Role of orbital selectivity on crystal structures and electronic states in BiMnO₃ and LaMnO₃ perovskites

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Correlated oxides, such as BiMnO3 and LaMnO3, show a complex interplay of electronic correlations and crystal structure exhibiting multiple first-order phase transitions, some without a clear order parameter. The quantitative theoretical description of this temperature-dependent electronic-structural interplay in the vicinity of a Mott transition is still a challenge. Here we address this issue by simultaneously considering both structural and electronic degrees of freedom, within a self-consistent density functional theory with embedded dynamical mean field theory. Our results show the existence of novel electronic states characterized by the coexistence of insulating, semimetallic, and metallic orbitals. This state is in agreement with resonant x-ray scattering. We also show that electronic entropy plays a decisive role in both electronic and structural phase transitions. By selfconsistent determination of both the electronic state and the corresponding crystal structure, we show that the temperature evolution of these phases can be quantitatively explained from first principles, thus demonstrating the predictive power of the theoretical method for both the structural and the electronic properties.

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I. INTRODUCTION

Manganese oxides with perovskite-type structure host a variety of interesting phenomena, among others the colossal magnetoresistance [1,2], insulator-to-metal transition [3], ferroelectricity coexisting with long-range magnetic order [4], and orbital and charge ordering [5]. The theoretical description of these functionalities is challenging, as the electronic correlations play an essential role, and the interplay between the electronic, orbital, and magnetic degrees of freedom with the crystal structure makes these materials extremely sensitive to small changes in external parameters, such as electric voltage, magnetic field, and temperature. However, this high sensitivity is a useful property, which is explored in the development of new electronic devices.

The theoretical methods for the ab initio description of strong correlations in solids have been recently developed, and among them the density functional theory with embedded dynamical mean field theory (DFT + eDMFT) [6–8] has demonstrated predictive power in many classes of correlated materials. For example, it was predicted that the magnetic moment in plutonium metal is screened at 800 K with valence fluctuations [9], which was subsequently verified by neutron scattering experiment [10]. An order of magnitude enhanced electron-phonon coupling in the FeSe superconductor as compared to the DFT result was predicted in Ref. [11]. and subsequently verified by pioneering experiment using a combination of time-resolved photoemission spectroscopy coupled with x-ray diffraction [12]. Similarly, predictions of large fluctuating moments in iron superconductors and their energy distribution [13] have been later verified by neutron scattering [14]. The prediction of the isostructural metal-

insulator transition under pressure in FeO has been confirmed in Ref. [15]. A novel type of long-ranged order with very high multipole ordering was proposed to explain the hidden order in URu₂Si₂ [16], which was confirmed by Raman scattering experiment [17]. Finally, structural relaxations by the eDMFT method have been recently developed [18], and it was demonstrated on the example of NdNiO₃ [19] and Fe₂Mo₃O₈ [20] that they can predict small changes of crystal structures near the Mott metal-insulator transition, the region which is challenging for other ab initio methods. In this paper we study the AMnO₃ class of materials with perovskite structure type to understand the role that correlations play in generating their rich phase diagrams [23–26]. We compare and contrast LaMnO₃ with BiMnO₃, which have attracted considerable interest due to the giant magnetoresistance phenomena in doped LaMnO₃ [1] and the potential multiferroic property in BiMnO₃ thin films [27,28].

The high-temperature (HT) phase (T > 800 K) of both LaMnO₃ and BiMnO₃ is a perovskite-type structure with orthorhombic symmetry [29-31] [see Fig. 1(a)] with similar bad-metallic conductivity [29,32-34]. At room temperature (RT) both compounds are insulating [29,35] and in LaMnO₃ the orthorhombic distortions increase with decreasing temperature but the symmetry remains orthorhombic [31], while in BiMnO₃ the structure distorts to monoclinic symmetry [36–38]. An abrupt change of resistivity is observed in both compounds around the transition, with the high-temperature phase exhibiting nearly temperature-independent resistivity, characteristic of a very bad metal. The pattern of structural distortions in both compounds is compatible with the longrange orbital ordering (OO) [31,36,39,40], which has been validated by resonant x-ray scattering (RXS) experiments

