.340(8)			2.339(9)			
Ca1-O7	2.359(9)			2.392(9)			
Ca1-O5	2.362(8)			2.336(8)			
Ca1-O4	2.375(8)			2.368(9)			
Ba1-O3	2.749(8)	3.150(5)	1.59	2.746(8)	3.151(7)	1.55	
Ba1-O4	2.799(6)			2.802(7)			
Ba1-O5	2.799(3)			2.805(6)			
Ba1-O7	2.852(3)			2.831(6)			
Ba1-O6	2.880(3)			2.893(6)			
Ba1-O2	2.936(6)			2.914(6)			
Ba1-O2	3.345(6)			3.350(6)			
Ba1-O6	3.462(3)			3.461(6)			
Ba1-O4	3.474(7)			3.483(7)			
Ba1-O7	3.490(3)			3.506(6)			
Ba1-O5	3.496(3)			3.489(6)			
Ba1-O3	3.513(8)			3.527(8)			

results. The discrepancy with the reported room-temperature structure for $CaBaCo_4O_7$ in Ref. 6 may be due to the fact that the latter is based on laboratory XRD data, whereas NPD (used here as well as in Refs. 7 and 24) yields more reliable positions for the oxygen atoms. Based on these considerations, we propose that the structural model obtained in this work for Sr-doped samples is better for the description of $CaBaCo_4O_7$ at room-temperature.

The interatomic distances and BVS reported in Tables II and III also confirm that the Sr-substitution does not disrupt the scheme of charge order between Co^{2+} at Co_{2} and Co_{3} sites, and the hybridized $\text{Co}_{3}^{3+}/\text{Co}_{2}^{2+}(3d^{7}\underline{\textbf{L}})$ at sites Co1 and Co4 proposed for the parent compound.

The effects of the Sr substitution over the lattice parameters at room temperature are presented in Fig. 2, as well as on the volume of the cell normalized per unit formula. An anisotropic distortion is observed with the increase in Sr-substitution, while the c parameter remains unchanged (within experimental uncertainty). Most importantly, there is an increase in the unit cell volume. This is already pointing that Sr is substituting for Ca, a smaller cation. It is interesting to compare the behavior of our samples with those of Ca(Ba_{1-x}Sr_x)Co₄O₇ reported recently by Seikh *et al.*²⁵ The latter, in contrast to ours, shows a decrease in the unit cell volume with the Sr-for-Ba substitution, which is consistent

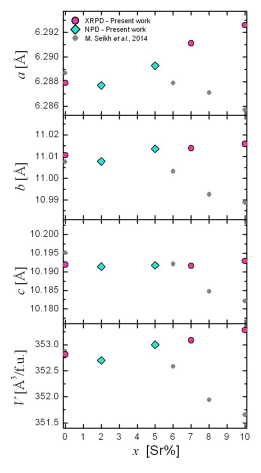


FIG. 2. Evolution of the room-temperature lattice parameters and volume per unit formula in the orthorhombic cell as a function of the Sr substitution (x) in $(Ca_{1-x}Sr_x)BaCo_4O_7$. Grey circles represent data for the $Ca(Ba_{1-x}Sr_x)Co_4O_7$ after Ref. 25.

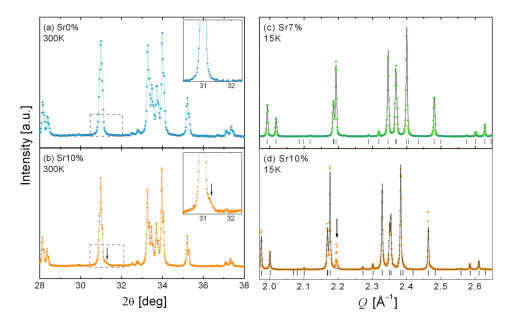


FIG. 3. (a) and (b) Room temperature laboratory XRPD data for sample Sr0% (a) and Sr10% (b) using Cu K- α radiation. The insets highlight the region where a reflection from a segregated phase appears for Sr10%, indicated with an arrow. (c) and (d) Synchrotron XRPD data at 15K for sample Sr7% (c) and Sr10% (d) collected at the LNLS synchrotron in Brazil. The solid line represents a Le Bail fit of the majority phase in the Pbn21 space group. The arrow indicates the segregated phase reflection. The LNLS data are plotted as a function of the scattering vector Q to allow comparison of different wavelengths.

with the smaller size of Sr⁺² with respect to Ba⁺² (grey circles in Fig. 2). Apparently, the structure is flexible enough to admit both substitutions, which makes it very important to ensure the correct substitution site before drawing any conclusions over the physical properties. To that end, a XAS study was conducted at the Sr K-edge and is presented below.

