Percolative transition on ferromagnetic insulator manganites: Uncorrelated to correlated polaron clusters

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We report an atomic-scale study on the ferromagnetic insulator manganite LaMnO$_3$$_{1.2}$ using $\gamma$-$\gamma$ perturbed angular correlation spectroscopy. Data analysis reveals a nanoscopic transition from an undistorted to a Jahn-Teller (JT) distorted local environment upon cooling. The percolation thresholds of the two local environments enclose a macroscopic structural transition (rhombohedral-orthorhombic). Two distinct regimes of JT distortions were found: a high-temperature regime where uncorrelated polaron clusters with severe distortions of the Mn$^{4+}$O$_6$ octahedra survive up to $T=800$ K and a low-temperature regime where correlated regions have a weaker JT-distorted symmetry.

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Intense experimental and theoretical work has been devoted to manganite systems due to their colossal magnetoresistance (CMR), polaron dynamics, and charge-orbital ordering phenomena. The undoped manganites (AMnO$_3$ where A is a trivalent ion of La, Pr, ...) typically show antiferromagnetic insulator behavior and cooperative Jahn-Teller (JT) distortion of MnO$_6$ octahedra. Oxygen excess or the presence of divalent ions at A sites reduce the static JT distortion by the creation of Mn$^{4+}$ ions. This effect favors the ferromagnetic interaction via dynamic electron transfer between Mn$^{3+}$ and Mn$^{4+}$, the so-called double-exchange (DE) interaction. Although DE interaction explains qualitatively the CMR, it does not fully account for the large resistivity of the perovskite A (La) site is specially appropriate to detect lattice distortions in the surrounding MnO$_6$ octahedra, because slight changes in the charge geometry will significantly alter the electric field gradient (EFG) parameters.

The $^{111}\text{m}\text{Cd}$ probes decay to $^{111}\text{m}\text{Cd}$ through an intermediate state by the emission of two consecutive $\gamma$ rays. The half-life for the $^{111}\text{m}\text{Cd}$ isomeric state is $T_{1/2}=48$ min, while...
for the intermediate state is $T_{1/2} = 84$ ns. The angular correlation between the two $\gamma$ rays can be perturbed by both the EFG and the magnetic hyperfine field (MHF). These fields respectively couple to the nuclear electric quadrupole ($Q$) and the magnetic dipole ($\mu$) moments of the intermediate nuclear state. The Hamiltonian for such static interactions, in the proper reference frame of the EFG tensor $V_{ij}$ with $|V_{zz}| \gg |V_{yy}| \gg |V_{xx}|$, reads

$$\mathcal{H} = \frac{\hbar}{6} \omega_0 \left[ 3I^2 - I(I+1) + \frac{1}{2} \eta (I_z^2 + I_x^2) \right] + \vec{\mu} \cdot \vec{B}_{hf},$$

where $\omega_0 = 3eQV_{zz}/[2(2I-1)\hbar]$ is the fundamental precession frequency, $I$ represents the nuclear spin of the probe intermediate state ($I=5/2$ for $^{111}\text{Cd}$), $\eta = (V_{xx} - V_{yy})/V_{zz}$ is the EFG asymmetry parameter, and $\vec{B}_{hf}$ is the magnetic hyperfine field.\textsuperscript{16} The perturbation of the $\gamma$-$\gamma$ directional correlation is described by the experimental $R(t)$ function, where $t$ is the time spent by the nucleus in the $^{111}\text{Cd}$ intermediate state. For a hyperfine interaction, $R(t)$ may be expanded as $R(t) = \Sigma A_{kk} G_{kk}(t)$ with $A_{kk}$ being the angular correlation coefficients. The perturbation factor $G_{kk}(t)$ is the signature of the fields interacting with the probes: MHF and an EFG in the ferromagnetic phase and EFG alone for $T > T_c$. Below $T_c$, in the presence of the two fields, we apply combined interaction theory to obtain the MHF and EFG parameters. Above $T_c$, on the other hand, $G_{kk}(t)$ may be expressed as\textsuperscript{16}

$$G_{kk}(t) = S_{k_0} + \sum_n S_{k_n} \cos(\omega_n t) e^{-\omega_n \Delta \hbar}$$

considering only pure electric quadrupole interactions. The frequencies $\omega_n$ and amplitudes $S_{k_n}$ are determined by the $\mathcal{H}$ diagonalization. For spin $I=5/2$, three frequencies are observable that are functions of $\omega_0$ and $\eta$.\textsuperscript{17} The exponential term in Eq. (2) accounts for an attenuation of the $R(t)$ function that appears in all spectra. This effect is due to randomly distributed intrinsic vacancies and defects that produce a Lorentzian distribution of static EFGs with central value $\omega_0$ and relative width $\delta$. Independently, in manganites, short-range charge diffusion coupled to lattice distortions (polarons) can lead to EFG fluctuations. These fluctuations contribute to further attenuate $R(t)$ when their time scale is comparable to the lifetime of the PAC probe intermediate state. When the characteristic fluctuation time ($\tau$) is shorter than the nuclear spin precession time ($2\pi/\omega_0$), the $R(t)$ function can be satisfactorily approximated by a single exponential damping term $e^{-\lambda \tau}$ multiplying the static expression (2) with $\lambda \approx \omega_0^2 \tau$.\textsuperscript{18}