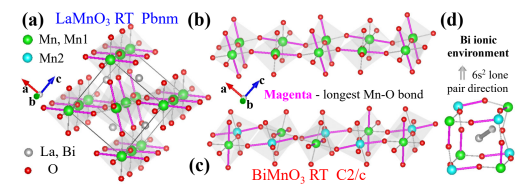


FIG. 1. Room temperature (RT) structural properties for LaMnO₃ and BiMnO₃. (a) Schematic view of the crystal structure for LaMnO₃. The longest Mn-O bond lengths are colored by magenta. (b) The octahedra distortion pattern in the (101) plane. (c) Distortions in BiMnO₃ in which Mn₁ ions show similar pattern to that in LaMnO₃, while in Mn₂ ions the long-bond distortions are confined to (101) plane. (d) Schematic view of Bi environment within a perovskite-like crystal structure; the gray arrow, pointing toward the face of the Mn₂O₆ octahedra, shows the direction of the $6s^2$ lone-pair electronic cloud within the Bi ionic environment [21]. The presence of the $6s^2$ lone pair is responsible for the appearance of two inequivalent Mn sites. Drawings were produced by VESTA [22].

[41–43] [see Figs. 1(b) and 1(c)]. The theoretical description of this Mott-type transition is most often described within the phenomenological models of the cooperative Jahn-Teller (JT) effect due to the $Mn-d^4$ configuration in octahedra, leaving a single electron in the e_g shell [21,24,32,44–47]. However, the two e_g orbitals are not degenerate in the high-temperature metallic phase, therefore the conventional JT mechanism does not apply, and better explanation of the transition is called for. Moreover, experimentally it was found that BiMnO3 undergoes two phase transitions: A structural phase transition from orthorhombic to monoclinic symmetry takes place at T = 768 K [29], followed by another isostructural phase transition at T = 474 K [29,36,48]. While orbital order is symmetry-allowed at the first transition, RXS experiments showed that the situation is more complicated [43]. There is a sharp mean-field type upturn of the forbidden Bragg peak intensity at T = 768 K followed by the saturation regime. Below the second transition (T = 474 K), there is a second sharp upturn of the intensity, signaling that the long-range orbital order is well established only below the second phase transition (T < 474). The forbidden Bragg peak intensity in the intermediate phase (474 K < T < 768 K) saturates to ≈0.6 of the room temperature intensity, which suggests that the intermediate phase has some type of partial order, but the nature of this partial order remains unknown. Both transitions are of first order, but the second isostructural transition does not have a clear order parameter.

The Mott transition and the Jahn-Teller distortion in LaMnO₃ have been previously studied by the dynamical mean field method using the minimal Hubbard model for the e_g electron, while the t_{2g} electrons were treated semiclassically, as a classical spin S with random orientation [49,50]. In this approach the effect of covalency is ignored, which was shown by x-ray absorption spectroscopy (XAS) to be important for interpretation of the XAS data [51]. Nevertheless, the metal-insulator transition was well captured by this minimal-model approach [49,50]. In Ref. [49] the transition temperature to the disordered state was somewhat underestimated, because the feedback effect of electronic correlations on crystal structure was ignored. This was partially corrected

in Ref. [50] by allowing JT distortion, although the full structural relaxation was not attempted. Nevertheless a much better estimate for the transition temperature was reported, despite the fact that the electronic entropy was ignored in the calculation of energy. On the other hand, the complex phase diagram of BiMnO₃ has not been theoretically addressed before.