B. Solubility limit

The presence of a segregated phase in the Sr10% sample $(x_{Sr} = 0.10)$ was first suggested by our XRPD measurements. Diffractograms at room-temperature for the Sr0% and Sr10% samples are shown in Figs. 3(a) and 3(b), respectively, collected using a laboratory X-ray diffractometer. A shoulder to the right of the (0 2 3)–(1 1 3) reflections located at 31° suggests the presence of an additional phase in Sr10% (Fig. 3(b)). This was confirmed in a synchrotron XRPD experiment performed at the LNLS in Brazil with a much better resolution.²⁶ In Figs. 3(c) and 3(d), we show the same Q-range in the LNLS diffractograms collected at 15 K. The presence of a segregated phase in the Sr10% sample is clearly visible (indicated with an arrow in Fig. 3(d)), while it is absent in the Sr7% sample, confirming that the solubility limit for Sr at the Ca site lies between 7 and 10 at. % Srdoping. Although the remaining reflections from the segregated phase are quite weak (not shown in the figure), the phase may be assigned to a perovskite belonging to the family $Sr_xBa_{1-x}CoO_{2.5}$, most likely with $x \sim 0.2$.^{27,2}

C. Sr substitution site

The local structure around Sr was explored by XAS. In Fig. 4, the room temperature crystal structure of CaBaCo₄O₇ is sketched. The tetrahedral coordination of Co atoms is shown in panel (a), whereas the coordination of Ca and Ba atoms is highlighted in panel (b). Calcium and Ba cations have very different first coordination shells: Ba atoms are coordinated with 12 oxygen neighbors at an average distance of 3.15 Å, and Ca atoms are coordinated with six oxygen

atoms at an average distance of 2.34 Å, as summarized in Table III. Since the resolution of our XAS data is about 0.05 Å ($\delta R = 0.2 \cdot \pi/k_{max}$, where $k_{max} \sim 13$ Å⁻¹), the first coordination shell of Sr at Ba or Ca sites should differ enough to be distinguished by XAS. For this experiment, we used the Sr7% sample to obtain the best signal from the Sr K-edge but without reaching the solubility limit.

The Sr K-edge XANES spectra corresponding to the two possible substitution sites, Sr@Ca and Sr@Ba, were

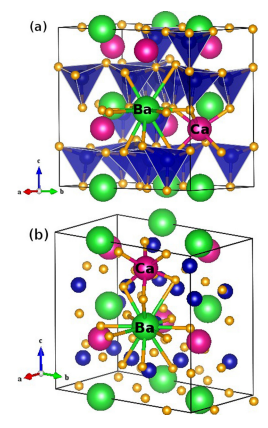


FIG. 4. Crystal structure of $CaBaCo_4O_7$ at room temperature as refined from neutron diffraction data. The oxygen coordination polyhedra around Co are represented in blue (a). (b) First neighbor's shell of Ca and Ba atoms.

modeled using FEFF8.4 calculations, starting from our NPD crystallographic data for the Sr5% sample. The simulated XANES spectra are shown in Fig. 5, together with the experimental spectrum of sample Sr7% which corresponds to the merge of the three collected scans aligned and calibrated to the Sr K-edge (16 105 eV). Although we could not fully reproduce the spectral features, it is evident that there is a better match between the experimental data and the XANES signal calculated for Sr atoms located at the Ca site and that all the features of the experimental spectrum are also qualitative present in the simulated Sr@Ca XANES signal. Therefore, we consider that the latter is the most suitable model to describe our sample, as the XANES signal for the Sr@Ba model shows a much worse agreement.

