Effect of Ti^{4+} substitution on structural, transport and magnetic properties of $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_3$.

Vishwajeet Kulkarni[†], K. R. Priolkar[†][§], P R Sarode[†], Rajeev Rawat[‡], Alok Banerjee[‡], and S. Emura[¶]

† Department of Physics, Goa University, Taliegao Plateau, Goa 403 206 India ‡ UGC-DAE Centre for Scientific Research, University Campus, Khandwa Road, Indore 452 017 India

¶ Institute of Scientific and Industrial Research, Osaka University, Mihoga-oka 8-1, Ibaraki, Osaka 567-0047, Japan.

E-mail: krp@unigoa.ac.in

Abstract. La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO₃ ($0 \le x \le 0.20$) polycrystalline materials are prepared by employing lower annealing temperature compared to the temperatures reported for the materials of the same composition. The transport and magnetic properties of these materials are significantly different from those compounds prepared at higher annealing temperature. Samples with x < 0.10, show metal-insulator transition and those with $x \ge 0.10$ exhibit insulating behavior over the entire temperature range investigated. A gradual transition occurs from ferromagneticmetallic state to ferromagnetic-insulator state with increasing Ti substitution. Lattice parameters and bond lengths of Mn and its near neighbours however do not change appreciably with the dopant content x in these materials. It is shown that Ti⁴⁺ doping in the low temperature annealed samples is inhomogeneous resulting in isolated Mn rich regions that are connected by a variable range hopping polaron.

PACS numbers: 75.47.Lx; 75.47.Gk; 71.30.+h; 87.64.Fb

Submitted to: J. Phys.: Condens. Matter

§ Author to whom correspondence should be addressed.

1. Introduction

Colossal magnetoresistive (CMR) manganites are among the most studied materials in condensed matter physics [1, 2]. Physical properties of manganites are very sensitive to the method of preparation, the type of symmetry of a unit cell, A and B site cations, size effects, concentration of the substitute, hetrovalent substitution, non stoichiometry etc. Most of the CMR properties are explained in terms of Double-exchange(DE) and/or superexchange interactions, charge localization via Jahn-Teller distortion with polaron formation, phase separation and site-disorder. Yet there is no agreement on the correct theoretical description of CMR properties, due to the complexities involved. Lately, there has been a greater focus on the effect of the synthesis procedures on CMR properties of these materials. It is well known that CMR materials exhibit significantly different properties when they are prepared with different synthesis procedures, starting from the same initial composition. Various defects like cation vacancies, oxygen nonstoichiometry and cation site disorder develop in the crystal structure due the synthesis procedure and the environment used to prepare these materials. However, the effect of synthesis procedure on the microscopic as well as macroscopic properties of these materials is quite complex and needs to be investigated further to understand the physics involved.

Effect of oxygen non-stoichiometry on properties of CMR samples has also been studied in great detail [3, 4, 5, 6, 7]. For example, the defect chemistry of LaMnO_{3± δ} is unique [8]. For LaMnO_{3+ δ}, the structure is GdFeO₃-type for $0 \leq \delta < 0.10$ and rhombohedral for $0.10 \leq \delta < 0.3$ [9]. Although traditionally lanthanum manganite is considered as an anion excess compound (LaMnO_{3+ δ}), detailed investigations by employing high-resolution electron microscopy and other cognate techniques have revealed the presence of metal vacancies $(La_{1-\gamma}Mn_{1-\gamma}O_3)$ instead of interstitial oxygen ions [10, 11]. LaMnO₃ perovskites has been shown to tolerate a considerable portion of vacancies in the A site (La site) giving rise to compositions of the type $La_{1-\delta}MnO_3$ with the charge compensated by formation of Mn^{4+} [12]. The Mn^{4+} content in lanthanum manganites can be varied by altering the firing temperature and atmosphere. Oxygenrich samples exhibit ferromagnetic-metallic to paramagnetic-insulating transition due to the holes doped by cation vacancies. However, in deliberately prepared cation deficient material like $La_{1-x}MnO_{3+\delta}$, it has been argued that charge deficit due to vacancies on La sites would be preferably compensated by the formation of oxygen vacancies, rather than by oxidation of Mn^{3+} into Mn^{4+} ions. Recently, XRD and EXAFS studies [13] on vacancy doped $La_{1-x}MnO_{3+\delta}$ have shown that for La/Mn ratio below 0.9, there is phase segregation of $La_{0.9}MnO_3$ and parasitic Mn_3O_4 and T_C remains almost constant as that of $La_{0.9}MnO_3$.

Strontium doped LaMnO₃ series also exhibits both oxygen rich as well as oxygen deficient non-stoichiometry. Trukhanov et al[14, 15] have studied oxygen deficient $\text{La}_{1-x}\text{Sr}_x\text{MnO}_{3-\frac{x}{2}}$ compounds wherein ferromagnetic indirect superexchange is considered to be the cause of the magnetism. There are several reports [7, 11, 16, 17] on $\operatorname{La}_{1-x}\operatorname{Sr}_{x}\operatorname{MnO}_{3+\delta}$ compounds, wherein cation vacancies in oxygen rich materials are considered to be equally distributed on A and B sites, given by $v = \frac{\delta}{(3+\delta)}$ whereas Tofield and Scott [3] and Mitchell *et al* [18] have shown that the A-site vacancies are predominant. Mizusaki *et al* have discussed various models for cation deficiency in $\operatorname{La}_{1-x}\operatorname{A}_{x}\operatorname{MnO}_{3}$ [19] such as A-site cation substituting B-site and vice-versa. However, there appears to be very little literature about B-site vacancies [20, 21] in these materials. Interestingly, Nakamura [22] has shown that a fraction of Mn ions enter into A-site cation vacancies in $\operatorname{La}_{1-\Delta}\operatorname{MnO}_{3+\delta}$ and affects properties of the material substantially.

In spite of the extensive work carried out on these materials, there is no clear understanding and consensus on the microscopic picture of the disorder caused by vacancies or dopant atoms in these materials. Moreover, the effect of dopant ions replacing hetrovalent ions has not been studied so far. Such a disorder can have tremendous effect on the properties of these materials and is expected to give some interesting results. Ti doped CMR manganites offer such an opportunity. Ionic radius of Ti^{4+} ion is known to be in between those of Mn^{4+} and Mn^{3+} ionic radii and there exists a distinct possibility that a fraction of Ti⁴⁺ ions substitute for Mn³⁺ ions leading to oxygen non stoichiometry (cation deficiency) or such inhomogeneities. The available literature on Ti substitution in manganites indicates that Ti⁴⁺ ions substitute the isovalent Mn⁴⁺ ions in these materials [24, 25, 26, 27, 28, 29, 30, 31, 32, 33]. Recently however, it has been shown that at high doping levels Ti^{4+} ions occupy Mn^{3+} sites in addition to Mn^{4+} sites [34]. Effect of lower temperature annealing has also been studied recently [35]. It has been shown that by suitably modifying the preparation procedure, Ti⁴⁺ ions can substitute Mn ions randomly in $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_3$ forming inhomogeneous short range ordered ferromagnetic clusters. Here we report a detailed investigation of structural, magnetic, transport and spectroscopic properties of lower temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_3$ ($0 \le x \le 0.2$) using techniques like XRD, resistivity, AC and DC susceptibilities, XPS, Mn and La K-edge EXAFS and IR spectroscopy. Since annealing temperature of these materials is substantially lower than that reported in [27, 29], we term these as low-temperature annealed materials. For comparison, x = 0.10 sample has been prepared at the same annealing temperature as in [27] and is referred to as high-temperature annealed sample.

2. Experimental

Polycrystalline samples of $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_3$ with x = 0.0, 0.03, 0.05, 0.10 and 0.20 were synthesized by conventional solid-state reaction method. The powders of La_2O_3 , SrCO₃, freshly prepared MnCO₃ and TiO₂ with proper stoichiometry were mixed, ground manually using agate mortar and pestle and fired in air at 1000°C for about 15 hours, reground to calcine at 1100°C for 20 hours. Finally they were pressed into pellets and sintered in air at 1200°C for 20 hours. Grinding period was about half hour every time. The x = 0.10 sample was prepared also by firing the ingredients in stoichiometric proportions at 1000°C, 1200°C and annealing at 1450°C which closely

match with those reported in [27, 29]. In this case however the intermediate grinding was done manually with agate mortar and pestle for about 3 hours.