II. METHODS

To understand the unusual Mott transition coupled with the partial orbital order from first principles, we use the DFT + eDMFT method [6,8]. Here the downfolding to Hubbard-type models is avoided, and the dynamic correlations are added only to the very localized 3d orbitals, which is the solution of the Schrödinger equation inside a sphere centered on the Mn ion (radius 1.93 bohrs). The double counting between the DFT and DMFT correlations is treated within a nominal scheme, where the double counting is given by $U(n^0 - 1) - 1$, with n^0 being the nominal occupancy of the d orbital [52]. The Coulomb repulsion on these quasiatomic orbitals used in this work has the Slater form in its fully rotational invariant form with its values estimated by constrained eDMFT to be U = 10 eV, J = 1.18 eV.

All crystal structures, except the high-temperature orthorhombic structure of BiMnO₃, were experimentally determined [30,31,36], and we checked that they agree very well with the theoretically determined optimized crystal structures (see Ref. [53]). We note that the force in our approach is calculated from the derivative of the free energy, rather than total energy [18]; therefore it contains important effects of electronic entropy at finite temperature, which we find to be essential to describe the orbitally disordered state in the orthorhombic structure, with finite but small orthorhombic distortions. We also note that the high-temperature orthorhombic BiMnO₃ crystal structure has not been determined before, and here is predicted for the first time (see Ref. [54]). It would be desirable to confirm it experimentally.

TABLE I. Occupancy of the Mn orbitals $(n_{e_{gl}}, n_{e_{g2}})$, where $|e_{gl}| \approx |3z^2 - 1i$ and $|e_{g2}| \approx |x^2 - y^2i$. Note that $n_{t_{2gl}} \approx n_{t_{2g2}} \approx n_{t_{2g3}} \approx 1$. The realized electronic states are band-Mott insulator (BMI), orbital-selective Mott (OSM) state, and site-orbital-selective Mott (SOSM) state.

LaMnO ₃	T = 300 K	T = 573 K	T = 798 K
Mn	(0.98, 0.45)	(0.97, 0.46)	(0.76, 0.66)
state	BMI	BMI	OSM
BiMnO ₃	T = 300 K	T = 550 K	T = 760 K
Mn_1	(0.98, 0.44)	(0.93, 0.47)	(0.72, 0.69)
Mn_2	(0.98, 0.44)	(0.73, 0.69)	(0.72, 0.69)
state	BMI	SOSM	OSM
	21111	200111	35111

III. RESULT AND DISCUSSION

A schematic view of the room temperature (RT) crystal structure of LaMnO₃ and BiMnO₃ compounds is shown in Fig. 1. It consist of one and two inequivalent corner-sharing MnO₆ octahedra in LaMnO₃ and BiMnO₃, respectively. The two inequivalent octahedra, denoted by Mn₁O₆ and Mn₂O₆, appear due to the lone pair of electrons derived from the Bi-6s orbital [21], whose direction is shown schematically by the gray arrow in Fig. 1(d). Note that the lone pair consist of electronic charge, occupying the space between the center of Bi and the Mn₂ ion [21], which we mark by the gray arrow.

To describe the Mn-3d orbitals, we use the local coordinate system attached to each Mn ion, in which the hybridization of each orbital with environment is minimal. As expected for octahedral environment, there are three t_{2g} and two e_g orbitals pointing toward the faces and corners of the octahedra, respectively. As shown in Fig. 1(d), the lone pairs are pointing toward the octahedron face of Mn₂ ions; thus for this ion type the t_{2g} orbitals are expected to hybridize with the lone pair strongly, while the orbitals of Mn₁ ion type should not directly feel the presence of the lone pair in BiMnO $_3$.

While the formal valence of the Mn ion is Mn^{3+} , our calculations show that due to covalency there are approximately $n_d \approx 4.4$ electrons on each Mn ion (Mn^{2.6+}), of which almost exactly three electrons are in the t_{2g} shell, and therefore $n_e \approx 1.4$. This is in good agreement with valence determined from XAS in Ref. [51] ($n_d \approx 4.5$), but it is in stark contrast with the earlier DMFT works in which the number of e_g electrons has been fixed to one electron per Mn.

