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OPEN Trends in $(LaMnO_3)_n/(SrTiO_3)_m$ superlattices with varying layer thicknesses

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We investigate the thickness dependence of the structural, electronic, and magnetic properties of $(LaMnO_3)_n/(SrTiO_3)_m$ (n, m = 2, 4, 6, 8) superlattices using density functional theory. The electronic structure turns out to be highly sensitive to the onsite Coulomb interaction. In contrast to bulk SrTiO₃, strongly distorted O octahedra are observed in the SrTiO₃ layers with a systematic off centering of the Ti atoms. The systems favour ferromagnetic spin ordering rather than the antiferromagnetic spin ordering of bulk LaMnO3 and all show half-metallicity, while a systematic reduction of the minority spin band gaps as a function of the LaMnO, and SrTiO, layer thicknesses originates from modifications of the Ti d_{xy} states.

A zoo of extraordinary phenomena, such as two dimensional quantum gases¹, superconductivity², magnetism³, magnetoresistance⁴, exchange bias^{5,6}, half-metallicity⁷, and ferroelectricity⁸, is hosted by perovskite oxide heterostructures. The lattice, charge, orbital, and spin degrees of freedom and their competition give rise to these properties⁹. In general, electrostatic and strain effects drive structural and electronic reconstructions at interfaces¹⁰, which thus often behave distinctly different from their component materials. Nowadays the thickness of superlattices can be determined and controlled down to the atomic scale, where interface effects become more and more important. Thus, the thickness is a key factor for tailoring the electronic and transport properties. For example, Nanda and coworkers¹¹ have studied the magnetism in LaMnO₃/SrMnO₃ superlattices with varying layer thicknesses, concluding that the charge reconstruction is confined to two unit cells around the interfaces. Volume changes have been observed in NaNbO₃/SrTiO₃ superlattices¹², for example, and it has been demonstrated that the appearance of metallicity correlates with the layer thicknesses¹³.

Manganites and titanates are popular perovskite oxides, displaying colossal magnetoresistance¹⁴ and superconductivity¹⁵. Colossal magnetoresistance was first observed in 1994 in $La_{1-x}Ca_xMnO_3^{16}$ and gained considerable attention thereafter. The parent compound, LaMnO₃, is an antiferromagnetic insulator, which becomes ferromagnetic under hole doping with Ca¹⁷, Sr¹⁸, and Ba¹⁹. Therefore, superlattices with SrTiO₃ can be expected to show an unusual magnetic behavior and accordingly are attracting interest both for fundamental and technological reasons. The authors of Ref. 20, for example, have reported significant doping effects in LaMnO₃/SrTiO₃ superlattices, which can extend over few unit cells²¹, and have studied their origin. In LaSrMnO₃/SrTiO₃ superlattices with SrTiO₃ layers thinner than 1 nm an interfacial charge transfer leads to finite Ti magnetic moments²². In general, the thicknesses of the component layers in a LaMnO₃/SrTiO₃ superlattice seem to strongly influence the magnetic and transport properties²³. This would be interesting, because the behavior of the system could be controlled by means of the layer thickness. However, it requires detailed understanding of the strong lattice distortions and their implications for the physics of this superlattice, which is the topic of the present study.

Results

 $(LaMnO_3)_n/(SrTiO_3)_m$ superlattices are modeled by a $\sqrt{2} \times \sqrt{2}$ in-plane supercell of the cubic perovskite structure for taking into account possible effects of octahedral rotations. We use the Vienna Ab-initio

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Figure 1. Total densities of states of bulk SrTiO₃ and strained bulk LaMnO₃.





Simulation Package²⁴ with projector augmented wave pseudopotentials and employ the generalized gradient approximation in the parametrization of Perdew, Burke, and Ernzerhof (including spin polarization). The density of states is calculated by the tetrahedron method (smearing 0.01 eV) with Blöchl corrections²⁵. Because of the correlated nature of the transition metal *d* orbitals, we add an onsite Coulomb interaction using the Lichtenstein scheme²⁶. The values for the *U* and *J* parameters are taken from Refs 27–29 as 4 eV and 1 eV for the Mn *d* states, 5 eV and 0.5 eV for the Ti *d* states, and 9 eV and 1 eV for the La *f* states. We have tested for the bulk compounds that these values are transferable to the parametrization of Perdew, Burke, and Ernzerhof by comparing the densities of states, which show no qualitative difference. The optimized lattice constant of SrTiO₃ is 3.97 Å, with a band gap of 2.6 eV, and the optimized in-plane and out-of-plane lattice constants of LaMnO₃ (half-metallic) are shown in Fig. 1.