The samples were characterized by X-ray powder diffraction recorded on Siemens D5000 diffractometer at room temperature between $2\theta = 10^{\circ}$ to 70° with a step 0.05°. For AC susceptibility measurements, the samples were cooled under zero magnetic field and measurements were carried out in the warming run under the applied field of 0.936 Oe and AC frequency 133.33 Hz from liquid nitrogen temperature to 310 K on an in-house AC Susceptometer [36]. For the samples with x = 0.00 and x = 0.03, the susceptibility measurements were carried out up to 365 K and 330 K respectively. Standard four-probe technique was used to measure resistivity of the samples in the temperature interval 300 K to 80 K. Magnetoresistance (MR) measurements were performed down to 30 K using the standard four-probe geometry in transverse magnetic fields up to 5 T using OXFORD Spectralab 10 T superconducting magnet. Infrared measurements were performed on Shimadzu FTIR-8900 spectrophotometer at room temperature in transmission mode in the range of 350 cm^{-1} to 1000 cm^{-1} . The samples were mixed with KBr in the ratio 1:100 by weight and pressed into pellets for IR measurements. XPS study was carried out in an ESCA-3 Mark II spectrometer (VG Scientific Ltd., England) employing Al K_{α} radiation (1486.6 eV) at a pass energy of 50 eV. The powder samples were made into pellets of 8 mm diameter and placed into an UHV chamber housing the analyser at 10^{-9} Torr. Before the measurements were carried out, the samples were kept at the preparation chamber for 5 hours for desorption of gases. Binding energies were measured with a precision of ± 0.1 eV. The charging effect was taken care of with respect to the C(1s) peak of adventitious carbon at 285 eV. XAFS spectra at La and Mn K-edges were recorded using BL01B1 XAFS beam-line at SPring-8. Si(111) crystal served as the monochromator for Mn K-edge EXAFS and Si(311) for La K-edge EXAFS. For Mn K-edge measurements, fine powder of the sample was brushed onto scotch tape. A number of layers of tape were stacked to obtain total absorption lengths $\mu x \approx 2.5$ (where x is the thickness of the sample). For La K-edge measurements, the absorbers were made by pressing the samples into pellets of 10 mm diameter with boron nitride as binder. The thickness(x) of the absorber was adjusted such that $\mu x \geq 1$. The pre-edge absorption was removed by fitting the data to a linear background and a simple cubic spline was used to simulate the embeddedatom absorption, μ_0 , above the edge. The XAFS oscillations χ were obtained as a function of photoelectron wave vector $k = \sqrt{2m(E-E_0)/h^2}$. E₀ was estimated from the first inflection point of the main edge. XAFS oscillations, $\chi(k) = (\mu - \mu_0)/\Delta\mu$ (k - photoelectron wavenumber, μ_0 - atomic absorption coefficient and $\Delta \mu$ - edge jump), were extracted following standard procedures. FEFFIT program [37] was used to fit Fourier transformed (FT) $k\chi(k)$ data in r space to the theoretical spectra calculated using FEFF6.01 [38]. The Mn and La K-edge EXAFS data were analysed in tandem, to improve the reliability of fits.

Table 1. Doping content, excess oxygen, lattice-parameters a and c, Susceptibility, calculated spin-only susceptibility, and T_C for low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$. * sign implies VRH supported ferromagnetic transition around room temperature.

х	δ	$a(\text{\AA})$	$c(\mathrm{\AA})$	χ_{exp} (μ_B /f.uOe)	$\chi_{spin-only}$ (μ_B /f.uOe)	$T_C(K)$
00	0.00	5.49(1)	13.35(1)	3.53	3.67	360
0.03	0.01	5.49(1)	13.36(1)	1.79	3.58	325
0.05	0.02	5.49(1)	13.34(1)	2.55	3.52	305
0.10	0.03	5.49(1)	13.34(1)	0.75	3.37	$\sim 300^*$
0.20	0.05	5.50(1)	13.36(1)	0.76	3.07	$\sim 300^*$

3. Results

3.1. X-ray Diffraction

The X-ray diffraction patterns of $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ ($0 \le x \le 0.20$) and the fits obtained from Rietveld refinement of the X-ray diffraction data are presented in Fig.1. All the samples crystallize in a single phase rhombohedral structure (space group R3C). Lattice parameters calculated for hexagonal setting of the space group are found to be a = 5.486 Å and c = 13.345 Å for the pristine sample [Table 1]. These values closely match with those reported earlier [27]. A monotonous increase in lattice parameters has been observed by Kallel et al [27] and Kim et al [29] for increasing doping content x in their Ti doped $La_{0.7}Sr_{0.3}MnO_3$ samples. They have ascribed the increase in lattice parameters to larger Ti^{4+} ions substituting smaller Mn^{4+} ions. In our samples, we observe non-systematic and relatively insignificant changes in the lattice parameters with the doping content x. Recently, Zhu et al [34] have reported that in thin films of $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$, the lattice parameters initially increase with doping content x for $x \leq 0.3$ and decrease for further doping. The decrease in the lattice parameters is said to be due to smaller Ti⁴⁺ ions substituting larger Mn³⁺ ions. The nearly constant lattice parameters observed in our samples, could therefore be possible if Ti⁴⁺ ions substitute both larger Mn³⁺ and smaller Mn⁴⁺ ions at random.

3.2. Infra-red Absorption Studies

To investigate the valence of Ti ions, IR transmission spectra of the samples were recorded [Fig. 2]. For the undoped sample, two broad absorption peaks are clearly seen between 350-500 cm⁻¹ and 500-750 cm⁻¹, respectively. The first peak between 350-500 cm⁻¹ is attributed to Mn-O-Mn bending mode and the second to Mn-O stretching mode of vibration [39, 40]. The widening of the stretching peak with doping content x may be induced due to the Mn-O and Ti-O stretching vibrations in Mn-O-Ti structure. Two additional shoulders at 670 cm⁻¹ and 540 cm⁻¹ appear in heavily doped Ti samples. These frequencies match closely with TiO₂ absorption peaks [23] indicating that Ti is primarily in tetravalent state. Recently temperature dependent IR absorption studies have been reported on some of these compounds [35].

3.3. Magnetic susceptibility

Plots of AC magnetic susceptibility as a function of temperature for $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_3$ $(0 \le x \le 0.20)$ samples are presented in Fig. 3. For $0.0 \le x \le 0.05$, a sharp rise in AC susceptibility is seen indicating paramagnetic to ferromagnetic transition. The ferromagnetic ordering temperature (T_C) decreases with increasing doping content. T_C obtained for the parent compound is close to 360 K which is as reported earlier [41] and it decreases to 325 K and 305 K for x = 0.03 and 0.05 respectively [Table 1]. However, AC susceptibility of x = 0.05 sample is greater than that of x = 0.03 sample. Such anomalies have been reported earlier [42, 43]. Blasco et al [42] have attributed it to the magnetic inhomogeneity in their samples. In our case too, the anomaly may perhaps be due to the non-uniform distribution of magnetic $Mn^{3+/4+}$ ions in the sample. It is also possible that the Mn³⁺ content in x = 0.03 sample is smaller than that in x = 0.05sample. Such a possibility does exist and can be seen in the error estimates of Mn^{3+} and Mn⁴⁺ contents in the two samples [Table 3]. For x = 0.10, AC susceptibility exhibits two distinct features close to 300 K and 120 K respectively. This sample has a ferromagnetic like transition just above 300 K but the magnitude of the susceptibility per formula unit (f.u.) is very much less than the calculated spin only moment value for a fully ordered sample [Table 1]. Susceptibility again shows a tendency to increase around 120 K. Such a behavior has been reported in $La_{1.4}Sr_{1.6}Mn_{2-y}Ti_yO_7$ [44]. For x =0.2, a similar weak and broad magnetic transition is observed at about 300 K with slope of the ac susceptibility-temperature curve being negative over the entire temperature range. It may be mentioned here that the calculated moment values shown in Table 1 are with an assumption that Ti⁴⁺ ions substitute only Mn⁴⁺ ions.