For the EXAFS analysis, *ab-initio* amplitude and phase scattering functions were generated using FEFF8 (Ref. 19) for both Sr@Ba and Sr@Ca models. To account for the lower resolution of XAS data, the first three coordination shells around Sr (Sr-O, Sr-Co and Sr-Ba/Ca) were averaged from the NPD data and are indicated in Table IV. Shell fitting was performed using only the above single-scattering paths. Coordination numbers were kept fixed to the crystallographic structures, while atomic distances, Debye-Waller factors, and the "global" parameters (the many body amplitude reduction factor, S_0^2 , and the energy shift, Δ E) were left free to vary during minimizations. The EXAFS signal and fit curve using the Sr@Ca model, together with their Fourier transforms, are shown in Fig. 6.

The structural parameters extracted from the EXAFS fit at the Sr K-edge are reported in Table IV and compared with the structures calculated from the Sr@Ca and Sr@Ba models based on averaged neutron diffraction data. The comparison reported in Table IV shows that the experimental data match well with the model where Sr substitutes at the Ca site (Sr@Ca). The first coordination shell (Sr-O) average distance is larger than in the NPD-model of Sr@Ca, in agreement with the larger size of Sr with respect to Ca. The second coordination shell (Sr-Co) average distance matches

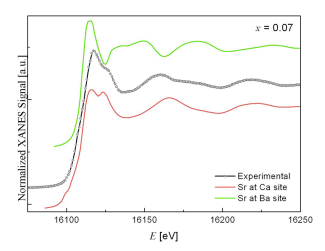


FIG. 5. Simulated Sr K-edge XANES spectra for the Sr@Ca and Sr@Ba models. The experimental spectrum for sample Sr7% (nominal $(Ca_{0.93}Sr_{0.07})$ Ba Co_4O_7) is shown for comparison using symbols. The spectra are vertically shifted for clarity.

TABLE IV. Comparison between the crystallographic structure around the Sr atom of the Sr@Ca and Sr@Ba models calculated from neutron diffraction data and the results of the EXAFS refinements for sample Ca(Ba_{0.93}Sr_{0.07})Co₄O₇ (Sr7%) (CN, coordination numbers; R, atomic distances; σ^2 , Debye-Waller factors; S₀, amplitude reduction factor; ΔE , energy shift; χ^2_ν , reduced chi-square). The numbers between parentheses represent the error on the last digit.

	Mo	EXAFS data		
Photoelectron path	Sr@Ca	Sr@Ba	Sample Sr7%	
	CN	CN	CN	
	R (Å)	R (Å)	R (Å)	
			$\sigma^2 (\mathring{\mathrm{A}}^2)$	
Sr-O	6	12	6	
	2.338	3.150	2.46(1)	
			0.009(1)	
Sr-Co	12	12	12	
	3.688	3.687	3.71(1)	
			0.018(2)	
Sr-Ba	4		4	
	3.856		3.90(5)	
			0.018(6)	
Sr-Ca		4		
		3.856		
			$S_0^2 = 0.9(1)$	
			$\Delta E = 1.2(9)$	
			$\chi^2_{\nu}=4.3$	

with both Sr@Ca and Sr@Ba models, in agreement with the fact that Sr "sees" the same cobalt environment from both Ca and Ba sites. The refinement of the third coordination shell would be conclusive in revealing the substitution site, because Sr would only "see" Ba if located at the Ca site (Sr@Ca) and only Ca if substituting at the Ba site (Sr@Ba). Both Sr-Ba and Sr-Ca contributions were used to fit the third coordination shell, but, although the match with experimental data improves when using Sr-Ba scattering paths, the results are not conclusive. In fact, even if, in principle, XAS would allow distinguishing between Sr-Ba and Sr-Ca contributions to the third shell, the correlation between parameters dramatically lowers its chemical sensitivity. On the other hand, all attempts to fit the data with the model Sr@Ba led to bad matches with the experimental data and to unphysical values of the S_0^2 parameter. Therefore, despite restrictions in analyzing the third shell by EXAFS, the complete XAS analysis allows to conclude that Sr sits at the Ca site in the structure, so we can be confident that the nominal composition is correct and the discussion on the effects of replacing Ca-by-Sr may rely on this result.