The degree of the orbital order is usually measured by occupancy ratio $n_{e_g}/(n_{e_g} + n_{e_g}) \equiv (c_2)^2$, which was estimated by neutron scattering to be $c_2 \approx 0.81$ at room temperature [31]. Table I gives the temperature dependence of the occupancies, as found by the eDMFT method. We note that the maximal orbital ordering is for $n_{e_g} \approx 1$, but due to covalency n_e does not vanish in this limit; hence c_2 always remains smaller than 1. In LaMnO₃ we get $c_2 = 0.826$, in good agreement with estimates from Ref. [31]. In LaMnO₃ the orbital polarization changes very little as long as the system has sizable charge gap, while orbital polarization becomes very small in the bad-metal phase above T = 798 K. Since the two e_g orbitals are not equivalent in the high-temperature structure, there remains a small difference in the occupancy as long as the system is in the orthorhombic phase. We note that in

the earlier DMFT work [49] such agreement with experiment could not be achieved, and it was assigned to negligence of the electron-phonon coupling. In the minimal Hubbard model the insulating state must have integer occupancy; hence the constrained $n_{e_g} + n_{e_g} = 1$ leads to $c_2 \approx 1$ when polarization is large $(n_{e_{g1}} \approx 1)$. Hence we now understand that the earlier disagreement with experiment was due to negligence of Mn covalency.

In BiMnO₃ there is an intermediate phase, in which electronic occupancies on the two Mn atoms are very different. Mn₁ keeps orbital polarization close to its room temperature value, while Mn₂ loses its polarization in this temperature range. As we will see later from structural optimization, such result is hard to guess from crystal structure alone, as the Mn₂ octahedra remain quite distorted in this intermediate phase.

If Fig. 2 we show temperature evolution of the density of states (DOS) for both t_{2g} and e_g orbitals. We also display structural information of the experimental structure. Each octahedron has three different types of Mn-O bond lengths, which are displayed as red horizontal bars. We label them as short (s; $\mathbb{P}[1.9]$), medium (m; $\mathbb{P}[2.0]$), and long (l; & 2.1). We also provide distortion parameters for octahedra [22], namely, the bond distortion index $\mathbf{D} = \begin{bmatrix} 1000 & 6 \\ i=1 & |l_i - l_{av}|/(6 l_{av}) \end{bmatrix}$ and the bond angle variance $\mathbf{S} = \begin{bmatrix} 8 \\ i=1 & (\varphi_i - 90^\circ)^2/7 \end{bmatrix}$. Here l_i and φ_i are the bond distances and bond angles, respectively. l_{av} is the average bond length. Compared to Ref. [22], we use a formula for \mathbf{D} with a multiplying factor of 1000. \mathbf{D} and \mathbf{S} are zero for an ideal octahedron.

At all temperatures and in both compounds t_{2g} electrons are in the Mott insulating state, and remain gapped even in the high-temperature metallic phase (see Figs. 2 and 3). This can be verified by looking at the t_{2g} self-energy, which has a pole inside the gap even when the e_g density of states shows metallic behavior (see Ref. [55]). This state is commonly called the *orbital-selective Mott state* [56–58], and was first introduced in Ref. [56] to explain the properties of Ca_{2-x}Sr_xRuO₄ in which spin-1/2 local moments coexist with metallic transport. Such a picture of frozen high-spin t2g electrons in LaMnO₃ was postulated already in the very early works on the double-exchange model [59-61], and was also assumed in the previous DMFT calculations within the minimal two-orbital Hubbard model [49,50]. In the limit of weak correlations among e_g electrons, coexisting with the high-spin core t2g states, the DMFT problem maps to the ferromagnetic Kondo model, which was shown in Ref. [62] to explain qualitatively the giant magnetoresistance in doped LaMnO₃. Nevertheless, to the best our knowledge, direct DMFT simulations with equal dynamic treatment of both t_{2g} and e_g orbitals was lacking; hence the theoretical photoemission spectra and position of the t_{2g} states was unknown. Moreover, as we will see in BiMnO₃ the approximation of t_{2g} electrons as frozen high-spin core is not really adequate.