To investigate the magnetic properties further, DC susceptibility measurements were carried out in the temperature range of 300 K to 10 K in an applied field of 100 Oe on $0.00 \le x \le 0.20$ samples, using a Faraday balance. Similar measurements were also carried out on the high-temperature annealed x = 0.10 sample. Susceptibility of the high-temperature annealed sample with x = 0.10 shows single ferromagnetic transition at temperature 220 K [Fig. 4(a)] whereas corresponding low-temperature annealed sample shows two transition like-features at 300 K and 120 K respectively [Fig. 4(b)]. Susceptibility at saturation is substantially more for high-temperature annealed sample compared to that of the corresponding low-temperature annealed sample. The weak susceptibility observed around 300 K in the low-temperature annealed sample is not visible in the high-temperature annealed sample.

In other Ti doped $\text{La}_{1-x}\text{B}_x\text{MnO}_3$ (B = Ca, Pb, Sr) compounds, in which Ti⁴⁺ ions is reported to substitute only Mn⁴⁺ ions, T_C continuously decreases as a function of dopant concentration x. This is due to the weakening of double exchange interaction between Mn³⁺-O-Mn⁴⁺ chains. In the present case, the T_C initially decreases up to x = 0.05 and then remains at about 300 K for the further doping $0.10 \ge x \ge 0.20$.

Excess oxygen concentration or equivalently cation vacancies[45] in the samples and non-magnetic Ti ions isolate the regions of DE pairs from one another and results in lower susceptibility of the samples. Yet, higher T_C for $x \ge 0.10$ samples observed in the samples $x \ge 0.10$ could be possible if such isolated DE pairs are ferromagnetically linked with one another via a hopping polaron. Polaronic conduction could probably answer the higher T_C for $x \ge 0.10$ samples. In order to ascertain the exact mechanism responsible for higher T_C for $x \ge 0.10$ samples, transport properties of this series have been studied.

3.4. Resistivity

The resistivity as a function of temperature is plotted in Fig. 5 for the Ti doped samples along with the parent compound $La_{0.67}Sr_{0.33}MnO_3$. The parent compound shows a metallic behavior in the entire temperature range studied which augurs well with its metal-insulator (M-I) transition temperature of about 330 K [41]. At x = 0.03 and 0.05, M-I transition takes place at 270 K and 260 K respectively. For $x \ge 0.10$, lowtemperature annealed sample exhibits insulating behavior throughout the temperature range studied while the high-temperature annealed sample (x = 0.10) shows the MI transition at 180 K [Fig. 4(c)]. Increase in resistivity with doping content x in the lowtemperature annealed samples is mainly due to the weakening of the Mn³⁺-O-Mn⁴⁺ DE bonds by interspersed Ti ions. At $x \ge 0.10$, the ferromagnetic double-exchange energy reduces to the extent that it does not become comparable to the thermal energy at any temperature down to 80 K so as to bring about the M-I transition. It is to be noted that corresponding high-temperature annealed sample exhibits M-I transition.

To study the transport mechanism of the charge carriers, generally three models are discussed for semiconductor like behavior; the band-gap model [46], a nearest neighbour hopping (NNH) model [47] for the transport of small polarons and Variable Range Hopping(VRH) model [48]. In the band gap model, resistivity of the sample is given by Arrhenius law $\rho = \rho_0 \exp(E_{\rho}/k_B T)$. This equation could not be fitted to the resistivity curves of our low temperature annealed samples indicating that the activated transport across the band gap does not occur in these samples in the temperature range studied. For the NNH model, the resistivity is given by $\rho = AT \exp(W_P/k_BT)$ with $W_P = E_P/(2-t)$. E_P denotes the polaron-formation energy and t the electronictransfer integral. In Mott's VRH model, the resistivity is expressed in the form of $\rho = \rho_{inf} \exp(T_0/T)^{1/4}$ [44]. Figure 6 shows that up to x = 0.05, the transport of the charge carriers in the insulating region can be explained by both, NNH and VRH. For x = 0.20, resistivity can be fitted only by VRH equation and for x = 0.10, resistivity deviates from VRH behavior below 130 K. For x = 0.20 resistivity could not be measured below 180 K due to its very high value. The details of the fitting ranges are presented in Table 2. Further, it may be noted that for x = 0.10 sample, the deviation of resistivity from VRH curve at 130 K matches closely with the second

Table 2. Doping content and lower temperature limit down to which NNH or VRH fit is possible for low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$, higher limit being 300 K. *Resistivity could not be measured below 180 K due to the high resistivity of the sample.

х	NNH	VRH
0.03	$276~{\rm K}$	$291 \mathrm{K}$
0.05	$282~{\rm K}$	$280 \mathrm{K}$
0.10	$217~{\rm K}$	$130 \mathrm{K}$
0.20	$236~{\rm K}$	Entire range [*]

transition observed in susceptibility plots in Figs. 3 and 4(b). The weak ferromagnetic transition around 300 K could then be due to the interaction between Mn ions via a hopping polaron. Below 130 K the hopping energy weakens and Mn rich ferromagnetic regions get isolated. These isolated ferromagnetic regions then order leading to the second ferromagnetic transition. In sample with x = 0.20, perhaps VRH mechanism extends to a still lower temperature and the second transition is therefore not distinctly observed.

It is to be noted that the charge-transport in the high-temperature annealed x = 0.10 sample clearly shows activated behavior across a band-gap. An exponential curve could be fitted to the resistivity-temperature plot of this sample between 228 K to room temperature. Thus the absence of M-I transition in the low temperature annealed x = 0.1 sample can be understood to be due to inhomogeneous substitution of Ti⁴⁺ ions. Whereas in the high temperature annealed samples, the decrease in the metal to insulator transition temperature with increasing Ti doping level can be ascribed to the replacement of some of the Mn^{3+} -O- Mn^{4+} bonds by the Mn^{3+} -O- Ti^{4+} bonds. This could be either due to grain boundary effect or the disorder in Ti doping at lower annealing temperatures. If grain boundaries are the cause of higher resistance, then the volume fraction sensitive properties like magnetization should not be affected [49]. However, it can be clearly seen that the behavior of magnetization for x = 0.1 of both high temperature annealed and low temperature annealed samples is different. Hence grain boundary effect cab be ruled out. A disorder in Ti substitution could create isolated Mn³⁺-O-Mn⁴⁺ pairs connected through a polaron with a variable hopping range. This fact highlights the importance of VRH in transport and magnetic properties, and especially, high T_C , in $x \ge 0.10$ low-temperature annealed samples.

3.5. Magnetoresistance (MR)

Isothermal MR is negative for all the low-temperature annealed samples $(0.00 \ge x \ge 0.20)$ [Fig. 7]. For x = 0.00, MR shows H² dependence in all the isothermals which is consistent with the ferromagnetic order of the sample. Magnetoresistance of La_{0.7}Sr_{0.3}MnO₃ has been reported[41] to be 45% percent at room temperature under the field of 6 T. Magnetoresistance observed for the parent sample La_{0.67}Sr_{0.33}MnO₃ in

our series is 38% at room temperature and under the field of 5 T. The extrapolation of the isothermal to 6 T shows that the MR observed matches closely with that reported in [41]. Isothermal MR for samples with x = 0.00, 0.03 and 0.05 increases gradually at all the temperatures at which MR is recorded. However for x = 0.10, the isothermal MR is smaller than those of x = 0.00, 0.03 and 0.05 up to 210 K and increases substantially between temperatures 210 K and 85 K. The sudden increase in MR points towards a ferromagnetic transition within this temperature range. It is clear from the susceptibility plots [Fig. 3 and 4(b)] that this sample has a second magnetic transition at about 120 K. The plot of resistivity in presence of magnetic field (H = 5 T) also shows a deviation at about the same temperature from the zero field resistivity curve [Fig. 8].

MR is shown to be much higher at ferromagnetic transition temperature in the case of $La_{0.815}Sr_{0.185}MnO_3$ [17] and cation deficient $LaMnO_3$ [45]. In this case however, MR for the low-temperature annealed x = 0.10 sample is very small from 300 K to 210 K in spite of a ferromagnetic transition at about 300 K. MR at 300 K is only of the order of 4% as compared to, 50% observed by S. Hébert et al [45] and around 45% to 66% for increasingly cation deficient samples studied by Bukowaski et al [17]. The low MR at 300K further rules out the grain boundary effect as it would induce intergrain tunneling MR. Moreover, the sudden increase in MR between 200 K and 80 K indicates that the ferromagnetic transition temperature lies somewhere within this range. These facts clearly indicate that ferromagnetic transition at room temperature observed in low-temperature annealed $x \ge 0.10$ samples is due to VRH transport of charge carriers that connects regions of DE pairs isolated from one another by non-magnetic Ti ions.

