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Metal-insulator phase transition and structural stability in 'Sb' doped CaMnO₃ perovskite

R. Kannan¹, *, D. Vanidha¹, A. Arun Kumar¹, K. U. Rama Tulasi¹, R. Sivakumar²

Email address:

kannan@pec.edu(R. Kannan)

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Abstract: Electron doped $Ca_{1-x}Sb_x$ MnO₃ (X=0 to 0.4) compound when prepared by high temperature solid state reaction, exhibits orthorhombic distorted perovskite structure. A systematic, continuous doping increases the metal – insulator transition temperature to above the room temperature (\approx 431 to 469 K); due to the mismatch of ionic radii in A site rising anti - site effect, which consequently changes the bond length. Doping induced metal – insulator transition accompanied by structural transition is reflected through the drastic changes in the parameters like cation size variance factor (σ), average ionic radius <r>, tolerance factor (t) and T_o -which is correlated to the band related transport properties. Structural transition from the phase of perovskite to Brownmillerite has been found for the compositions x=0.3 and 0.4, which is attributed due to the ionic radii mismatch. Doping exerts chemical pressure by modifying compression of A-O bond and relaxation of B-O bond, giving rise to strain similar to external pressure in ABO₃ perovskite. The dependence of electrical transport at high temperature has been studied, by employing variable range hopping and small polaron hopping model as an account for the experimental observation due to disorder induced localization.

Keywords: Phase Transition, Pervoskite, High T_c

1. Introduction

In manganites, optimum grain size for preferential orientations the need of the hour for low field magnetoresistance (LFMR) applications at ambience in the applied magnetic field for 100% spin polarization between the grains. Storage and retrieval of information from computer hard disk were revolutionized after the discovery of colossal magneto resistance (CMR) based manganite read/write heads. Miniaturization, Storage increase and High magneto resistance (MR) through LFMR, kept further renewing the interest in CMR even after three decades of fundamental researches on manganite [1, 2]. However, the origin of conductivity in doping induced disordered oxide materials is not yet clear due to charge localization. The parameters like dopant atom, particle size and pressure are varied in perovskite manganites to achieve metal insulator transition temperature (T_{MI}) above the room temperature.

CaMnO₃is an antiferromagnetic Mott-Hubbard insulator known for becoming metallic in nature at narrow doping

levels itself,asCaMnO3 is at the edge of Metal-Insulator (M-I) transition. For electron doped CMR compounds, due the high efficiency and minimum disorder AMnO₃(A=Ba,Ca,Sr) is most preferred when compared to LaMnO₃. In those compounds anti-ferromagnetic (AFM) to paramagnetic (PM)transition was reported by Esa Bose et al [3]. Doping induced structural changes are reported by Qingdi Zhou et al [4] in SrMnO₃ since MR is intrinsically connected to metal - insulator transition and charge orbital ordering. Electron doping induced cubic to tetragonel and cubic to orthorhombic phases was reported by Brendan J.Kennedy et al [5] in tetravalent Cerium (Ce⁴⁺) doped SrMnO₃ due to Jahn Teller co-operative tilting of MnO₆ octahedra. High temperatureresistivity of Sb doped CaMnO₃ [6] exhibits decrease in conductivity with increase in temperature and $d\sigma/dT$ became negative at around $600^{\circ}C$ indicating a transition from semiconducting to metallic.

CaMnO₃ when doped with Sb⁵⁺ion produces excess electrons through Mn⁴⁺ to Mn³⁺ conversion. Ping Duan et al [7] proved the existence of mixed valence states of Mn (Mn³⁺ and Mn²⁺) through XPS (X-ray photoemission

¹Department of Physics, Pondicherry Engineering College, Pondicherry-605 014, India

²Department of Physics, Pondicherry University, Pondicherry-605 014, India

spectroscopy) and upon Sb doping Curie temperature (T_c) reaches a value of 245 K in La_{1-x}Sb_xMnO₃ for x = 0.1. High phase transition temperature ($\approx T_p$ =>temperature at which maximum resistivity occurs) of 289 K has been recorded for La doped Mn²⁺ rich compound BaMnO₃ for x=0.3 [1].