Figure 2(a) shows that the room temperature gap is between e_g states, in which the occupied part is strongly admixed with oxygen states. The orbital with larger occupation (e_{g1}) points along the longest Mn-O bond, while the less occupied orbital (e_{g2}) points toward the shorter bonds. The gap between e_g is not of Mott type; hence we call this state the band-Mott insulator. The peak-to peak splitting between these low-energy states is between 1.8–2 eV in the orbital

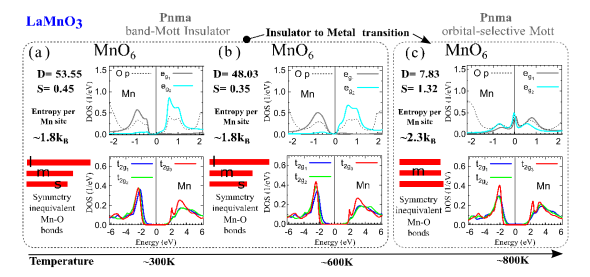


FIG. 2. Temperature-dependent electronic states in LaMnO₃. Panels (a) to (c) show for each inequivalent Mn octahedra the following properties: (1) values of the distortion indices, **D** and **S**; (2) electronic entropy per Mn site; (3) schematic view of the symmetry-inequivalent Mn-O bonds represented by red horizontal bars [the horizontal size of the bar is proportional to the bond length, short (s), medium (m), long (l); in panel (c) the three bonds are slightly distorted around the average bond length]; (4) e_g and t_{2g} projected density of states per formula unit versus energy in eV, for each electronic phase at a given temperature. On top of each panel we give the crystallographic symmetry at a given temperature and a name characterizing the electronic properties of the predicted electronic phases. The panels are connected by arrows suggesting the order parameter of the phase transitions.

ordered state, and the gap collapses around 800 K in the high-temperature orthorhombic structure, in which the three Mn-O bonds become very similar, although not strictly equal. Consequently, the two e_g orbitals are close to degenerate, but still different [see Fig. 2(c)]. The phase transition around 800 K is thus a first-order metal-insulator transition, strongly coupled with the structural evolution in which the distortion of the octahedra is strongly reduced, but the symmetry of the crystal structure does not change. This calculation shows not only that the details of the crystal structure and the lattice distortions have large impact on the electronic state, but also that the feedback effect of the electronic correlations on the crystal structure is crucial to properly describe the temperature evolution of the system. To understand whether structural distortions or electronic correlations drive the transition, we also calculate the electronic entropy [63] per Mn site at three temperatures as shown in Fig. 2. While normally the metal-lic phase has small entropy, and the Mott insulating state carries large entropy, here the situation is very different. In the insulating state we find the entropy to be around $1.8k_B$ per Mn site, while in the bad-metallic state it is increased to $2.3k_B$, which corresponds to 35 meV entropy gain across the transition, which is a decisive contribution to the free energy for the metal-insulator transition.

The important consequence of the Mott insulating t_{2g} states coexisting with the metallic e_g states at high temperature is the existence of the so-called *orbital-selective Mott state* (OSM), in which e_g electrons have enormous scattering rate (of the order of eV), and consequently small mobility so that there are no real bands present in the photoemission spectra, but rather just a very incoherent spectral weight (see Fig. 4). Photoemission on LaMnO₃ revealed that the system is of charge transfer type insulator, with strong admixture of oxygen p and Mn- e_g weight around 1 eV below the Fermi level