3.6. XPS Studies

From the above studies it is clear that in the low temperature annealed samples, doping of Ti⁴⁺ for Mn ions results in isolated Mn rich regions wherein DE is active. If this is indeed the case then the ratio of Mn^{3+} to Mn^{4+} should not increase exponentially due to depletion of Mn⁴⁺ by Ti⁴⁺. To estimate Mn⁴⁺ and Mn³⁺ contents, X-ray photoelectron spectra of Mn 2p levels has been recorded for all the samples. Figs. [9 and 10] show the Mn 2p X-ray photoelectron spectra of the samples in the range $0.0 \le x \le 0.20$. The Mn 2p spectra exhibit two main peaks around 642.5 eV and 654 eV. The two main peaks correspond to the spin-orbit split $2p_{\frac{3}{2}}$ and $2p_{\frac{1}{2}}$ levels, whereas the weak structure at around 24 eV from the main peak is the satellite of the $2p_{\frac{1}{2}}$ peak. The satellite of the $2p_{\frac{3}{2}}$ peak is not visible because it overlaps with the $2p_{\frac{1}{2}}$ peaks. The two main peak features were deconvoluted, into those corresponding to $2p_{\frac{1}{2}}$ and $2p_{\frac{3}{2}}$ states of Mn^{3+} , Mn^{4+} ions and their satellites, with the help of a curve fitting Peakfit software program. The peaks for $2p_{\frac{3}{2}}$ ions are higher in binding energy than those for $2p_{\frac{1}{2}}$ states by about 11.6 eV. The Mn^{4+} and Mn^{3+} contents are calculated from areas under the curves corresponding to Mn^{4+} and Mn^{3+} ions and are presented in Table 3. Results obtained clearly indicate the presence of greater Mn⁴⁺ content in the samples compared to a case wherein Ti⁴⁺ replaces only Mn⁴⁺ and maintains charge balance. Higher Mn⁴⁺

Table 3. Percentage Mn^{4+} and Mn^{3+} contents in low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ samples obtained from XPS data. Ti content was kept fixed to its nominal value. Figures in the bracket indicate the maximum estimated error.

Doping content (x)	Mn^{4+}	Mn^{3+}	Ti^{4+}
0.03	32.2(0.9)	64.8(0.9)	3
0.05	32.9(0.7)	64.1(0.7)	5
0.10	27.7(0.8)	62.3(0.8)	10
0.20	23.7(0.8)	56.3(0.8)	20

content is possible only if substitution of Ti⁴⁺ is inhomogeneous such that it results in regions of sample that are rich in Mn perhaps surrounded by Ti rich regions.

3.7. XANES

Normalized Mn K-edge XANES spectra of $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ samples, recorded at room temperature, are shown in Fig. 11. The data was recorded with a step of 1eV, primarily for EXAFS studies. The main peak arising due to 1s - 4p electronic transition and a prepeak at about 15 eV below the main peak are clearly visible in all the spectra. Positions of the main peak and the prepeak are almost the same over the entire doping range ($0.00 \le x \le 0.20$). All peak profiles are very similar to each other. Surprisingly, chemical shift of the main edge is not observed over the entire doping range. The chemical shift of the inflection point of the main absorption edge is reported to be 4.2 eV for Mn³⁺ and Mn⁴⁺ ions in LaMnO₃ and CaMnO₃, respectively [50, 13]. If Ti⁴⁺ ions substitute Mn⁴⁺ ions alone, the effective valence of the Mn ion will shift towards +3 with the increasing doping content x and such a change should be observed in the form of shift of the main edge in Mn K-edge XANES spectra. Such effect has been reported in Ti doped La_{0.7}Ca_{0.3}Mn_{1-x}Ti_xO₃ samples [51]. The absence of a chemical shift in our sample indicates that average valence of Mn ions does not change appreciably in our samples which again supports findings from XPS study.

3.8. EXAFS

EXAFS investigations of the low-temperature annealed samples have been undertaken to see the changes in octahedral environment of Mn due to Ti substitution. Our interest lies mainly in La-(Mn/Ti), and Mn-(Mn/Ti) bond lengths, as these correlations can reveal whether the substitution of Ti⁴⁺ ions is homogeneous or not. Both La and Mn K-edge EXAFS were fitted initially for La_{0.67}Sr_{0.33}MnO₃ sample. Bond lengths were varied first, followed by their Debye Waller Factors (σ^2). Coordination numbers were initially fixed at the known crystallographic values while fitting Mn K-edge EXAFS and in case of La K-edge EXAFS, twelve La-O bond lengths were divided into three different groups of coordination numbers 3, 6 and 3 respectively on the basis of closeness of their values. Later, coordination numbers had to be varied slightly to improve the quality of fit. The Mn K-edge and La K-edge EXAFS were fitted in tandem so that equal La-Mn bond lengths were obtained from both the EXAFS.

Fourier transforms of La K-edge EXAFS are presented in Fig. 12. The FT spectra are not corrected for phase shift, however, the values of bond lengths reported in the text and tables are phase corrected corrected values. La K-edge EXAFS were fitted with k-weighting = 1, in R-range 1.5 Å to 3.5 Å and k-range 3 Å⁻¹ to 16 Å⁻¹. Fits of inverse Fourier transforms of La K-edge EXAFS for x = 0.00, 0.05, 0.10 and 0.20 samples are presented in Fig. 13. Results of La K-edge EXAFS analysis are presented in Table 4. Nearly constant bond lengths in these materials are in tune with the almost similar lattice parameters obtained by analysis of X-ray diffraction spectra of these materials [Table 1]. As La-(Mn/Ti) correlation in La K-edge EXAFS includes scattering from $Mn^{3+/4+}$ and Ti^{4+} ions, if Ti^{4+} ions were to selectively replace Mn^{4+} ions, the bond length and σ^2 of this correlation should have systematically increased with doping content x. However, no such change is seen either in the bond length or σ^2 . Furthermore, the changes in La-O bond lengths and their Debye Waller Factors are small and non-systematic and can be understood in terms of random distribution of Ti^{4+} , Mn^{3+} and Mn^{4+} ions in the sample and resulting oxygen displacement that is permissible in rhombohedral $(R\bar{3}C)$ structure. In this structure, effect of oxygen displacements will be evident, more on $La-O_1$ and LaO_3 bonds compared to that on $La-O_2$ which can be seen from the values reported in Table 4.

Mn K-edge EXAFS were fitted with k-weighting, in R-range 1Å to 3.6Å and krange 3Å to 12\AA^{-1} . Fits of the inverse Fourier transforms of Mn K-edge EXAFS for x = 0.00, 0.03, 0.10 and 0.20 are presented in Fig. 14 and the results of the Mn K-edge analysis are presented in Table 5. Mn-O, Mn-Mn as well as Mn-La bond lengths for the samples, obtained by fitting Mn K-edge EXAFS, are almost the same over the entire doping range $0.00 \le x \le 0.20$. Ti substitution should have seriously affected the Mn-O bond length at least at 10% and 20% doping level due to ionic size difference. Therefore a constant bond length could only imply that for most Mn ions, the local environment remains unaltered. This is possible only if in the low temperature annealed samples the Ti substitution is inhomogeneous leading to creation of Mn rich regions over bulk of the sample separated by Ti rich regions. This is further supported from the fact that Mn-(Mn/Ti) distance obtained from Mn K-edge EXAFS and La-Mn distance obtained from Mn K-edge as well as La K-edge EXAFS remains almost the same throughout the series. Therefore in the case of inhomogeneous doping, even at high doping level of 20%, the local environment around La and Mn ions would be similar to that in the undoped sample giving constant bond lengths through out the series.