Some experimental $R(t)$ curves are displayed in Fig. 1 for the compound with $\Delta=0.12$. We find in the temperature range from 10 to 776 K the coexistence of three main local environments ($u, d, r$), i.e., three fractions of probes ($f_u, f_d, f_r$) interacting with different local EFG distributions. The environment $r$ is detected by a low residual fraction of the Cd probes (5%), which is temperature independent. Its EFG parameters are approximately $V_{zz} = 102$ V/Å$^2$ and $\eta_r = 0.9$ at room temperature. This highly axial asymmetric EFG might be related to probes located at the vicinity of Mn/La vacancies and/or other defects. Actually, assuming that the positions of the vacancies are not correlated, the probability that a Cd siting in a La vacancy has in its surroundings a Mn or next shell La vacancy is roughly 2%.

In Fig. 2, the temperature dependence of the EFG asymmetry parameter $\eta$ (top) and principal component $V_{zz}$ (bottom) for the $u$ and $d$ environments is displayed. For comparison, the EFG parameters found in $\Delta=0.08$ and $\Delta=0$ samples are also included in the same figure. The $u$ environment that is dominant at high $T$ shows an almost axially symmetric

**FIG. 1.** Representative $R(t)$ experimental functions and the correspondent fits for LaMnO$_3$.\textsuperscript{12} Corresponding Fourier transforms are displayed on the right side.

**FIG. 2.** Asymmetry parameter $\eta$ (top) and EFG principal component $V_{zz}$ (bottom) for LaMnO$_3$ as a function of $T$. EFG parameters for $\Delta=0.08$ and $\Delta=0$ are also shown. Inset: $T$ dependence of the MHF for the $d$ environment.
The temperature dependence of the probe volume fractions $f_u$, $f_d$, and $f_r$. Triangles: orthorhombic phase percentage from x-ray diffraction. The shaded region is limited by the temperatures where the percolation thresholds occur and the dashed horizontal line stands for the percolation threshold. Inset: log-log plot of $(f_u-f_u^0)$ vs $(T-T_s)$.

EFG ($\eta_\mu = 0$). This value characterizes an EFG with an axis of threefold or higher rotational symmetry, which is compatible with the rhombohedral lattice structure observed at high temperatures. The MnO$_6$ octahedra in the $R$ structure are constrained by symmetry to be JT undistorted (equal Mn-O bond lengths), thus we will name this local environment undistorted. In contrast, the $d$ (distorted) environment is characterized by a weaker $V_{zz}$ (Ref. 19) and highly axial asymmetric EFG ($\eta_\mu > 0.45$). At high $T$, the values of $\eta_\mu$ and $V_{zz}^d$ coincide with the ones observed for the undoped fully JT-distorted orthorhombic system, $\Delta = 0$ (full circles in Fig. 2). Consequently, at high temperatures, the $d$ local environment must be characterized by a distortion involving several (minimum eight) Mn$^{3+}$O$_6$ octahedra similar to the collective JT-distorted lattice of the orbital-ordered LaMnO$_3$. Lowering $T$ below 321 K, the asymmetry parameter $\eta_\mu$ decreases stabilizing at a value close to that observed for the $\Delta = 0.08$ sample (solid stars in Fig. 2). This behavior suggests that the JT distortions are weakening till they reach a similar degree as in the $\Delta = 0.08$ sample. The EFG principal components $V_{zz}^u$ and $V_{zz}^d$ slightly increase with decreasing temperature. This is a typical feature of perovskite and related systems. Below $T^*_u = 145$ K both $d$ and $u$ local environments experience increasing magnetic hyperfine fields upon decreasing temperature (inset of Fig. 2), presenting at 10 K values of $B_{hf}^d = 3.8(2)T$ and $B_{hf}^u = 4.0(3)T$ compatible with a full ferromagnetic environment of the surrounding Mn ions.

Further insight into the behavior of $d$ and $u$ environments may be achieved by studying the $T$ dependence of the volume fractions $f_u$ and $f_d$. As may be seen in Fig. 3, the $u$ environment is dominant at very high temperatures ($f_u = 86\%$ at $T = 776$ K), though $d$ regions survive up to that $T$ ($f_d = 9\%$). This confirms the high stability of the inhomogeneous phase-segregated state. Our data, at high $T$, are compatible with a scenario where random distributed JT-distorted nanoclusters are embedded in an undistorted matrix as predicted by Refs. 10 and 24, setting a lower bound for the temperature where the polaron clusters start to form.