[51,64]. In Ref. [65] a strong peak at approximately -3 eV and a shoulder between -1.5 eV and the Fermi level was identified, which we now interpret as the t_{2g} and e_g peak in the density of states, respectively. Even more revealing are optical experiments, which show a strongly temperature-dependent peak around 2 eV present only in the orbitally ordered state [66], and replaced by a broad distribution of optical weight above 800 K. By analyzing low-temperature data in Ref. [67] it was found that the 2 eV peak in optics comes from e_g - e_g transitions, and that t_{2g} - e_g transitions appear at 3.8 eV. This is all consistent with our spectra in Fig. 2(a), as the e_g - e_g distance is close to 2 eV and the t_{2g} - e_g distance is close to 4 eV at room temperature, while in disordered state above 800 K the gap disappears and the scattering rate broadens low-energy excitations. We note that in earlier DMFT works, the e_g - e_g splitting in the ordered state was somewhat too large (≈3.5 eV [49] and ≈ 4 eV [50]), pointing again to the shortcomings of the low-energy Hubbard-type modeling. Finally we note that theoretical DOSs in Fig. 2 are plotted using experimental crystal structures, but the results do not change appreciably when the theoretical crystal structures are used.

While a large amount of experimental data is available for LaMnO₃, very little is known about BiMnO₃. Our calculations (Fig. 3) show a strong similarity with LaMnO₃, but also some important differences.

In particular, there is stronger anisotropy within the DOS of the t_{2g} and e_g orbitals for the Mn_2 ion compared to the Mn_1 ion [see Figs. 3(a) and 3(b)]. For example, there is a very sharp peak in the DOS of two t_{2g} orbitals at -2.0 eV and the e_g orbital at -0.6 eV, corresponding to the Mn_2 ion [Fig. 3(b)]. This is a consequence of the anisotropic hybridization between the lone-pair O-p and $Mn_2 - d$ orbitals. We recall that the lone pair of the Bi-6s state points toward the face of the distorted Mn_2O_6 octahedra as shown in

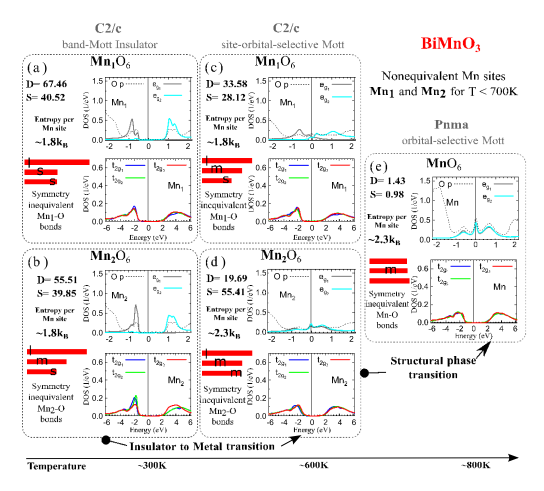


FIG. 3. Temperature-dependent electronic states in BiMnO₃. Panels (a) to (e) show for each inequivalent Mn octahedra the following properties: (1) values of the distortion indices, **D** and **S**; (2) electronic entropy per Mn site; (3) schematic view of the symmetry-inequivalent Mn-O bonds represented by red horizontal bars [the horizontal size of the bar is proportional to the bond length, short (s), medium (m), long (l); in panel (e) the three bonds are slightly distorted around the average bond length]; (4) e_g and t_{2g} projected density of states per formula unit versus energy in eV, for each electronic phase at a given temperature. On top of each panel we give the crystallographic symmetry at a given temperature and a name characterizing the electronic properties of the predicted electronic phases. The panels are connected by arrows suggesting the order parameter of the phase transitions.

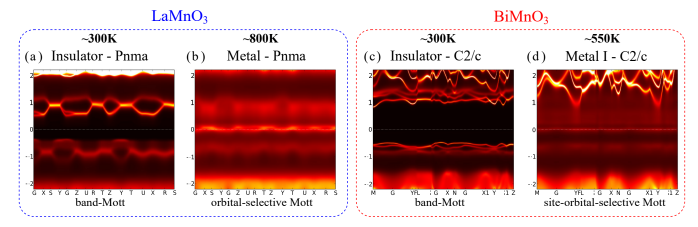


FIG. 4. Spectral functions for various novel phases in (Bi,La)MnO₃. For both compounds we give the spectral functions showing the metal-to-insulator phase transitions. In panels (a) and (b) we show the spectral functions for the band-Mott and orbital-selective Mott states for LaMnO₃; in panels (c) and (d) we show the spectral functions of the band-Mott and site-orbital-selective Mott states for BiMnO₃.

