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Band structures, magnetic properties, and enhanced magnetoresistance in the high pressure phase of Gd and Y doped two-dimensional perovskite Sr2CoO4 compounds

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Abstract
The authors present their studies on the band structures and on the magnetic and magnetotransport properties of the high pressure phase of Sr$_{2-x}$RE$_x$CoO$_4$ [rare earth (RE)=Gd and Y, x=0.1-0.5] compounds which were synthesized by a high pressure and high temperature technique. The authors found that as x increases, the magnetoresistance $\frac{(\rho_H - \rho_0)}{\rho_0}$ increases up to -17% at 5 K and 7 T, which is 2.5 times higher than that for undoped Sr$_2$CoO$_4$, although the ferromagnetic transition drops from 255 to 200 K for the Gd doping with x=0.3. The saturation moments at low temperature are significantly enhanced for the Gd doped Sr$_2$CoO$_4$. Observation of a close correlation between resistance and field revealed a strong spin-dependent tunneling magnetoresistance. First-principles band structure calculations indicate that high spin polarization is present for both undoped and doped compounds.

Keywords
Band, structures, magnetic, properties, enhanced, magnetoresistance, high, pressure, phase, doped, two, dimensional, perovskite, Sr2CoO4, compounds

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Band structures, magnetic properties, and enhanced magnetoresistance in the high pressure phase of Gd and Y doped two-dimensional perovskite Sr$_2$CoO$_4$ compounds

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The authors present their studies on the band structures and on the magnetic and magnetotransport properties of the high pressure phase of Sr$_{2-x}$RE$_x$CoO$_4$ [rare earth (RE)=Gd and Y, $x=0.1-0.5$] compounds which were synthesized by a high pressure and high temperature technique. The authors found that as $x$ increases, the magnetoresistance $(\Delta \rho / \rho_0)$ increases up to $-17\%$ at 5 K and 7 T, which is 2.5 times higher than that for undoped Sr$_2$CoO$_4$, although the ferromagnetic transition drops from 255 to 200 K for the Gd doping with $x=0.3$. The saturation moments at low temperature are significantly enhanced for the Gd doped Sr$_2$CoO$_4$. Observation of a close correlation between resistance and field revealed a strong spin-dependent tunneling magnetoresistance. First-principles band structure calculations indicate that high spin polarization is present for both undoped and doped compounds. © 2007 American Institute of Physics.

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Finding novel ferromagnetic materials showing negative magnetoresistance and studies of polarized spin-dependent tunneling magnetoresistance are hot topics within spintronics. So far, several magnetoresistance (MR) oxide systems have been found and extensively investigated. (1) The perovskite ferromagnetic manganites in the form of $\text{RE}_n\text{A}_m\text{MnO}_3$, with RE standing for a rare earth and A for a divalent ion such as Ca, Sr, or Ba, exhibit the well known type of colossal magnetoresistance in fields of several tesla. (2) Pyrochlore-type $\text{T}_2\text{Mn}_2\text{O}_7$ shows large MR near the Curie temperature of 140 K. The double-perovskite ferromagnets, Sr$_2$FeMoO$_6$ and Sr$_2$FeReO$_6$, exhibit large low-field MR below high Curie temperatures $T_C$ of about 400 K. $\text{Fe}_2\text{O}_4$ is a good candidate for room temperature device applications because it has a very high $T_C$ of 858 K. Analogous to the perovskite manganites, giant magnetoresistance is also observed in the cobalt based perovskite compounds $\text{RE}_1\text{A}_1\text{CoO}_3$, as well as in oxygen deficient LnBa$_2$Co$_2$O$_{6-x}$ (Ln=Eu and Gd) compounds. When the dimensionality is reduced from three to two dimensions, the MnO$_2$ or CoO$_2$ network is confined to the $ab$ plane. Spin-related property changes are also expected. Layered perovskite manganites, defined as the Ruddlesden-Popper family of compounds (RE)$_n$(A)$_m\text{MnO}_{3n+1}$, consist of $n$ layers of MnO$_6$ octahedra separated by $n$ layers of REO$_2$.