Sb₂O₃ during the synthesis acts not only as reactant but also as an additive for decreasing the reaction temperature for phase formation from 1200°C to 900°C. The decreased reaction temperature for phase formation is due to the change in the eutectic point [8] of the mixture of all the reactants since additives provide liquid - phase grain growth at low temperature [9, 10]. Thus the synthesis at a temperature less than the eutectic temperature of a mixture enhances the phase formation and control of particle size [11]. High reaction temperature for phase formation, nonhomogeneous mixing and agglomeration are commonly found in synthesis of oxide materials by solid state reaction [12]. Hence, the decreased reaction temperature is one of the reasons for increased Metal - Insulator transition temperature to 431 K compared to literature value 420 K [13].

In the present work we are discussing the electrical transport properties of the 'Sb' doped Ca_{1-x}Sb_x MnO₃ (x=0 to 0.4), fabricated by low temperature sintering method. Doping induced structural phase transition accompanied by metal - insulator phase change has lead to a record high temperature of 431 K, as the T_{MI} for x=0.2 in 'Sb' doped CaMnO₃ is a consequence of A site doping. This is attributed to very large difference in ionic radii, 'Sb' (0.74 Å) atom occupies Mn (0.53 Å) site instead of Ca (1.12 Å) site making A/B cation ratio deviating from perovskite structure.Doping induced charge localization and along with phase transition leading to changes in the density of states and resistivity are found from the T₀ values [14].

2. Experiment

The electron doped bulk samples of Ca_{1-x}Sb_xMnO₃ were prepared by high temperature solid-state reaction method. Highly pure CaCO₃ (Merck, 99.9%), Sb₂O₃ (Merck, 99.9%) and MnO₂(Merck, 99.9%) were taken in the stoichiometry ratio and thoroughly mixed and ground until uniform mixing and homogeneous distribution is achieved. The finely grounded samples were taken in an alumina crucible and calcined in air at 900° C for 12 hours. The resulting powders were pelletized using hydraulic press in to small disks of 8 mm diameter and1 mm to 2mm thickness under a pressure of 3 to 5 tons. All those pellets prepared were sintered at 900° C continuously or 36 hours and cooled to room temperature.

All the five samples of Ca_{1-x}Sb_xMnO₃(x=0 to x=0.4) were prepared under exact heating conditions as described above. The samples are characterized by powder XRD (Rich-Seifert, Germany) using Cu Kα radiation. Electrical transport studies were carried out using Silver electrodes of 99.9% purity with Keithley electrometer (Model 617).

3. Results and Discussion

3.1. XRD Analysis

XRD pattern of polycrystalline $Ca_{1-x}Sb_xMnO_3(X = 0 \text{ to})$ 0.4) samples belonging to orthorhombic structure are shown in fig 1(a). In 'Sb' substituted samples, splitting of the X-ray peak (112) into (002) and (112) for x = 0.1, 0.2,0.3 and x = 0.4 was observed and attributed to the increased orthorhombicity of the compound. Average grain size is estimated using Scherrer's formula $D=k\lambda/\beta\cos\theta$ [14] and it is found to be increasing as a function of increasing composition of "x" which are tabulated in table 1. Crystallite size changes consistently with doping as 12 nm for x=0.1, 21 nm for x=0.2, 27 nm for x=0.3. Extreme end members x=0,i.e. undoped compound crystallizes at 15nm and for x=0.4, it is found to be 66 nm, due to phase segregation and limited solubility. Smaller particle size of 12-27 nm comparable to wet chemical methods areattributed due to long time sintering for 36 hours at low temperature (i.e) 900 °C. The smaller particle size due to low sintering temperature are attributed to the assistance of Sb₂O₃ as sintering aidlike Bi₂O₃ and Li₂CO₃ in synthesis of dielectrics [9,15].

