Local polarization in oxygen-deficient LaMnO$_3$ induced by charge localization in the Jahn-Teller distorted structure

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The functional properties of transition metal perovskite oxides are known to result from a complex interplay of magnetism, polarization, strain, and stoichiometry. Here, we show that for materials with a cooperative Jahn-Teller distortion, such as LaMnO$_3$ (LMO), the orbital order can also couple to the defect chemistry and induce novel material properties. At low temperatures, LMO exhibits a strong Jahn-Teller distortion that splits the $e_g$ orbitals of the high-spin Mn$^{3+}$ ions and leads to alternating long, short, and intermediate Mn-O bonds. Our DFT+U calculations show that, as a result of this orbital order, the charge localization in LMO upon oxygen vacancy formation differs from other manganites, such as SrMnO$_3$, where the two extra electrons reduce the two Mn sites adjacent to the vacancy. In LMO, relaxations around the defect depend on which type of Mn-O bond is broken, affecting the $d$-orbital energies and leading to asymmetric and hence polar excess-electron localization with respect to the vacancy. Moreover, we show that the Mn-O bond lengths, the orbital order, and consequently the charge localization and polarity are tunable via strain.

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The family of doped (La,Ca/Sr)MnO$_3$ perovskite manganites has attracted great interest due to its very rich phase diagram and its unusual functional properties (colossal magnetoresistance [1–5], photoinduced infrared absorption [6], and efficient hole conductivity [7]) with strong potential for new applications in the fields of electronics, spintronics, and energy conversion. The orbital order plays a significant role in determining these properties and is strongly coupled with the other electronic, structural, and spin degrees of freedom [8,9]. LaMnO$_3$ (LMO) as the end member of this family also exhibits interesting properties such as a pressure-induced insulator-to-metal transition [10] and a dielectric anomaly [11], that also depend on the orbital order [12,13].

Below 750 K, LMO adopts a distorted orthorhombic (Pbnm) perovskite structure [see Fig. 1(a)] with an $A$-type antiferromagnetic order ($A$-AFM, Néel temperature $T_N \approx 140$ K [14]), with Mn atoms coupled ferromagnetically in the $ac$ plane and antiferromagnetically along the $b$ axis [15–17]. The stabilization of this orbital-ordered insulating state is a consequence of the high-spin Mn$^{3+}$ ($t^3_{2g} e^1_g$) ions inducing a strong cooperative Jahn-Teller (JT) distortion that splits the $e_g$ orbitals ($d_{z^2}/d_{x^2-y^2}$ orbitals are alternately occupied within the orthorhombic $ac$ plane) and lead to alternating long and short in-plane Mn-O bonds, along with intermediate bonds along $c$.

![FIG. 1. (a) Pbnm unit cell of $A$-AFM LaMnO$_3$ (LMO). (b) Definition of the rotation and tilt angles and of the short (Mn-O$_s$), intermediate (Mn-O$_i$), and long (Mn-O$_l$) Mn-O bonds. (c) The $(2 \times 2 \times 2)$ LMO supercell used in this work with the two inequivalent $V_{0}$ positions: in the $ac$ plane (IP) and perpendicular to it (OP). (d) Definition of the Jahn-Teller distortion amplitude $Q_3$: $d_{36}$, $d_{14}$, and $d_{25}$ are distances between the corresponding O atoms. Axes $a'$, $b'$, and $c'$ are orthorhombic, while $a$, $b$, and $c$ are pseudocubic.](https://doi.org/10.1103/PhysRevResearch.2.042040)

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the $b$ axis [see Fig. 1(b)]. There are two symmetry-distinct oxygen atoms in this structure: the in-plane O (OIP) in the $ac$ plane with one short and one long Mn-O bond, and the out-of-plane O (OOP) along the $b$ direction with two intermediate Mn-O bonds [see Fig. 1(c)].

Unlike most other perovskite oxides, bulk LMO can exhibit oxygen superstoichiometry (LaMnO$_{3+\delta}$), which is accommodated through cation vacancies rather than oxygen interstitials [18–21]. LaMnO$_{3+\delta}$ has an extremely complex magnetic phase diagram exhibiting paramagnetic, ferromagnetic (FM), and canted AFM order as well as a spin-glass state as a function of temperature and O superstoichiometry [19]. Topotactic low-temperature reactions [22,23] were shown to lead to oxygen substoichiometric LaMnO$_{3-t}$ with oxygen vacancies ($V_o$), where for $0.00 \leq \delta \leq 0.20$, the $A$-AFM order persists below $T_N \approx 140$ K [23]. The FM behavior often observed in LMO thin films [24–31] was explained by Mn$^{3+}$–Mn$^{4+}$ double exchange due to the cation deficiency generally observed in these films. It was suggested that, below a critical thickness of 6 unit cells, LMO thin films become AFM [32], but several authors also reported thicker films with bulklike AFM behavior obtained either through growth [33,34] or high-temperature annealing [35] under conditions favoring the formation of $V_o$.