4. Discussion

Structural, transport and magnetic properties of Ti doped low-temperature annealed $La_{0.67}Sr_{0.33}MnO_3$ samples have been investigated in the present study. Samples with

		x=0.00	x = 0.05	x=0.10	x=0.20
$La-O_1$	R(Å)	2.48(1)	2.43(1)	2.45(1)	2.44(1)
	$\sigma^2({ m \AA}^2)$	0.008(2)	0.007(2)	0.006(1)	0.005(1)
	n	3.3(5)	3.3(5)	3.3(3)	3.3(2)
$La-O_2$	$ m R(m \AA)$	2.64(1)	2.63(1)	2.63(1)	2.62(1)
	$\sigma^2(\text{\AA}^2)$	0.010(2)	0.008(3)	0.008(1)	0.009(1)
	n	6.2(7)	6.2(6)	6.2(4)	6.2(3)
$La-O_3$	$ m R(m \AA)$	2.84(2)	2.85(3)	2.86(1)	2.84(1)
	$\sigma^2({ m \AA}^2)$	0.005(2)	0.006(3)	0.006(6)	0.009(2)
	n	2.7(6)	2.7(7)	2.7(3)	2.7(5)
La-Mn/Ti	$ m R(m \AA)$	3.39(1)	3.39(1)	3.39(1)	3.39(1)
	$\sigma^2({ m \AA}^2)$	0.005(1)	0.005(1)	0.006(1)	0.007(1)
	n	8.1(4)	8.1(4)	8.1(2)	8.1(2)

Table 4. Structural parameters obtained from La K-edge EXAFS for lowtemperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ samples. Figures in brackets indicate uncertainties in the last digit.

Table 5. Structural parameters obtained from Mn K-edge EXAFS for lowtemperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ samples. Figures in brackets indicate uncertainties in the last digit.

		x=0.00	x=0.03	x=0.10	x=0.20
Mn-O	R(Å)	1.92(0)	1.92(0)	1.92(1)	1.92(1)
	$\sigma^2(\text{\AA}^2)$	0.007(2)	0.007(1)	0.007(2)	0.009(1)
	n	5.5(6)	5.5(3)	5.5(4)	5.7(5)
Mn-(La/Sr)	$ m R(m \AA)$	3.39(1)	3.39(0)	3.39(0)	3.39(0)
	$\sigma^2({ m \AA}^2)$	0.009(1)	0.010(1)	0.010(1)	0.010(1)
	n	8.4(7)	8.4(5)	8.4(5)	8.6(5)
Mn-(Mn/Ti)	$ m R(m \AA)$	3.81(1)	3.82(0)	3.81(0)	3.82(0)
	σ^2	0.004(1)	0.002(0)	0.003(1)	0.002(1)
	n	5.6(5)	5.6(2)	5.6(2)	5.6(2)
Mn-O-(Mn/Ti)	$ m R(m \AA)$	3.83(1)	3.83(1)	3.83(1)	3.83(1)
	$\sigma^2(\text{\AA}^2)$	0.007(5)	0.007()	0.007(1)	0.007(0)
	n	12.4(6)	12.4(3)	12.4(3)	12.3(4)

10 % Ti doping were prepared using two different preparation schedules. High temperature annealed sample shows transport and magnetic properties similar to those reported in [27] whereas low temperature annealed sample has significantly different properties. With increasing doping content x, substitution of Ti⁴⁺ (3d⁰) ions in Mn⁴⁺-O-Mn³⁺ chains is expected to increase resistivity of the samples. However, lowtemperature and high-temperature annealed x = 0.10 samples exhibit significantly different resistivities. Moreover, high-temperature annealed x = 0.10 sample exhibits metal-insulator transition whereas low-temperature annealed sample is insulating over the entire temperature range investigated. This fact can be understood in terms of additional scattering centres like cation vacancies or substitutional inhomogeneities in the low-temperature annealed sample. The short-range order of $Mn^{4+}-O-Mn^{3+}$ chain, caused by interspersed Ti ions and cation vacancies, gives rise to the insulating behavior of low-temperature annealed $x \ge 0.10$ samples. Strong localizing effect of random cation vacancies is well known [52]. The random potential fluctuations due to cation vacancies and dopant Ti ions favor Anderson type localization, but the localized wavepackets are large enough for the e_g electrons to extend over several sites to provide the ferromagnetic interaction. The samples with $x \le 0.05$ exhibit sharp rise in susceptibility indicating paramagnetic to ferromagnetic transition and the ferromagnetic ordering temperature (T_C) decreases with increasing doping content, indicating the gradual weakening of double-exchange coupling. However, for x = 0.10 and 0.20 samples, T_C remains at around 200 K, which is significantly higher than these reported for Ti depend mangemitics

 (T_C) decreases with increasing doping content, indicating the gradual weakening of double-exchange coupling. However, for x = 0.10 and 0.20 samples, T_C remains at around 300 K, which is significantly higher than those reported for Ti doped manganites [23, 24, 27, 29, 25]. Higher T_C of these samples can not be simply attributed to the presence of cation vacancies. It is to be noted that T_C decreases with increasing cation deficiency in La_{1-x}Mn_{1-y}O₃ [53]. In case of La_{0.815}Sr_{0.185}MnO_{3+ δ} samples, T_C shifts marginally on the higher temperature side for small δ and shifts towards lower temperatures for higher δ [17]. Hence, high T_C in our low temperature annealed x = 0.10and x = 0.20 samples may not be due to cation vacancies alone, it could be related to Ti doping as well. Moreover, susceptibility (emu/f.u.) of these x = 0.10 and 0.20 samples is significantly smaller and can not be explained only on the basis of substitution of Mn⁴⁺ ions by non-magnetic $Ti^{4+}(3d^0)$ ions. Decrease in susceptibility of such a magnitude is not observed in Ti doped samples of the same composition [27, 29, 26]. T_C of our hightemperature annealed x = 0.10 sample matches closely with the reported value of T_C [27]. In fact susceptibility is found to be higher in cation deficient samples than that in stoichiometric samples [54] and similar magnitudes in [17]. Reduced susceptibility in our low-temperature annealed samples can be explained to be due to formation of isolated Mn rich regions where in DE is active separated by Ti rich regions. These regions are created due to inhomogeneous Ti substitution.

IR spectra indicate that Ti is predominantly in tetravalent state in these compounds. Moreover, IR spectra exhibit a slight shift in the stretching mode absorption peak in our samples. It has been argued on the basis of fixed Mn-O stretching mode absorption peaks in $La_{0.67}Ca_{0.33}Mn_{1-x}Ti_xO_3$ that Jahn-Teller distortions are not affected by doping of Ti ions in their materials[23]. In the case of low temperature annealed samples, the 600cm⁻¹ mode is a quite broadened. This can be related to the presence of disorder in the Ti doping. In this sample Ti^{4+} substitution causes regions wherein double exchange is active and regions which are devoid of double exchange pairs. Perhaps the two regions are in such a proportion that the strengths of respective stretching modes are nearly same resulting in a broad absorption dip. XRD studies indicate the complete solid solubility of Ti in the parent $R\bar{3}C$ structure of $La_{0.67}Sr_{0.33}MnO_3$ with lattice parameters of the low-temperature annealed samples remaining nearly the same over the entire doping range. This also supports the above idea of inhomogeneous substitution of Ti^{4+} ions. Such a substitution would result in formation of Mn rich regions that are isolated from each other by regions that are rich in Ti. This would increase the magnitude of

14

resistivity due to additional scattering from the Ti rich regions, inhibit insulator to metal transition, suppress magnetic order as well as magnetoresistance, all of which are seen in low temperature annealed x = 0.1 and 0.2 samples. The weak ferromagnetic transition seen in these two samples at about 300K could then be explained to be due to ordering of Mn rich regions connected with each other by a variable range hopping polaron. The VRH of charge carriers strengthens ferromagnetic exchange-coupling between isolated ferromagnetic regions and manifests in a weak ferromagnetic-like transition at room temperature.

EXAFS results provide the further evidence for the inhomogeneous substitution of Ti ions. Local structure around La and Mn ions is found to be almost the same in doped as well as the undoped samples [Table (4), (5)]. This indicates that even at 20% dopant concentration, large parts of the sample have Mn ions in exactly similar environment as in the undoped sample. In low-temperature annealed x = 0.10 sample, Mn⁴⁺ and Mn³⁺ contents are found to be nearly 28% and 62% respectively [Table 3]. If the substitution was homogeneous, 10% Ti ions would have depleted the Mn⁴⁺ concentration so as to maintain charge balance. Instead, in the low temperature annealed samples, it can be seen that Mn⁴⁺ concentration decreases at much slower rate indicating the presence of isolated Mn rich regions.

5. Conclusions

In this paper studies of structural, magnetic, transport and spectroscopic properties of low temperature annealed $\text{La}_{0.67}\text{Sr}_{0.33}\text{Mn}_{1-x}\text{Ti}_x\text{O}_3$ ($0 \leq x \leq 0.20$) is presented. The lower annealing temperature results in an inhomogeneous substitution of Ti ions resulting in a nearly similar crystal structure and local structure around La and Mn ions but vastly different magnetic and transport properties as compared to the undoped sample. This is perhaps due to the formation of isolated ferromagnetic clusters linked to each other by a variable range hopping polaron.