IV. MAGNETIC PROPERTIES

The effects of Sr-substitution on the magnetism of the CaBaCo₄O₇ system were studied using magnetometry techniques. For the lowest level of substitution studied here, corresponding to 2 at.% Sr-for-Ca, we observe a very similar magnetic behaviour as for the parent compound. The comparison between the Sr0% and Sr2% samples is presented in Fig. 7. It may be inferred from this figure that the Sr2% sample also undergoes a transition to a ferrimagnetic phase with

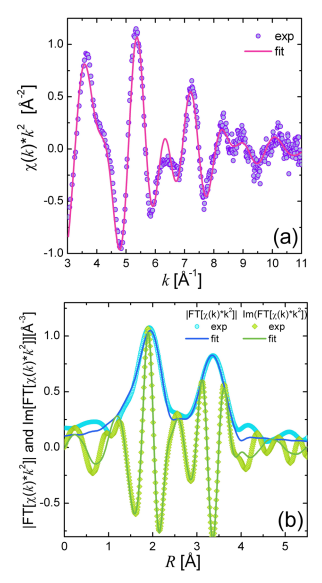


FIG. 6. (a) k^2 -weighted EXAFS signal (symbols) and fit (line) of sample Sr7% (nominal (Ca_{0.93}Sr_{0.07})BaCo₄O₇). (b) Modulus (|FT|) and imaginary parts (Im) of the Fourier transformed (1–4 Å Hanning window) EXAFS signal and fit reported in panel (a). Symbols represent experimental data and solid lines are the fit curves. Fourier transformed data are k^2 -weighted and not corrected for the phase shift.

a similar saturation value of magnetization at the lowest temperature and with a T_C slightly lower than for the parent compound. The inset in Fig. 7 shows the ac-magnetization $(\gamma'(T))$ for the two samples under different frequencies. For the parent compound, we observe the expected curve for a stoichiometric sample, 15,24 with a unique sharp peak centered at $T_C = 60 \,\mathrm{K}$ which is frequency-independent. No further features at all are observed at lower temperatures, which have been reported to appear in cases where the stoichiometry is not perfect, being $\chi'(T)$ one of the most sensitive properties to the presence of new magnetic interactions introduced by doping and/or stoichiometry deviations.^{24,29} Despite the similarities with the parent compound, the $\gamma'(T)$ curve for the Sr2% sample reveals some indications of additional magnetic interactions in the system. To begin with, the peak corresponding to the ferrimagnetic transition has become less sharp and has shifted to a lower temperature. In

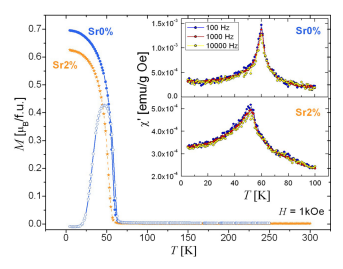


FIG. 7. DC-magnetization as a function of temperature under an applied field of $1000\,\mathrm{Oe}$ for $\mathrm{CaBaCo_4O_7}$ and $\mathrm{Ca_{0.98}Sr_{0.02}BaCo_4O_7}$. Open symbols correspond to a zero-field cooled (ZFC) magnetization curve collected on warming, whereas solid symbols correspond to the magnetization collected on cooling under an applied field (FCC). The insets show the temperature dependence of the real part of the ac-susceptibility under an applied field $H_{ac}=10\,\mathrm{Oe}$ and $H_{dc}=0\,\mathrm{Oe}$ measured at different frequencies.

addition, a small kink has appeared at ${\sim}80$ K, which has been repeatedly reported for several kinds of substitutions in the CaBaCo₄O₇ system^{13,24,30,31}

More insight into the origin of these features may be derived from the evolution of the magnetization curves upon further substitution with Sr. In Fig. 8, we show the magnetization as a function of temperature for the whole series of Sr-substituted samples with $0.02 \le x \le 0.10$. Note that the data are presented in a logarithmic scale, to stress the dramatic decrease in the net magnetization upon Sr substitution. It is immediately clear that the feature at ~ 80 K is systematic, and gradually evolves into a "plateau" of the magnetization in a temperature range whose lower limit is x-dependent. This "plateau" suggests the existence of antiferromagnetic (AFM)-like order in competition with the