Very recently, metallic ferromagnetism with a Curie temperature $T_C$ of 250–255 K has been discovered in the K$_2\text{NiF}_4$-type two-dimensional layered perovskite Sr$_2$CoO$_4$ in the form of both single crystalline films fabricated by pulsed laser deposition (PLD) and bulks produced by a high pressure and high temperature technique. A metal insulator transition with large negative MR values was observed in the vicinity of $T_C$. The MR increases at low temperatures and reaches a maximum at the coercive field $H_c$. $\text{Y}^{3+}$ doping into Sr$^{2+}$ made the Sr$_2$CO$_4$ gradually change from a ferromagnetic metal to an antiferromagnetic semiconductor. Band structure calculations by Lee and Pickett revealed that the Sr$_2$CoO$_4$ can be either a ferromagnetic metal for the thin film samples or a half metal for the high pressure phase samples. They pointed out that the newly discovered Sr$_2$CoO$_4$ in the form of bulks fabricated by the high pressure technique and thin films made by PLD introduces new transition metal oxide physics and may be useful in spin electronics devices. In addition, it should be mentioned that the MR is only about $-7.5\%$ at 7 T and 5 K for the high pressure phase Sr$_2$CoO$_4$ samples. Using the high pressure method it would be interesting to see how the physical properties such as the MR values and Curie temperatures as well as the electronic structures of the Sr$_2$CoO$_4$ can be changed or improved by doping with other rare earth elements, in particular, with those having large magnetic moments or no magnetic moments. In this letter, we present our observations on the magnetic properties and the improvement of the negative magnetoresistance, as well as the electron band structures of the Gd or Y doped Sr$_2$CoO$_4$.

Polycrystalline samples of Sr$_{2-x}$RE$_x$CoO$_4$ (RE=Y or Gd, $x=0,0.1,0.3,0.5$) were prepared by a high pressure and high temperature technique. Fine and pure powders of SrO$_2$, Co, and Y$_2$O$_3$ or Gd$_2$O$_3$ were well mixed in the ratio of Sr:RE:Co=2-$x$:$x$:1. The samples were compressed at...
6 GPa in a high pressure apparatus and then heated at 1000–1350 °C for 1–3 h and quenched to room temperature, with the quenching followed by a release of pressure. Structure refinements were carried out by the Rietveld method. Magnetic and magnetotransport properties were investigated using commercial Quantum Design magnetic and physical properties measurement systems between 2 and 330 K in fields up to 7 T.

The temperature dependence of the magnetization reveals that the ferromagnetic transition of Sr\(_2\)CoO\(_4\) decreases from 255 to 230 K for Gd doping with \(x=0.1\) and to 200 K for both Gd and Y with \(x=0.3\), as shown in Fig. 1. An upturn at low temperatures in the Gd doped sample is believed to come from the contribution of paramagnetic Gd\(^{3+}\) ions. The temperature dependence of the resistivity of those samples is shown in Fig. 1(a). Among these samples, the Sr\(_2\)CoO\(_4\) has the lowest resistivity (0.04 \(\Omega\) cm) over the wide range of temperatures measured. The resistivity for samples with \(x=0.1\) and 0.3 for both Gd and Y doping showed a very gradual increase in resistivity with decreasing temperature and does not increase at low temperatures. This indicates that the resistance is dominated by the carrier scattering at grain boundaries, as is widely seen for polycrystalline samples.