In this Rapid Communication we investigate strained stoichiometric and oxygen-deficient LMO by density functional theory (DFT) calculations within the QUANTUM ESPRESSO package [36,37] (see Supplemental Material [38] Sec. S1 for details). At the PBCsol+U [39,40] level of theory with a self-consistently computed Hubbard $U$ for Mn-3d states [41,42], we correctly describe the JT distorted structure, which is crucial to accurately predict magnetic and electronic properties [13,43–45]. As can be seen from Supplemental Material [38] Table S3, not only are the computed lattice parameters in good agreement with experiment (relative error below 1%), but we also obtain a good description of the JT distortion (quantified via the magnitude of the $Q_3$ mode [see Fig. 1(d)]) and reproduce the Mn-O bond lengths to within 2% of experiment. We find the orbitally ordered insulting (see Supplemental Material [38] Fig. S1) state with $d_{x^2-y^2}$ orbital being alternately occupied within the orthorhombic $ac$ plane, which is consistent with the alternating Mn-O$_a$ and Mn-O$_b$ bonds. The direct gap of about 1 eV is in good agreement with experiments, while the indirect gap of 0.69 eV is lower than reported in previous theoretical studies [13,43], but still larger than experiment [46].

Under experimentally relevant conditions, i.e., in the oxygen-poor limit and for a Fermi energy close to the experimental indirect band gap of 0.24 eV [46], the singly charged oxygen vacancy ($V_o^-$ in Kröger-Vink notation [47]) is the favored charge state for $V_o$ in bulk LMO (see Fig. 2), the neutral defect ($V_o^{0\infty}$) being less stable by about 0.2 eV. However, already for 0% strain, corresponding to a cubic substrate with the same $ac$ area as bulk LMO, epitaxial strain renders $V_o^{0\infty}$ and $V_o^-$ formation energies nearly degenerate, the neutral defect being further stabilized under compressive strain, as we will discuss in more detail later. For these reasons and being mainly interested in LaMnO$_{3-t}$ thin film properties, we consider here only neutral oxygen vacancies, for simplicity referring to them as $V_o$ in the following. Singly ($V_o^+$) and doubly charged ($V_o^{+2}$) vacancies are described in Supplemental Material [38] Secs. S5 and S6.

Upon neutral oxygen-vacancy formation, the JT distorted structure leads to excess charge localization different from that observed, for example, in CaMnO$_3$ and SrMnO$_3$ [42,48], where the two excess electrons reduce Mn in nearest-neighbor (NN) positions to the defect. Figure 3(c) schematically illustrates the changes in the Mn-O framework induced by the relaxations around an out-of-plane oxygen vacancy ($V_{O_{\text{OIP}}}$). After breaking the two equivalent Mn-O$_b$ bonds along the $b$ axis, the structure relaxes by shortening the Mn-O$_b$ bonds along $c$ of the two undercoordinated NN Mn atoms [Mn$_1$ and Mn$_5$ in Fig. 3(c)] and significantly expanding the former Mn-O$_a$ bonds of the next-nearest-neighbor (NNN) Mn atoms [Mn$_4$ and Mn$_8$ in Fig. 3(c)] along $c$. As a result of these relaxations, the energy of the $d_{x^2}$ orbital increases for Mn$_1$ and Mn$_5$ and decreases for Mn$_4$ and Mn$_8$ [see Fig. 3(d)], localizing the two excess electrons resulting from $V_o$ formation on these orbitals and reducing the NNN Mn$_4$ and Mn$_8$ atoms [see Fig. 3(c)].