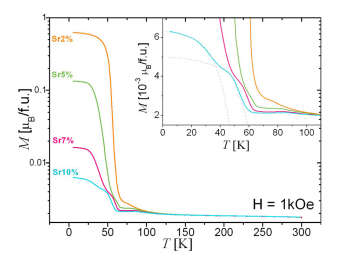


FIG. 8. Magnetization as a function of temperature under an applied field of 1000 Oe for $Ca_{1-x}Sr_xBaCo_4O_7$ with $x=0.02,\ 0.05,\ 0.07,\ and\ 0.10$. Data were collected with a FCC protocol. The inset shows a magnification of the low temperature data in a linear scale. The dashed lines on the Sr10% curve are only guides to the eye.

ferrimagnetic phase, preventing and/or delaying the sharp raise in magnetization at the T_C .

The similar raise in magnetization below 60 K for the Sr2% sample suggests that the ferrimagnetic phase characteristic of the parent compound is preserved. On the other hand, the curve for the Sr5% sample already indicates that the order has changed dramatically at low temperature, and further substitution enhances this idea. Moreover, the inset in Fig. 8 shows a magnification of the low-temperature region to highlight what we believe are two different transitions characterized by different T_C values and saturation magnetizations. We have outlined this feature with dashed lines for the Sr10% sample in the inset. Although the x = 0.10 sample has a small amount of a segregated perovskite phase, its magnetization behaviour is completely in line with the systematics of Sr-addition, and the evidence of two different transitions at temperatures below the "plateau" is also observed in samples with x = 0.07 and 0.05 in which there are no segregated phases at all. We would finally like to stress the similarities yet differences with the behavior reported in Ref. 29 for the Sr-for-Ba substitution. The presence of the magnetization "plateau" is clearly a common feature, however below 60 K, the curves do not look similar. The authors propose a scenario of competition between the ferrimagnetic phase of the parent compound and a frustrated magnetic phase with a frequency dependent ac-susceptibility which is not at all compatible with our data. The present magnetization curves at low temperature are rather more similar to those reported for Li-substituted samples³¹ which were reported as a Li-for-Co substitution. It is evident that the variety of transitions suggested by the dc- and acmagnetization curves deserves a more profound study, which is now in progress and will be reported separately. It is also clear that a deep discussion on the reasons for the impressive effects of substitution must rely on the correct identification of substitution sites in the structure, which is the main objective of the present work.

V. CONCLUSIONS

The Ca-based "114" cobaltates have opened up exciting possibilities for the study of magnetoelectricity, magnetic frustrated states, ferrotoroidicity, etc. 10-12 Cation substitution has proven to trigger even more interesting questions regarding the phase competition in the system, as it deeply affects their magnetism, charge order, and transport properties. The accurate knowledge of the substitution site usually remains a standing issue, despite being of crucial importance to understand the magnetic phenomena.

In this work, we have investigated the cationic arrangements in Sr-doped Ca-"114" cobaltates combining long-range structural probes, such as neutron and x-ray diffraction, with short-range chemical selective tools, such as EXAFS and XANES simulations. By applying this multitechnique approach, we demonstrated that Sr can substitute at the Ca site with a solubility limit lying around $x \simeq 0.08$. However, in the light of the results by other groups, ²⁵ a Sr-for-Ba substitution might also be possible in this system. It is therefore important to correctly characterize the samples

before drawing conclusions on the effects of doping on the magnetic, structural, and other properties. The present results support our nominal doping site and constitute the base to analyze the impressive effects of the Sr-for-Ca substitution on the magnetic properties. We observe that a small amount of Sr addition ($x \simeq 0.02$) preserves the ferrimagnetic phase of the parent compound although there are indications that other magnetic interactions start to compete in the system. This is firmly established by further increasing the Sr-for-Ca substitution, which results in a decrease in magnetization of two orders of magnitude for a 10 at. % substitution. An AFM state in a temperature region around $60 \,\mathrm{K} \lesssim T \lesssim 80 \,\mathrm{K}$ is shown to get stabilized with Sr content. At lower temperature, the ferrimagnetic phase of the parent compound seems to be replaced by other types of order involving even a third transition which needs to be further studied.

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