Acknowledgments

K R Priolkar acknowledges financial assistance from DST under SR/FTP/PS-19/2003. Thanks are also due, to Prof M S Hegde, Indian Institute of Science, Bangalore for useful discussions and constant encouragement and to Dr N Y Vasanthacharya for DC susceptibility measurements. XAFS measurements were performed at SPring-8 under the proposal no. 2003A0028-Cx-np with financial assistance for travel from DST, New Delhi.

References

 Tokura Y eds Colossal Magnetorestive Oxides, (Gordon and Breach Science Publishers, New York, 2000).

- [2] Rao C N R and Raveau B eds Colossal Magnetoresistance, Charge Ordering and Related Properties of Manganese Oxides (World Scientific Publishing Company Pvt.Ltd., Singapore, 1998.)
- [3] Tofield B C and Scott W R 1974 J. Solid State Chem. 10 183
- [4] Sakai N, Fjellvåg H , Lebech B and Fernandez-Diaz M T 1997 Acta Chem. Scand. 51 904
- [5] Maignan A, Michel C, Hervieu M and Raveau B 1997 Solid State Commun. 104 277
- [6] Geller S (1956) J. Chem. Phys. 24 1236
- [7] van Roosmalen J A M, Cordfunke E H P, Helmholdt R B and Zandgergen H W 1994 J. Solid State Chem. 110 100.
- [8] Mizusaki J, Yonemura Y, Kamata H, Ohyama K, Mori N, Takai H, Tagawa H, Dokiya M, Naraya K, Sasamoto T, Inaba H and Hashimoto T 2000 Solid State Ionics 132 167
- [9] Maguire E T, Coats A M, Skakle J M S and West A R (1999) J. Mater. Chem. 9 1337
- [10] Hervieu M, Mahesh R, Rangavittal N and Rao C N R (1995) Eur. J. Solid State Inorg. Chem. 32 79
- [11] van Roosmalen J A M and Cordfunke E J P (1994) J.Solid State Chem. 110 106
- [12] Ferris V, Brohan L, Ganne M and Tournoux M (1995) Eur. J. Solid State Inorganic Chem. 32 131
- [13] Dezanneau D, Audier M, Vincent H, Meneghini C and Djurado E (2004) Phys. Rev. B 69 014412
- [14] Trukhanov S V, Bushinsky M V, Troyanchuk I O, and Szymczak H (2004) J. Exp. and Theo. Phys. 99 756
- [15] Trukhanov S V, Lobanovski L V, Bushinsky M V, Khomchenko M A, Pushkarev N V, Troyanchuk I O, Maignan A, Flahaut D, Szymczak H and Szymczak R (2004) Eur. Phy. J. B 42 51.
- [16] Dabrowski B, Rogacki K, Xiong X, Klamut P W, Dybzinski R, Shaffer J and Jorgensen J D (1998) Phys. Rev.B 58 2716.
- [17] Bukowski Z, Dabrowski B, Mais J, Klamut P W, Kolesnik S and Chmaissem O (2000) J. Appl. Phys. 87 5031
- [18] Mitchell J F, Argyriou D N, Potter C D, Hinks D G, Jorgensen J D and Bader S D (1996) Phys. Rev.B 54 6172.
- [19] Mizusaki J, Mori N, Takai H, Yonemura Y, Minamiue H, Tagawa H, Dokiya M, Inaba H, Naraya K, Sasamoto T and Hashimoto T 2000 Solid State ionics 129 163
- [20] Roosmalen J A M, van Vlaaderen P, Cordfunke E H P, Ijdo W L and Ijdo D J W 1995 J. Solid State Chem. 114 516
- [21] Toepfer J and Goodenough J B 1997 Solid State Ionics 101-103 1215
- [22] Nakamura K (2003) J. Solid State Chem. 173 299
- [23] Liu X, Xu X and Zhang Y 2000 Phys. Rev. B 62 15112
- [24] Sahana M, Venimadhav A, Hegde M S, Nenkov K, Roβler U K, Dorr K and Muller K -H 2003 J. Magn. Magn. Mater. 260 361
- [25] Hu J, Qin H, Chen J and Wang Z 2002 Mat. Sci. Engg. B 90 146
- [26] Troyanchuk I O, Bushinsky M V, Szymczak H, Barner H and Maignan A 2002 Eur. J. Biochem. 28 75
- [27] Kallel N, Dezanneau G, Dhahri J, Oumezzine M and Vincent H 2003 J. Magn. Magn. Mater. 261 56
- [28] Liu Y -H, Huang B -X, Zhang R -Z, Yuan X -B, Wang C -J and Mei L -M 2004 J. Magn. Magn. Mater. 269 398
- [29] Kim M S, Yang J B, Cai J, Zhou X D, James W J, Yelon W B, Parris P E, Buddhikot D and Malik S K 2005 Phys. Rev. B 71 014433.
- [30] Ulyanov A N, Kang Y -M, Yoo S -I, Yang D -S, Park H M, Lee K -W and Yu S -C 2006 J. Magn. Magn. Mater. 304 e331.
- [31] Ulyanov A N, Yang D -S, Lee K -W, Greneche J -M, Chau N and Yu S -C 2006 J. Magn. Magn. Mater. 300 175.
- [32] Alvarez-Serrano I, Lopez M L, Pico C and Viega M L 2006 Sol. State Sci. 8 37.
- [33] Nam D N H, Bau L V, Khiem N V, Dai N V, Hong L V, Phuc N X, Newrock R S and Nordblad

P 2006 Phys. Rev. B 73 184430.

- [34] Zhu X B, Sun Y P, Ang R, Zhao B C and Song W H 2006 J. Phys D: Appl. Phys. 39 625.
- [35] Priolkar K R and Rawat R 2007 J. Magn. Magn. Mater article in press.
- [36] Bajpai A and Bannerjee A 1997 Rev. Sci. Instrum. 68 4075.
- [37] Newville M, Livins P, Yacoby Y, Rehr J J and Stern E A 1993 Phys. Rev. B 47 14126.
- [38] Rehr J J, Albers R C, and Zabinsky S I 1992 Phys. Rev. Lett. 69 3397.
- [39] Popovič Z V, Cantarero A, Thijssen W H A, Paunovič N, Dohcevič-Mitrovič N and Sapina F 2005 J.Phys.:Condens.Matter 17 351.
- [40] De Marzi G, Popovič Z V, Cantarero A, Dohcevič-Mitrovič Z, Paunovič N, Bok J and Sapina F (2003) Phys. Rev. B 68 064302.
- [41] Mahendiran R, Tiwary S K, Raychoudhuri A K, Ramadrishnan T V, Mahesh R, Rangavittal N and Rao C N R 1996 Phys. Rev. B. 53 3348.
- [42] Blasco J, Garcia J, Teresa J M, Ibarra M R, Perez J, Algarabel P A, Marquina C and Ritter C 1997 Phys. Rev. B 55 8905.
- [43] Sundaresan A, Paulose P L, Mallik R and Sampathkumaran E V 1998 Phys. Rev. B 57 2690.
- [44] Zhu H, Liu X, Ruan K, and Zhang Y 2002 Phys. Rev. B. 65 104424.
- [45] Hébert S, Wang B, Maignan A, Martin C, Retoux R, Raveau B 2002 Solid State Commun. 123 311.
- [46] Kusters R M, Singleton D A, Keen D A, McGreevy R and Hayes W 1989 Physica B 155 362.
- [47] Snyder G J, Hiskes R, DiCarolis S, Beasley M R and Geballe T H 1996 Phys. Rev. B 53 14434.
- [48] Viret M, Ranno L and Coey J M D 1997 Phys. Rev. B 55 8067.
- [49] Hwang H, Cheong S -W, Ong N P and Batlogg B 1996 Phys. Rev. Lett. 77 2041.
- [50] Subías G, García J, Proietti M G, and Blasco J 1997 Phys. Rev. B 56 8183.
- [51] Cao D 2001 Phys. Rev. B 64 184409.
- [52] Coey J M D, Viret M, Ranno L and Ounadjela K 1995 Phys. Rev. Lett. 75 3910.
- [53] de Silva P S I P N, Richards F M, Cohen L F, Alonso J A, Martinez-Lope M J, Casais M T, Thomas K A, MacManus-Driscoll J L 1998 J. Appl. Phys. 83 395.
- [54] Ranno I, Viret M, Mari A, Thomas R M, Coey J M D 1996 J.Phys:Condens.Matter 8 L33.