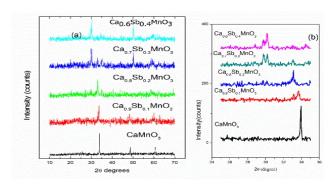


Fig 1. (a) XRD pattern of $Ca_{1-x}Sb_xMnO_3(0 \ge X \le 0.4)$ (b) Shifting of 112 peak of $Ca_{1-x}Sb_xMnO_3(0 \ge X \le 0.4)$

Lattice parameters of orthorhombic crystal structure as a function of Sb doping was found to be increasing from 4.8403 Å to 5.3213 Å for 'a', indicating that Ca^{2+} being replaced by Sb^{5+} along 'a' axis. This produces a corresponding compression in 'b' axis from 5.8426 Å ⁰ to 5.3834 Å making the tolerance factor "t" stable [Table 2]. The 'c' axis also found to undergo an elongation from 7.5401 Å to 7.7405 Å for doping concentration of $0.3 \le x \ge 0.1$. Increase in the lattice parameter for the concentration x = 0.2 to 0.4, indicates low chemical pressure induced by doping giving rise to high transition temperature [16].

Doping induced structural disorder was found from the XRD for concentrations x = 0.3 and x = 0.4. Due to large difference in ionic radii, Sb (0.74 Å) atom occupies Mn (0.53 Å) site instead of Ca (1.12 Å) site making A/B cation ratio deviating from one. Brownmillerite phase [17] of stoichiometry composition $(Ca_{1-x}Sb_x)[Mn]O_3$ becomes $(Ca_{1-x})[Sb_xMn]O_3$, which was ascertained from the disappearance of characteristic perovskite 112 peak at $2\theta =$

34° and appearance of (002) peak at $2\theta = 31.5°$ degrees of Brownmillerite phase as shown in fig1(b). This gives rise to a non-uniform change in the lattice parameters due to chemical doping pressure and cell volume 213.2 (Å)³ to 249.9(Å)³. The structural distortion produced by both the phases having orthorhombic in common is a measure of average A-size cationic radius <rable radius <rable

3.2. Electrical Transport Properties

Resistivity (ρ) as a function of temperature(T) from 303 to 673 K is shown in the Fig.2. All the ρ -T curves exhibit the same behavior indicating similar structure and phase

formation undergoing M-I phase transition at 303 K, 431 K, 435 K and 468 K for x =0.1 to 0.4 respectively. For x = 0, which corresponds undoped compound do not show T_{MI} since it is AFM insulator. The increased T_{MI} for concentrations x = 0.3 and 0.4 is attributed to anti-site doping disorder, at which perovskite phase become Brownmillerite phase [17]. Low sintering temperature, presence of excess Mn^{4+} due to Sb doping (increase in Mn^{3+} / Mn^{4+}) ratio, nature of 'Sb' as both reactant and additive and structural changes due to lattice mismatch between 'Sb' and 'Ca' ionic radii are all cannot be denied as possible causesfor increased T_{MI} .

Table 1: Lattice Parameter, Unit Cell Volume, Metal insulator Transition Temperature (T_{MI}) , And Crystalline Size of $Ca_{1:X}Sb_XMnO_3(0 \ge X \le 0.4)$.

Crystalline size (nm)	15	12	21	27	66
ρat Tp K Ω cm	-	9.34	171.17	172.23	61.042
pat 303K K.Ω cm	16.910	8.542	0.1219	22.298	0.355
TMI (K)	-	303	431	435	468
V (Å)	219.9	213.2	223.7	221.8	249.9
c (Å)	7.306	7.540	7.658	7.740	7.5146
b(Å)	5.789	5.589	5.507	5.383	6.000
a(Å)	5.200	4.840	5.303	5.323	5.424
System	CaMnO3	Ca0.9Sb0.1MnO3	Ca0.8Sb0.2MnO3	Ca0.7Sb0.3MnO3	Ca0.6Sb0.4MnO3

Table2: Ionic Radius, Size Variance Factor, Tolerance Factor, T_o , Activation Energy of $Ca_{1:x}Sb_xMnO_3(0 \ge X \le 0.4)$.