When the temperature changes, $f_u$ (symmetrically $f_d$) suffers a smooth variation leading from an undistorted to a JT-distorted dominant microscopic environment. If we assume that this variation is a continuous phase transition, the order parameter would be $f_u - f_u^0$ and must follow a power law behavior $f_u - f_u^0 \sim (T-T_s)^\beta$ when the critical temperature $T_s$ is approached from above. To check this possibility, we display $f_u$ in a log-log plot in Fig. 3 (inset). The data adjust pretty well to a power law with $T_s = 170 \pm 10$ K (relatively close to $T_c$) and $\beta = 0.42 \pm 0.02$. Associated with the transition, there must also exist a correlation length, a correlation of the $d$ spatial distribution, that must diverge at $T_c$. As may be seen in Fig. 2, when $T$ decreases $\eta_\mu$ starts to fall as the $d$ component percolates [at $f_d = 31.16\%$ (Ref. 26)] and only stabilizes around $T_c$. Macroscopically, on the other hand, x-ray measurements detect a structural transition $(R-O)$ that lies exactly between the temperatures corresponding to the percolation thresholds of the two main nanoscopic components. These are precisely the temperatures in which the minority invading cluster suffers a sudden size divergence becoming macroscopically observable.

The temperature dependence of the attenuation of $R(t)$ provides additional information about the dynamics of the $u$ and $d$ environments. A complete sketch of the dynamic and static attenuation for $R(t)$ in both environments is depicted in Fig. 4. The best fit to the $R(t)$ spectra discards the presence of time-dependent interactions for the $u$ environment $(\delta_u = 0\%$ independently of $T$ and $\lambda_u = 0)$. Thus, in all temperature ranges, the charge transfer between Mn$^{3+}$ and Mn$^{4+}$ (activated hopping) in this environment should occur with a frequency higher than we can probe. For the $d$ environment, on the other side, the best fits were obtained admitting a fluctuating EFG $(\lambda_d \neq 0$ and $\delta_d = 2\%)$ in the temperature region spanning from $T = 266$ to $T = 350$ K. Notice that these time-dependent effects cannot be attributed to Cd/O and/or defects diffusion because they would be detected in both...
fractions. The temperature dependence of the dynamic attenuation parameter, \( \lambda_d \), allows us to estimate an activation energy \( E_a \). This energy is obtained from \( \lambda_d = \lambda_{\text{d}} e^{E_a/kT} \), and was found to be \( E_a \approx 0.31 \text{ eV} \) (see the inset of Fig. 4), close to the polaron binding energy reported in the literature for low-doped manganites.\(^{27,28}\) We identify such EFG fluctuations with polaron diffusion related to charge (hole) transport. The EFG fluctuation time \( (\tau) \) can be estimated from the maximum of \( \lambda_d(T) \).\(^{29}\) Considering that a carrier (hole) can hop to any of the eight octahedra around a La site (eight possible EFG states), we find \( \tau \approx 0.5 \mu \text{s} \) at \( T = 266 \text{ K} \) corresponding to ultraslow polaron diffusion. Similar polaron residence times have been recently reported in Ref. 22, although the \( E_a \) measured there was smaller possibly due to the intense magnetic field (7 T) needed to perform the NMR measurements. The competition of the distinct dynamics of the \( u \) (fast hopping) and \( d \) (related to polaronic conduction) environments is responsible for the macroscopic ferromagnetic insulator behavior observed in these systems.\(^{25}\) Below \( T_{c_1} \), both local environments become ferromagnetic and a phase coexistence between metallic \( (u) \) and insulator \( (d) \) regions exists. However, the majority fraction \( (d) \) is characterized by the ultraslow diffusion of charge carriers imposing an overall insulator behavior.

In conclusion, we report an extensive nanoscopic analysis on the prototypical FMI manganite \( \text{LaMnO}_3 \) using \( \gamma-\gamma \) PAC spectroscopy. The undeniable separation of two distinct local environments, consistent with a phase coexistence scenario, allowed us to study the detailed structure, stability, and evolution of the JT-polaron clusters in manganites. We find that uncorrelated JT-polaron clusters survive up to a remarkably high \( T \) within the \( R \) crystallographic phase. Our results define a lower bound for the polaron cluster formation temperature \( (T_{\text{clusters}} \approx 776 \text{ K}) \). When \( T \) is lowered, PAC measurements reveal a smooth increase in the JT-distorted nanoscopic environment density until it becomes dominant. This change may be described as a continuous phase transition with the transition temperature \( T_{c_1} \) marking the point where the polaron cluster correlation length diverges. In parallel, a macroscopically \( O-R \) structural transition takes place as a consequence of the macroscopic changes; actually it occurs between the percolation thresholds of the two main local components.

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