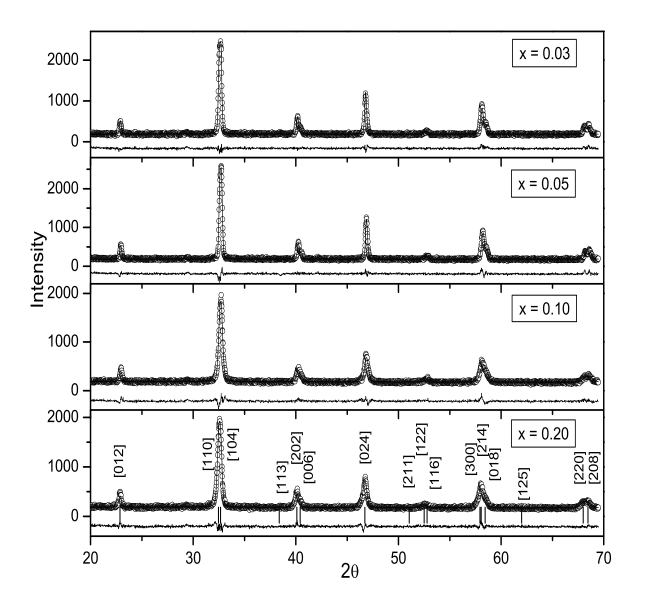


Figure 1. X-ray diffraction patterns (shown with circles) for low-temperature annealed $\text{La}_{0.67}\text{Sr}_{0.33}\text{Mn}_{1-x}\text{Ti}_x\text{O}_{3+\delta}$ ($0 \le x \le 0.20$)samples and their Rietveld fits (shown with line). Vertical lines in the bottom panel indicate the Bragg reflection positions.

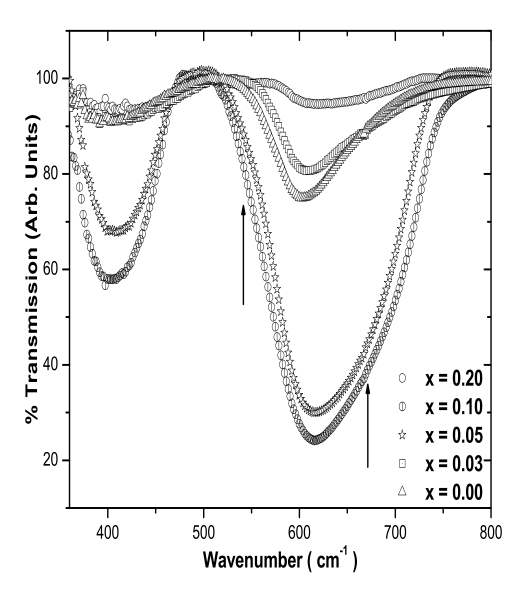


Figure 2. IR transmission spectra of low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_3$ ($0 \le x \le 0.20$). Arrows indicate Ti-O stretching modes.

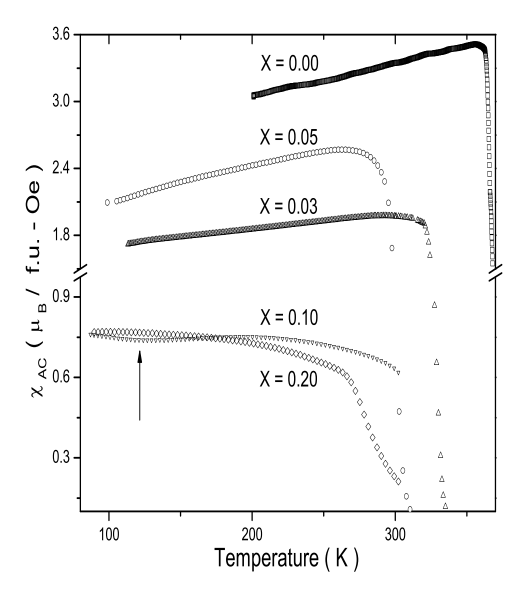


Figure 3. AC susceptibility-temperature plots of low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ ($0 \le x \le 0.20$). Arrow indicates the second ferromagnetic transition in x = 0.10 sample.

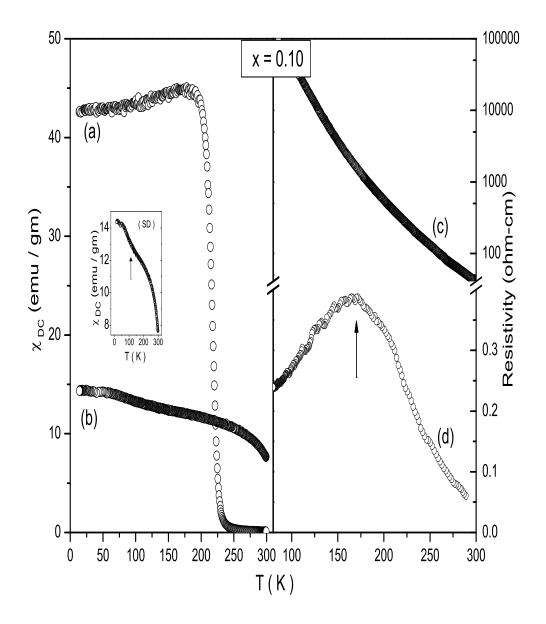


Figure 4. Temperature dependence of DC susceptibility for $La_{0.67}Sr_{0.33}Mn_{0.90}Ti_{0.10}O_{3+\delta}$ (a) high-temperature annealed, (b) low-temperature annealed samples and resistivity of (c) low-temperature annealed and (d) low-temperature annealed samples. Second ferromagnetic transition in the low-temperature annealed sample is indicated by an arrow in the inset.

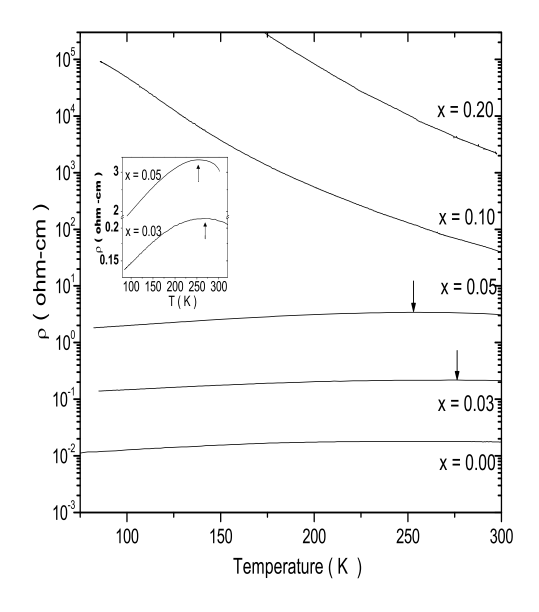


Figure 5. Resistivity versus temperature plots of low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ ($0 \le x \le 0.20$). Arrows in the inset show the metal-insulator transition in x = 0.03 and 0.05 samples.