Fig. 1(d); thus this anisotropic hybridization is present in the system (see Ref. [68]). The anisotropic hybridization of the t_{2g} orbitals with the lone pair in the case of Mn_2 atoms is also confirmed by the fact that the Mn_1 atoms have similar distortions, but the corresponding DOS of the t_{2g} orbitals is isotropic [Fig. 3(a)]. The highly distorted monoclinic crystal structure of the BiMnO₃ compound is a consequence of the competition between the Jahn-Teller octahedra distortions that are squeezing the lone pair of the Bi-6s state. This is different from the case of the LaMnO₃ compound, where the lone pair is absent; thus the Jahn-Teller octahedra distortions appear in the orthorhombic crystal structure.

At room temperature, the local Jahn-Teller distortions for both inequivalent Mn ions are large enough to decrease the e_{g1} hybridization with the O-p states allowing localization of e_g electrons, even though the e_g orbitals also hybridize with the lone pair. These local distortions are depicted by red segments and given in terms of distortion indices in Figs. 3(a) and 3(b).

Note that larger values of the bond distortion index D in octahedra corresponds to more anisotropic hybridization between the two e_g orbitals and O-p states, causing orbital polarization and consequently localization of electrons. With increasing temperature our theoretical method reveals that the local Jahn-Teller distortions decrease (meaning smaller **D** values); thus the e_{g1} hybridization with the O-p states is increasing while the e_{g2} hybridization is decreasing (see Ref. [69]). At the same time the lone pair is exerting pres-sure on the neighboring ions to gain more space; thus it is contributing further to decreasing local distortions. Since the lone pair points toward the face of the Mn₂O₆ distorted octahedra, the local distortions of the Mn₂ site get reduced faster with increasing temperature than for the Mn₁ site. These local distortions are depicted by red segments and given in terms of distortion indices in Figs. 3(c) and 3(d). Consequently, when the local distortions are small enough, e_{g1} and e_{g2} hybridizations with the O-p states become similar; thus the e_g gap on Mn₂ ion collapses, and the system becomes bad metal. While decreasing the local distortions, the anisotropic hybridization of the t_{2g} with the lone-pair and O-p orbitals disappears; i.e., the resonance in the t_{2g} DOS disappears going from Figs. 3(b) to 3(d). However, the Mn_1 ion remains close to its low-temperature state, for one thing, its orbital occupancy in Table I remains unchanged, and the e_g states show DOS with a pseudogap [Fig. 3(c)]. Due to the qualitative difference in physical state on the two Mn ions, we call this state the siteorbital-selective Mott state, and we note that the orbital order remains strong only on the Mn₁ ion. Such partial orbital order on half of Mn ions is consistent with RXS experiments [43], in which the room temperature signal was roughly a factor of two stronger than in the intermediate state. Surprisingly, the entropy on Mn atoms is very weakly temperature depen-dent between the room temperature and the first transition (around 600 K), at which the entropy on Mn₂ increases for $0.5k_B$, while the entropy on Mn₁ remains almost unchanged. Only at the second transition around 800 K the entropy of Mn₁ increases to its saturated value within the bad metallic

We also note that the electronic spectral weight of the t_{2g} orbitals pointing toward the lone pair is strongly temper-ature dependent; therefore treating t_{2g} electrons as a fixed

core spin would not result in correct prediction of the crystal structures.