System	Ionic radius <ra> (Å)</ra>	Size variance factor σ2 (Å)	Tolerance factor 't'	T0 from VRH model X108	Small polaron hopping Activation Energy in (eV)
CaMnO3	-	-	-	8.78	0.3875
Ca0.9Sb0.1MnO3	1.136	0.0175	0.9315	2.44	0.5596
Ca0.8Sb0.2MnO3	1.092	0.0310	0.9152	1.722	1.0441
Ca0.7Sb0.3MnO3	1.048	0.0405	0.8989	1.694	0.8582
Ca0.6Sb0.4MnO3	1.004	0.0464	0.8826	2.3868	1.0786

The room temperature resistivity of the samples decreases from 16.95 K Ω cm (x = 0.0) to 0.1219 K Ω cm (x=0.2) upon systematic doping, is attributed to the increase in the charge carrier concentration (Sb⁵⁺ introduces twice as many electrons per Ca²⁺ atom) enabling double exchange (DE) mechanism [7]. Every Sb⁵⁺ dopant atom makes two Mn⁴⁺ ions converted into two Mn³⁺ ions and electron hopping between 'Mn' ions is responsible for electrical conductivity in perovskite manganites, and it also indicates that the dopant atom 'Sb' exactly replaces 'Ca' making the double exchange possible up to x= 0.2. For increased concentrations, resistivity increases to 23.205 K Ω cm and then decreases to 0.355 K Ω cm respectively for x=0.3 and x

= 0.4 as shown in table 1. The abrupt rise in resistance for x=0.3 is assigned to Jahn-Teller distortion indicating that no more double exchange is possible, favoring the insulating state due to localization of electrons [18,19]. The decreased resistivity of three order for x=0.4, is due to increased grain size of almost three times (27 nm to 66 nm respectively for x=0.3 to x=0.4) and also due to the appearance of Brownmillerite phase different from the perovskite phase. The peak resistivity is also found to follow the similar trend with the exception for x=0.2. The abrupt change in size variance factor (σ ²) from 0.0175 to 0.0310respectively for x=0.1 and x=0.2 is responsible for the observed change that are tabulated in table 2.

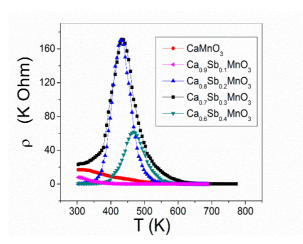


Fig 2. Resistivity versus temperature of $Ca_{1-x}Sb_xMnO_3(0 \ge X \le 0.4)$

The origin of Brownmillerite phase is attributed to limited solubility which leads to anti-site effect. Increased dopant concentration produces a chemical pressure, giving rise to a compression in the A-O band and relaxation in B-O bond making similar to external pressure which modifies the local structure of distorted orthorhombic perovskite to distorted orthorhombic Brownmillerite structure(Pbam)[17,20]. Abrupt reduction in the peak height and broadening in the $(\rho-T)$ curve for the composition x=0.3 and x=0.4 is also another form of evidence for the chemical pressure induced formation of Brownmillerite phase. Hence, the limit of increase of T_{MI} with doping is subjected to structural stability and beyond which it cannot be increased.

From the above experimental observations, we may conclude that the low sintering temperature and flux wrapping of Sb_2O_3 disfavors the formation of large grains [21,22]. Therefore, the average grain size decreases (12 nm to 27 nm for x=0 to x=0.3 and 66 nm for x=0.4) compared to normal and the grain boundary effect increases. This prevents the charge carriers from crossing the boundary between the neighboring grains leading to charge localization. Because of the increased grain boundary effect the resistivity is largely increased over the whole range of temperature studied [14]. This may be inferred from the peak broadening in XRD and consequently due to smaller particle size, grain boundary volume ratio is greater than the grains giving rise to high resistance.

High value of resistance ($M\Omega$) in Semiconducting-Insulating region is attributed to lattice mismatch (Ca /Sb) and appearance of Brownmillerite phase due to anti-site doping induced structural distortion; which in turn related to distortion favours insulating state due to localization of electrons as discussed earlier[18,19].Klien et al[21] reported that the lattice strain is mainly responsible forthe anomalies like phase segregation and magnetic interphase disorder, which gives rise to high resistance at room temperature. Resistance dependence of temperature gives information of disordering across the grain boundries.