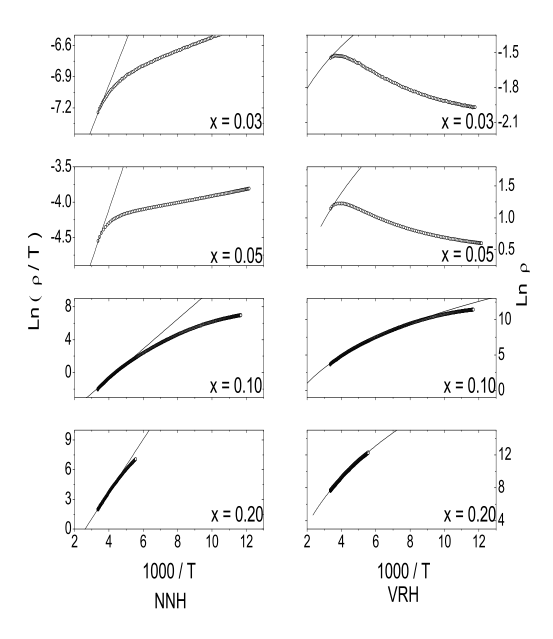


Figure 6. Nearest-neighbour and Variable Range Hopping fits shown with $\ln \rho$ and $\ln (\rho/T)$ as function of (1000 / T) for low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ (0.03 $\leq x \leq 0.20$)

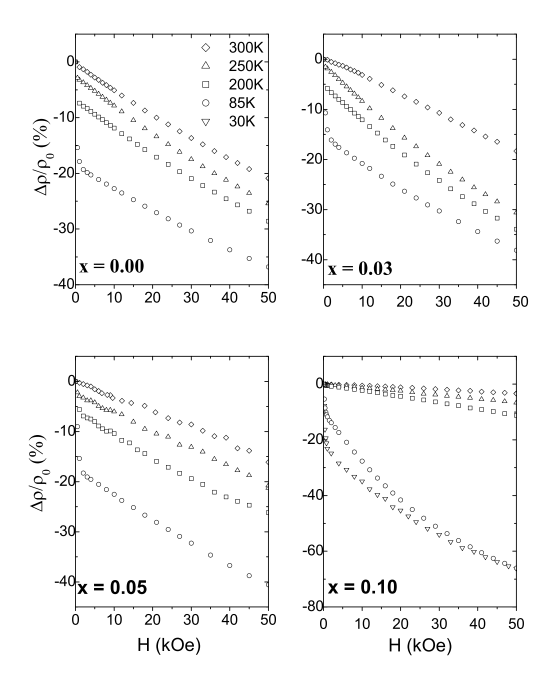


Figure 7. Magnetoresistance versus applied magnetic field plots of low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ ($0 \le x \le 0.10$)

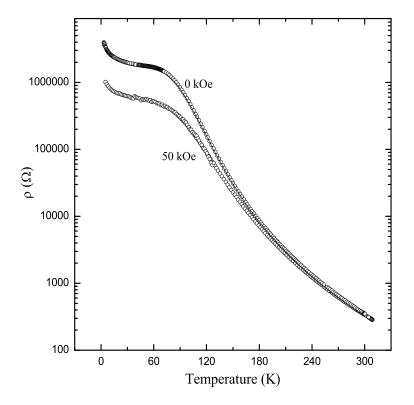


Figure 8. Resistivity under zero-field and 5T field for low-temperature annealed x = 0.10 sample.

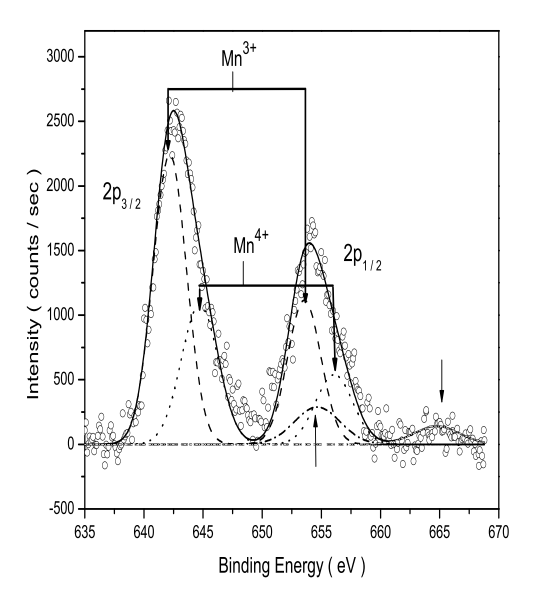
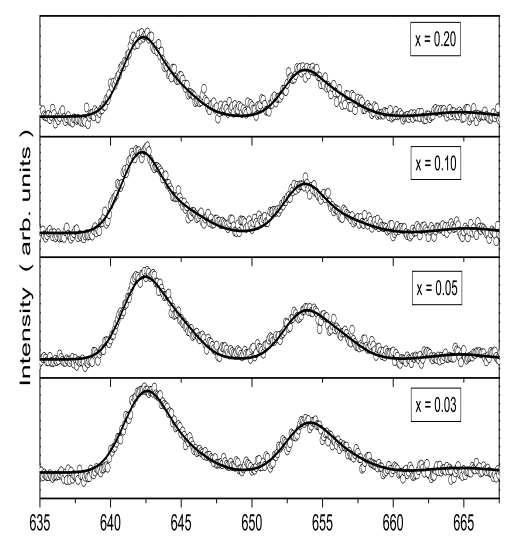


Figure 9. XPS spectrum (circles) of $La_{0.67}Sr_{0.33}MnO_3$ sample along with fitted curve (line). Discontinuous lines represent the deconvoluted $2p_{3/2}$ and $2p_{1/2}$ spectra corresponding to Mn^{3+} and Mn^{4+} ions. Arrows indicate the satellite peaks.



Binding Energy (eV)

Figure 10. XPS spectra (circles) for site-disordered $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ samples along with the fitted curve (line)

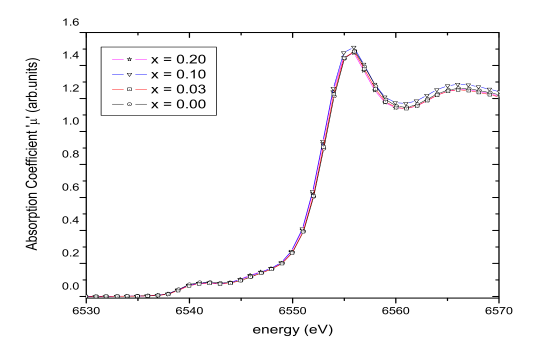


Figure 11. Normallised Xanes spectra of low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ ($0 \le x \le 0.20$).

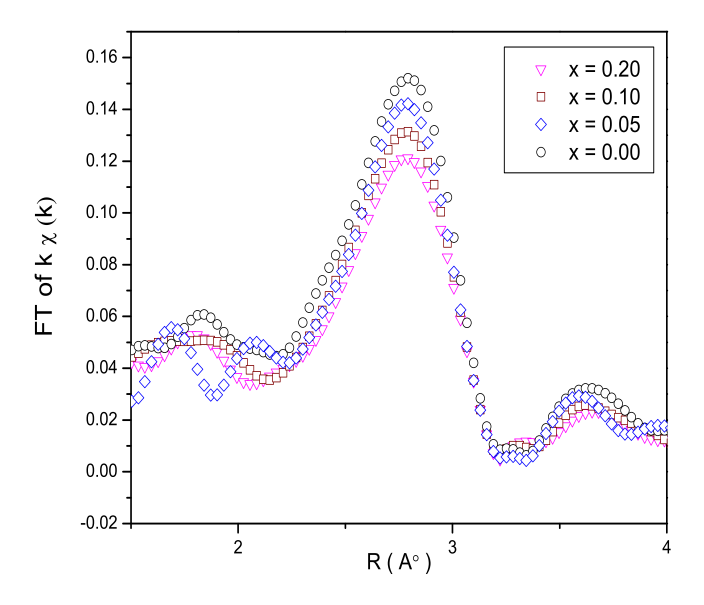


Figure 12. La K-edge Fourier Transforms of the low-temperature annealed $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$ samples.

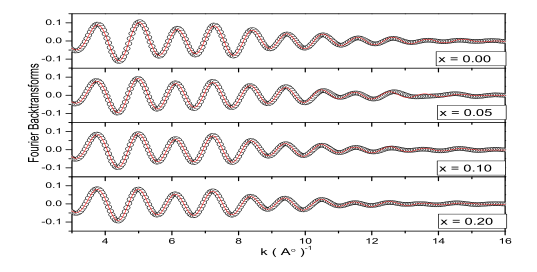


Figure 13. k - weighted backtransformed La K-edge EXAFS spectra (circles) along with fitted curves (line) in $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$.

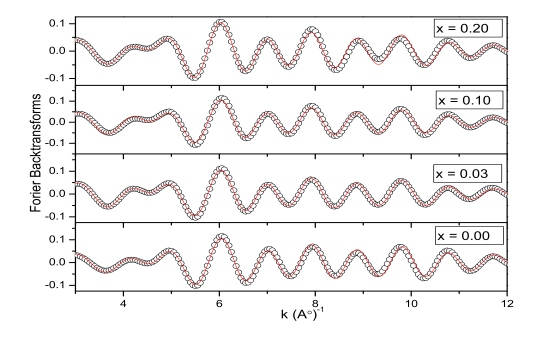


Figure 14. k - weighted backtransformed Mn K-edge EXAFS spectra (circles) along with fitted curves (line) in $La_{0.67}Sr_{0.33}Mn_{1-x}Ti_xO_{3+\delta}$.