The doping effect on the MR is clearly revealed in Fig. 2. The MR values at 50 K increase from −5% up to −8.5%, as the Y doping level is raised from 0.1 to 0.5. The MR values increase to −13% and −14% for the \(x=0.1\) and 0.3 samples at 5 K and 7 T. The Gd doping has a similar effect on the MR to that of the Y doping. For Sr\(_{1.7}\)Gd\(_{0.3}\)CoO\(_4\), the MR is −4% at 100 K and 7 T and increases to −12% at 20 K and 7 T, as shown in Fig. 3. It should be mentioned that the MR is only about 7.5% at 7 T and 5 K for undoped Sr\(_2\)CoO\(_4\). The MR is much enhanced by both Y and Gd dopings at 5 K. As shown in Fig. 3, the MR values increased to −14% and −17% for the Sr\(_{1.7}\)Y\(_{0.3}\)CoO\(_4\) and Sr\(_{1.7}\)Gd\(_{0.3}\)CoO\(_4\) samples, respectively. With decreasing temperatures, a field-hysteretic MR corresponding to the magnetic hysteresis loop gradually appeared and became much more pronounced at 5 K, as shown in Fig. 3 (lower panel). It can be seen that the maximum field-hysteric MR occurred at fields of about 2.5, 0.5, and 0.3 T for pure Sr\(_2\)CoO\(_4\), Sr\(_{1.7}\)Y\(_{0.3}\)CoO\(_4\), and Sr\(_{1.7}\)Gd\(_{0.3}\)CoO\(_4\), respectively, coinciding with the magnetization process with domain rotation at coercive fields of 2.5, 0.5, and 0.3 T, respectively.

Such a close correlation between the MR and magnetic domain rotation can be well explained in terms of spin-dependent tunneling MR at grain boundaries, which has been well established for granular manganites and other MR oxides. Under the application of magnetic field, the hopping of spin-polarized electrons between grains (domains) is pre-
dominantly affected by the relative angle of the magnetization direction and controlled by the external field through the domain-rotation process. Therefore, the observed field-hysteretic MR is ascribed to the field suppression of the spin-dependent scattering at grain (domain) boundaries. It should be pointed out that the enhanced MR value of $-17\%$ in Sr$_{1.7}$Gd$_{0.3}$CoO$_4$ is almost two times smaller than that for pure $a$-axis oriented Sr$_2$CoO$_4$ single crystal thin films, but it is two times greater than that in Sr$_2$CoO$_4$ polycrystalline bulks.\cite{12,13} It is obvious that the grain boundary tunneling between oriented grains is much greater than that between the randomly oriented grains. Therefore, it is expected that the MR values will be further enhanced in textured thin films of either Sr$_{1.7}$Gd$_{0.3}$CoO$_4$ or Sr$_{1.7}$Gd$_{0.3}$CoO$_4$, with much reduced grain boundary scattering.

First-principles band structure calculations were performed using a density functional theory.\cite{15} We adopted the standard generalized gradient approximation, with the Perdew-Burke-Ernzerhof functional. Lattice parameters and atomic positions obtained from the Rietveld refinement for pure and doped samples prepared under 6 GPa were used as the structural parameters for the band structure calculations. A 14 atom supercell of one Sr$_2$CoO$_4$ unit cell containing one Y or Gd atom substituted at the Sr site was used for the calculation. The Y or Gd concentration is thus 25 at. %. A schematic of the crystal structure of the Sr$_2$CoO$_4$ is shown in Fig. 4. A $7 \times 7 \times 2$ mesh was used for Brillouin sampling with a cutoff energy of 340 eV. A calculation for pure Sr$_2$CoO$_4$ was also performed as a comparison. Figure 5 shows the total density of states with majority “up” and minority “down” spins. For the undoped and doped Sr$_2$CoO$_4$, both the spin up and spin down states are partially occupied around the Fermi energy set at zero.

![Fig. 4.](Image) Crystal structure and distribution of spin densities (blue dots) of the pure (left) and Gd (right) doped Sr$_2$CoO$_4$.

![Fig. 5.](Image) Total density of states of pure, Gd, and Y doped Sr$_2$CoO$_4$, with the Fermi energy set at zero.

doped Sr$_2$CoO$_4$. It can be seen that the significantly enhanced magnetization at 5 K in the Gd doped Sr$_2$CoO$_4$, as shown in Fig. 3, mainly comes from Gd\cite{16} as well as from the oxygen and cobalt ions. The observation of enhanced magnetoresistance in the Y and Gd doped two-dimensional layer structured Sr$_2$CoO$_4$ suggests their potential applications in spintronics and in exploring novel magnetoresistance/spintronics materials in lower dimensional structured systems. Further enhancements in MR values and the Curie temperature are most likely present in other rare earth doped Sr$_2$CoO$_4$.

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