3.3. Modeling

The experimentally observed ρ -T curves of Ca₁. _xSb_xMnO₃ samples are considered as the superimposition of two different transport phenomena named AFM metal-like behavior below T_c and PM insulating behavior above T_c . The metal like nature below T_c is explained with Zener's DE mechanism with a resistivity equation of the type $\rho(T)$ = $\rho_0 + \rho_2 T^2 + \rho_n T^n [23]$ and pure magnon scattering is represented by the equation $\rho(T) = \rho_0 + \rho_2 T^{\alpha} [24,25]$ are fitted separately here. The corresponding ρ_0 values are $0.185{\times}10^6$ (0.1168×10^6) Ωcm for Ca_{0.8}Sb_{0.2}MnO₃, $0.270 \times 10^6 (0.166 \times 10^6) \Omega cm for Ca_{0.7} Sb_{0.3} MnO_3 and$ $0.069 \times 10^{6} (0.044 \times 10^{6}) \Omega$ cmfor $Ca_{0.6}Sb_{0.4}MnO_{3}$ for both the models found to be high due to disorder induced localization.

At above T_c in all the ρ -T curves, it is observed a deviation from the linearity when the temperature is close to 550 K, where the rising of peak resistivity starts. This deviation indicates a crossover from activated carrier to another transport mechanism. The sudden drop in resistivity over a very narrow temperature range of 431 K to 523 K and saturation over wide range of temperature 550 K to 773 K, constitutes a clear evidence of change in the transport mechanism. Hence, the semiconducting insulating part of the ρ (T) curve was found to exhibit two different regimes. This abrupt change in resistivity when it is close to the peak is explained by two different models inan account of the actual experimental observations. It explains, to understand the transport mechanism in doping induced disordered oxide CMR materials at high temperature regime [14,26,27] as a variable range hopping $\rho = \rho_0 \exp (-T_0 / T)$ and a small polaron hopping p $=BTexp(Ep/k_BT)$

3.4. Variable Range Hopping Mechanism (VRH)

Conducting behaviour in manganites is explained based on doping induced disorder which gives rise to localization of energy states close to fermi level between conduction band and valence band. At high temperatures, carriers are thermally activated to their nearest neighbour site with high mobility. But when temperature decreases, thermally activated behaviour decreases and hopping process dominates, and conduction takes place by hopping of charge carriers between the localized states distributed unevenly. Normally carrier localization at the grain boundary changes the conductivity to semiconducting or insulating and carrier delocalization leads to metal like behaviour. The hopping is characterized by the parameter T_0 which is related to electronic density of states N(E_f) and radius of localization length(a) by $T_0 = 18/k_BN(E_F)a^3$. High T₀ value indicates multiphonon hopping with a large increase in the localization length and increase in barrier height at the grain boundaries leads to insulating and if T₀ decreases, based on hopping range and energy, the material remains in the metallic state of mott's transition[8,19]. Size of the grain and width of grain boundary in the order of nanometers (or the localization length 'a' $(1/\alpha)$) will correspond to 100% spin polarization by tunneling mechanism [28,29].

The resistivity data in insulator regime were fitted to the variable range hopping (VRH) proposed by Mott as shown in Fig.3 linearizing the equation $\rho = \rho_0 \exp(T_0/T)^{1/4}$ and from the slope T_o values are obtained. In the present work, for every one 'Sb⁵⁺' doped in CaMnO₃,makes two Mn⁴⁺ ions which is transferred to two Mn³⁺ ions due to rich in electron carrier makes the carrier densityN(E_F) of the system fixed. Since VRH takes place due to excitation of charge carrier across coulomb gap(W/U>>1), Ponnambalam et al [30] reported VRH mechanism over a very large temperature range of 100 K to 900 K.

Variation of T_0 with doping concentration (8.78 $\times 10^8$ to 1.694 $\times 10^8$), decreases uniformly for x=0 to x=0.3 indicating the reduction in the disorder, which in turn changes the Mn-O-Mn bond and leads to increase in the carrier concentration which is attributed to altering of (t_{2g} - e_g) orbitals by giving rise to drastic change in theband width due to electron filling [31,32] except for x=0.4. And

for x = 0.4, T_o further increases to 2.3868 $X10^8$ indicating Jahn-Teller distortion along with structural phase transition to Brownmillerite phase for x > 0.3. The decrease of T_o due to low sintering temperature indicates the improvement in the crystallinity and hence the quantum mechanical tunelling between the disorder induced localized state increases reducing the resistivity.