The case of an in-plane defect ($V_{O_{\text{OIP}}}$), is more complex since both a Mn$_1$-O$_a$ and a Mn$_5$-O$_b$ bond are broken. As can be seen from Fig. 3(e), the structural relaxations upon $V_{O_{\text{OIP}}}$ formation primarily involve one of the two NN Mn ions [Mn$_6$ and Mn$_7$ in Fig. 3(e)] where a $V_{O_{\text{OIP}}}$ created along the $a$ axis is shown. The NN Mn atom where a Mn$_1$-O$_a$ bond was broken [Mn$_5$ in Fig. 3(e)] shortens its remaining Mn-O$_a$ bonds along the $c$ and $b$ axis, resulting in elongated Mn-O$_a$ bonds for NNN Mn sites around Mn$_5$: Mn$_1$ along $b$ and Mn$_8$ along $c$ [see Fig. 3(e)]. These bond-length changes are also reflected in the $d_{x^2}$ and $d_{z^2}$ orbital energies of Mn$_1$ and Mn$_5$, respectively, that are increased due to breaking the Mn$_3$-O$_b$–Mn$_7$ bond, while the $d_{x^2}$ orbital of Mn$_4$ and Mn$_8$, respectively, are stabilized and filled by the excess electrons. Therefore, for the $V_{O_{\text{OIP}}}$, the reduction happens on NNN Mn sites [see Figs. 3(e) and 3(f)].

![FIG. 2. Oxygen vacancy formation energy ($E_f$) in bulk LMO and for 0% epitaxial strain under oxygen-poor conditions ($\Delta\mu_o = -2.21$ eV) and in different charge states computed as a function of the Fermi energy ($E_F$) with respect to the valence band maximum of stoichiometric LMO. Only the most stable charge state is reported as indicated next to each line. The vertical dashed lines indicate $E_f$ values corresponding to the experimental and theoretical direct and indirect band gaps (see Supplemental Material [38] Table S3).](image_url)
Furthermore, the average lengths of the Mn-O shown in panels (d) and (f) are not noticeably affected by the defect. For both vacancies, the reduced sites show almost equivalent Mn-O bond lengths (between 2.03 and 2.17 Å), in good agreement with the structural parameters reported by Has-teen et al. [23] for LaMnO$_2$$_{0.8}$ that has a V$_O$ concentration very close to our calculations. Previous DFT calculations of oxygen vacancies in the high-temperature non-JT distorted G-AFM phase [49,50], reported reduction of the two NN Mn atoms, supporting that our findings stem from the JT distortion and orbital order in the A-AFM phase.

Interestingly, when the reduction does not take place on the Mn sites adjacent to the vacancy, the localization of the excess electrons with respect to the defect can be asymmetric and hence polar. We quantify this polarization considering the magnitude and orientation of the Mn$^2+$/V$^{2+\ast}$ dipoles, identifying the reduced Mn sites based on structural changes and oxidation states calculated according to Ref. [51]. More details can be found in Supplemental Material [38] Sec. S1. For V$_{OIP}$ a large polarization of 24.38 μC/cm$^2$ in the ac plane is observed while the out-of-plane contribution is zero, in line with the reduced Mn$_4$ and Mn$_8$, being symmetrically arranged with respect to the V$_O$ along the b axis [Fig. 3(c)]. A polarization with in- and out-of-plane components of 17.08 and 10.20 μC/cm$^2$, respectively, is computed, instead, for the V$_{OIP}$, in agreement with the larger asymmetry of the charge localization along the b axis. We note here that the local polarizations resulting from this novel mechanism are of similar magnitude as in conventional ferroelectrics such as BaTiO$_3$ as discussed in more detail in Supplemental Material [38] Sec. S1. We emphasize that the JT distortion is pivotal to obtain local polarizations of such magnitude. In non-JT distorted materials, vacancy formation on a Wyckoff position without an associated (local) inversion center can also result in a local dipole, the magnitude of which is, however, much smaller than in our case. With our model, we estimate a local polarization of only 3.6 μC/cm$^2$, assuming that the excess electrons localize on two Mn sites adjacent to the vacancy as in the non-JT-distorted G-AFM phase of LMO [49,50]. The key ingredient of our novel mechanisms is therefore the JT orbital-order induced excess charge localization on Mn atoms far from the oxygen vacancy, which together with the asymmetric charge arrangement yields the observed large local dipoles.

For both vacancies, the reduced sites show almost equivalent Mn-O bond lengths (between 2.03 and 2.17 Å), in line with the fact that no JT distortion is expected for a Mn$^{2+}$ ion (Mn$^+$ in Kröger-Vink notation [47]) in an octahedral crystal field. Furthermore, the average lengths of the Mn-O$_2$, Mn- O$_i$ and Mn-O$_j$ bonds are 1.98, 2.17, and 2.20 Å in good agreement with the structural parameters reported by Has-teen et al. [23] for LaMnO$_2$$_{0.8}$ that has a V$_O$ concentration very close to our calculations. Previous DFT calculations of oxygen vacancies in the high-temperature non-JT distorted A-AFM phase [49,50], reported reduction of the two NN Mn atoms, supporting that our findings stem from the JT distortion and orbital order in the A-AFM phase.