Figure 4 displays the single-particle spectral functions $A(\mathbf{K}, \omega)$ in the insulating room temperature state, and in the first metallic state. The room temperature direct gaps in LaMnO₃ and BiMnO₃ are ≈0.8 eV and 1.2 eV, which are in decent agreement with optical experiments (0.9 eV in LaMnO₃ [66] and 1.2 eV in BiMnO₃ [70,71]). We note that the prediction of the gap size in BiMnO₃ seems to be very challenging for other ab initio methods, as the gap size is either too large (hybrid functional 1.7 eV [28]) or too small (modified Becke-Johnson exchange functional 0.75 eV [72]; 0.5 eV or less by various LDA + U calculations [73–76]). Finally, the orbital-selective Mott phase, in which the orbital order is melted away, is displayed in Fig. 4(b). The spectral function has no sharp bands, only the incoherent spectral weight; hence the charge mobility is very low. This state is also realized in the ferromagnetic Kondo model of Ref. [62]; therefore it might show giant magnetoresistance. Figure 4(d) shows the new phase site-orbital-selective Mott state, realized in the intermediate temperature range where Mn₁ shows orbital ordering and Mn2 does not. This state is a superposition of a very incoherent orbital-selective Mott phase and the insulating phase, which shows somewhat more sharp bands above 1 eV.

Figure 5 shows the comparison between the theoretical and experimental temperature dependence of the distortion index **D**. In LaMnO₃ it changes only for 20% between the room temperature and 550 K, while it collapses abruptly around 800 K. We note an excellent agreement between the theoretical and experimental distortion index. In BiMnO₃ the two Mn sites show very different behavior: The changes on Mn₁ are comparable to LaMnO₃ (20%), while on Mn₂ ion the distortion is almost complete at 550 K (80%). In the hightemperature phase the distortion is small but finite. The reason for such markedly different behavior of the two sites is that the Mn₂ ion hybridizes with the Bi-6s lone pair, while Mn₁ does not (see Ref. [77]). Finally, we note that the measured and theoretical distortion indexes have similar trend, except that the disproportionation between Mn₁ and Mn₂ seems a bit weaker in experiment than in theory [78].

IV. CONCLUSIONS

In conclusion, using eDMFT methods we found novel electronic states in the Mn perovskite system, such as the site-orbital-selective Mott state, which hosts Mott insulating t_{2g} states, and a combination of semimetallic and metallic e_g states on two inequivalent Mn atoms. We showed that the isostructural transition in BiMnO₃ at 474 K should be characterized as the insulator-metal transition at which half of Mn atoms give up the orbital order, while the fully disordered state is reached only at the structural transition of 768 K. We showed that the electronic entropy plays the decisive role in the metal-insulator transition in both compounds. By selfconsistent determination of both the correlated electronic state and the corresponding relaxed crystal structure we showed that the temperature evolution of phases in LaMnO₃ and BiMnO₃ can be properly explained from first principles, thus demonstrating the predictive power of the eDMFT method for

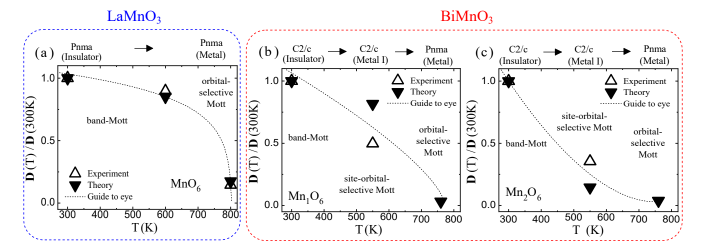


FIG. 5. Structural relaxations within LDA + eDMFT for all the temperature-dependent novel electronic phases in (Bi,La)MnO₃. In each panel we plot the bond distortion (**D**) index versus temperature, normalized to its values at room temperature. In panel (a) we plot **D** for the Mn site in LaMnO₃. In panels (b) and (c) we plot **D** for the two inequivalent Mn sites in BiMnO₃. Above each panel we give the temperature-dependent crystal symmetry and the properties of that electronic state (metal or insulator). The arrows point to the type of phase transition (structural or metal-insulator phase transition).

structural and electronic properties. In addition, we predict the temperature dependence of the spectral functions for the novel phases found in Mn perovskite systems, predictions that can be tested experimentally by ARPES.

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G.L.P. carried out the calculations. K.H. developed the code. G.L.P. and K.H. analyzed the results and wrote the paper.

The authors declare no competing interests.

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