This variations in T_0 is attributed to the variations of (σ^2) compared to CaMnO₃ and SrMnO₃ as shown in the Table 3.The increase in T_0 value observed for our system is correlated to the structural variations caused due to the doping induced cation size disorder(σ^2)[33,34]

The overall fitting of our experimental data observation indicates that at temperatures where the thermal energy is very low, the VRH model perfectly fits perfectly to the equation and at the temperature where the thermally activated process dominates the experimental data deviates from the fitted data as shown in the Fig.3. Moreover, the fitted model of VRH proves localizations of charge carriers created by doping induced disorder are mainly responsible for the electrical conduction process in manganites.

	System	То	Activation energy	Cation size variance σ2=Σyiri2- <ra>2</ra>	Reference
	Ca1-xSbxMnO3	2.333x108 to	0.3875 to 1.0786 (ev)	0.0175 to 0.0464	Present Work
(Sb=0,.1,,0.2,0.3,0.4)		1.88x109	0.3873 to 1.0780 (CV)	0.0173 to 0.0404	Trescrit Work
	LaMnO3	1.23x102 to	163to248.9 (meV)(VRH)		Ref 29
	LaivillO3	1.601x106	103t0248.9 (IIIe v)(v K11)		Kei 29
	La(NiFe)O3	1.87x102 to	10 to 206(meV)(VRH)		Ref 32.
		1.7x107			
	(NdLnSr)MnO3	0.42431x106 to	61.48to 160.03(meV)(SPH)		
	(INCLISI)INIIOS	18.4x106	01.40t0 100.05(me v)(5111)		Ref 35.
	LaCaMnO3		92 to 116 (meV)(SPH)		Ref 36.
	LaSrMnO3	4.3x107 to			
	Lastivinos	1.3x108			Ref 21.
	Ca1-xRxMnO3				
	(R=La,Pr,Nd,Sm,Eu,Gd,Tb,		56.9(meV) to 65.6(meV)(SPH)	1.54x10-3 to 3.6x10-7	Ref 40.
	Dy,Ho,Er,Yb) $(x=0.1)$				
	Sr1-xAxMnO3			2.5x10-3 to 9x10-4	Ref 33.
(x=0 to 1)(A=Ca)				2.3X10-3 to 9X10-4	KCI 33.

Table 3: To, Cation Size Variance Factor, Activation Energies of some reported systems along with present work.

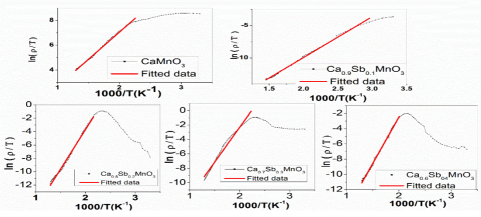


Fig. 3. Variable range hopping of $Ca_{1-x}Sb_xMnO_3(0 \ge X \le 0.4)$ [Ln (\mathbf{Q}) vs $1/T^1/4$]

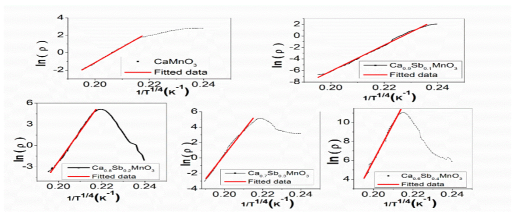


Fig. 4. Adiabatic small polaron hopping of $Ca_{1-x}Sb_xMnO_3(0 \ge X \le 0.4)$ [Ln (\bigcap /T) vs 1000/T]

3.5. Small Polaron Hopping Model

Zener's double-exchange model will not be sufficient enough to explain the very large change in resistivity in the Semiconducting-Insulating phase i.e., at high temperature region, since it does not consider spin–lattice or charge–lattice interactions, i.e., Jahn–Teller (JT) interaction and polarons, which significantly contribute [14] to the conduction process in the Semiconducting-Insulating phase.Therefore, it is very essential to understand the electrical conduction in the Semiconducting-Insulating region (where T>> $T_{\rm MI}$) which is governed by polaronic effects[20,21]. The conduction mechanism in manganites at high temperatures is governed by thermally activated small polarons based on the polaronic models.