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Different energetically equivalent patterns to accommodate the charges are possible. This is particularly apparent for the sketch in Fig. 3(c), where the charges are shown to localize on Mn sites along the positive c axis. Accommodation of charges on the Mn$_4$ and Mn$_8$ sites along the negative c axis, corresponds to an energetically equivalent situation, in which the local polarization, however, points in the opposite direction. For the V$_{OIP}$ there are two such states, while for the V$_{OIP}$ we expect four states, since equivalent positions exist along $b$ for Mn$_1$ as well as along $c$ for Mn$_8$. Given the high defect concentration achievable in experiment [22,23] it is conceivable that these local dipoles will couple, leading to an ordered ferroelectric phase. As such we expect the possibility of a switchable polarization that could lead to defect-induced ferroelectricity in reduced LMO.

Based on the preceding discussion, it is reasonable to expect that any external parameter affecting the Mn-O bond lengths and/or the orbital order will have a strong impact also on the charge localization and the resulting polarization. As one such external parameter, we investigate the effect of strain, both isostatic and biaxial, the latter induced, for example, by heteroepitaxial growth on a substrate with a different lattice parameter. As shown in Supplemental Material [38] Figs. S2 and S3, both types of strain are
accommodated in stoichiometric LMO by changes in octahedral tilt angles and bond lengths [52]. While Mn-O, and Mn-Oi bonds are relatively insensitive to strain, the Mn-Ot bonds are more strongly affected especially under tensile strain. This results in an increase of the JT mode amplitude $Q_3$ under tensile strain, while compressive strain reduces the $Q_3$ amplitude.

In Fig. 4 we show results for the formation energies, excess charge localization, and polarization under epitaxial strain. Results for isostatic strain are shown in Supplemental Material [38] Fig. S4. The localization of the excess charge away from the NN sites results in a strain dependence of the formation energy different from chemical expansion arguments [48,53,54] as discussed in Supplemental Material [38] Secs. S4–S6. While compressive strains favor neutral vacancies with slightly smaller formation energies for $V_{OOP}$ than $V_{OP}$, tensile strain results in singly charged $V_{OXP}$ and a mixture of singly and doubly charged $V_{OXP}$, the latter not contributing to the local polarization due to the absence of excess charge. This implies that to obtain large polarizations, compressive strain seems more promising and that under these conditions neutral $V_{OOP}$ will be predominantly stable, followed by neutral $V_{OXP}$.

Epitaxial strain leads to changes in the excess-charge localization as a function of the strain magnitude. Interestingly, for both neutral $V_{OOP}$ [Fig. 4(a)] and $V_{OP}$ [Fig. 4(b)], imposing the constraint of a cubic substrate with equal $a$ and $c$ lattice parameters that result in the same $ac$ area as bulk LMO (0% strain) induces a more asymmetric charge localization on NNN sites compared to the bulk. This excess-charge localization is maintained under compressive strain for both defects, with a slight variation for $-1$ and $-2\%$ strain for $V_{OXP}$. This variation shows that energetically nearly degenerate excess-charge localization patterns exist. Under tensile strain, the excess charge localizes more symmetrically, for $V_{OXP}$ first assuming the one of the bulk for $1$ and $2\%$ strain, before localizing one (for $3\%$ strain) and then two (for $4\%$ strain) of the excess electrons on NN Mn sites. For $V_{OP}$ the localization remains on NNN sites even for $4\%$ tensile strain.

These strain-induced changes in charge localization affect also the local polarization, which depends on the relative arrangement of the two $Mn'\cdot V_{O}^\bullet$ dipoles. For $V_{OXP}$ the OP component of the polarization is almost zero throughout the whole investigated range of strains. The IP component reaches sizable values that depend on the excess charge localization pattern. For $V_{OP}$ the OP component is nonzero and fairly constant, while the IP component is significantly reduced for $+3\%$ and larger strain due to the change of the relative $Mn'$ orientation in the $ac$ plane. Similar results for less stable $V_{O}^\bullet$ are discussed in Supplemental Material [38] Sec. S5.

In summary, we have shown that, in the case of Jahn-Teller distorted perovskite oxides, the orbital order is an additional parameter to take into account during defect-based design of functional material properties. As we have shown for the example of oxygen vacancies in LaMnO$_3$, unexpected excess-charge localization results from the orbital order and can lead to the emergence of local polarization. The charge localization and hence the polarization magnitude are tunable by strain, suggesting the possibility to engineer ferroelectricity in LaMnO$_3$ when local dipoles couple for high enough, yet experimentally attainable, defect concentrations.
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