The resistivity data in Semiconducting-Insulating region were fitted to the adiabatic small polaron hopping(ASPH) equation $\rho = BTexp(E_p/k_BT)$ where $B = [k_B/v_{ph}Ne^2R^2C(1-v_{ph}Ne^$ C) $[\exp(2\alpha R)]$, k_B is the Boltzmann constant, and T is the absolute temperature, N is the number of ion sites per unit volume, R is the average intersite spacing obtained from the relation $R=(1/N)^{1/3}$, C is the fraction of sites occupied by the polaron, α is the electron wave function decay constant, ν_{ph} is the optical phonon frequency $(v_{ph}=k_B\theta_D/h)$ and E_p is the activation energy following Padmavathi et al[35], Sayani Bhatacharyar[36], Abdel-Khalek et al[37] and Wasi Khan et al [29] for our system as shown in Fig.4. The temperature dependent polaron hopping is verified by calculating $\theta_D/2(\theta_D = \text{Debye's temperature})$, where deviation from linearity occurs in the $\ln (\rho/T \text{ vs } 1/T)$ plot and the estimated phonon frequency. Leaving the undoped compound for concentrations x =0.1 to 0.4, $\theta_D/2$ and v_{ph} (optical phonon frequency) are found to be varying linearly from 356 K, 459 K, 480 K, 495 K and 7.44X10¹², 9.53 X10¹², 9.9 X10¹²,10.35 X10¹² respectively and in agreement with the reported values[30, 33].

The activation energy E_p was found to be increasing continuously with the doping concentration from 0.5596 to 1.044 eV for x=0.1 to 0.2 indicating the presence of perovskite phase and it varies from 0.8582 to 1.0786 eV for x=0.3 and x=0.4, which is attributed to the appearance of brownmillerite phase and increase in the localization of

charge carriers[38]. The increase in the activation energy is attributed to structural phase transition associated with the inhomogenity or disorder in the barrier height and width. As well, the iinsufficient crystallization of 'Sb' in the system due to low sintering temperature (900°C) also cannot be ruled out.

Very large variations in (σ^2) compared to CaMnO₃ and SrMnO₃ as listed in the Table 3 is attributed to the large difference in the ionic radii between Ca (Ca=1.12Å) and Sb atom (Sb=0.74Å) which is due to anti-site doping induced cation disorder[28,39] and appearance of brownmillerite phase at higher 'Sb' concentrations. This gives rise to large value of activation energy compared to other system as listed in Table 3. Hence, this results indicate structural parameter σ^2 is closely related to electrical transport properties. Existence of higher activation energy is attributed to crystal distortions which increases the potential barrier height for polaron hopping [40]. When σ^2 varies from 0.0175 to 0.0464 activation energy is found to be 387.5 to 1078.6 (meV)(SPH) where as Yang Wang et al [40] reported the activation energy in the range of 56.9 to 65.6 (meV) for the σ^2 in the range of 1.54x10⁻³ to 3.6x10⁻⁷. In fact, this may be compared with Ca/Sb ratio to Ca/RE ionic radii ratio.

4. Conclusion

We have investigated the effect of 'Sb' doping on the structural and electric transport properties of CaMnO₃ in orthorhombic structure. The doping of 'Sb' for the first time increases T_{MI} to a maximum of 468K for x=0.3. The shifting of T_{MI} towards high temperature is attributed to the presence of Brownmillerite phase due to limited solubility of 'Sb' at 'Ca' site and increase in the lattice parameter for the concentration x =0.2 to x = 0.4, which indicates low chemical pressure induced by doping. Increased concentration of 'Sb' produces doping induced structural distortion from perovskite phase to Brownmillerite phase which is confirmed from the disappearance of characteristic perovskite at high intense 112 peak, which is very well reflected in the tolerance factor(t), average ionic radii $< r_A >$ and size variance factor σ^2 . Low sintering temperature

favors large change in resistance ratio, increased T_{MI} , liquid-phase grain growth and flux wrapping of 'Sb' yields smaller sized crystallites. Also, increased T_{MI} is limited only to the structural stability.

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