		n-IF	p-IF		n-IF	<i>p</i> -IF
Mn-O	2:2	2.12	2.13	4.4	2.17	2.00
Ti-O		2.04	1.91	4.4	2.01	1.90
Mn-O	2:4	2.17	2.03	4.2	2.14	2.07
Ti-O		2.00	1.93	4:2	2.04	1.88
Mn-O	2:6	2.12	2.01	6.2	2.14	2.07
Ti-O		1.95	1.96	0:2	2.04	1.91
Mn-O	2:8	2.12	1.99	0.7	2.13	2.13
Ti-O		1.95	1.97	0:2	2.08	1.87

Table 1. Bond distances (in Å) at the n-IF and p-IF for different n:m.



Figure 3. Total densities of states for thicknesses n:m = 2:2, 2:8, and 8:2.

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Structural optimization of the superlattices with and without onsite interaction is found to yield substantial differences in the electronic structure, especially near the Fermi level. More specifically, the minority channel shows a metallic character without onsite interaction for n:m = 2:2 (2 unit cells of LaMnO₃ alternate with 2 unit cells of SrTiO₃), whereas with onsite interaction we obtain a half-metallic character. The supercells have C_{2l_1} point group symmetry (tetragonal perovskite structure). An optimized in-plane lattice constant of 5.56 Å is obtained for the 2:2 system and is used for the larger superlattices. The out-of-plane lattice constant is optimized individually in each case. Starting from the 2:2 system, we fix the LaMnO₃ thickness and increase the SrTiO₃ thickness (2:4, 2:6, 2:8) or fix the SrTiO₃ thickness and increase the LaMnO₃ thickness (4:2, 6:2, 8:2). The 4:4 system is also considered for comparison. Figure 2(left) illustrates the 2:4 system as an example. There are two types of interfaces, the *n*-type LaO/ TiO₂ interface and the *p*-type SrO/MnO² interface, denoted in the following as n-IF and p-IF, respectively. The nomenclature refers to the compensating charges formed at the $(La^{3+}O^{2-})^+/(Ti^{4+}O_2^{2-})^0$ and $(Sr^{2+}O^{2-})^0/(Mn^{3+}O_2^{2-})^-$ contacts, respectively.

The Ti-O-Mn bond angle at the n-IF (where the MnO_6 and TiO_6 octahedra are connected) is found to be 159.4°, whereas at the p-IF it has a larger value of 164.8°, which corresponds to a reduced tilting. While substantial modifications of the structure are expected in the vicinity of an interface, strong effects are also observed for the O octahedra further apart. For instance, the Mn-O-Mn bond angles increase by up to 8.3° with respect to the bulk value (154.6°), which supports double exchange. On the other hand, the Ti-O-Ti bond angles strongly decrease throughout the SrTiO₃ layer, reflecting huge distortions as compared to the bulk cubic structure. In general, the Ti-O-Ti (Mn-O-Mn) bond angles near the p-IF (n-IF) are less modified, because the distance to the interface is larger. The tiltings of the O octahedra are found to vary distinctly for different thicknesses *n:m*, and so do the electronic properties.



Figure 4. Total densities of states near the Fermi energy for thicknesses n:m = 2:2, 2:4, 2:6, 2:8, 4:2, 6:2, and 8:2.

For increasing *m* the Mn-O (Ti-O) bond length along the *z* axis at the p-IF decreases (increases) from 2.13 Å to 1.99 Å (1.91 Å to 1.97 Å), while at the n-IF the Mn-O bond length stays close to 2.13 Å and the Ti-O bond length decreases from 2.04 Å to 1.95 Å, see Table 1. A schematic view of the Mn-O and Ti-O bonds is given in Fig. 2(right). For increasing *n* the Mn-O bond length at the p-IF maintains a value of 2.10 Å, while the Ti-O bond length decreases (bulk value 1.95 Å) from 1.91 Å in the 2:2 system to 1.87 Å in the 8:2 system. At the n-IF the Mn-O bond length shows no clear trend, while the Ti-O bond length grows significantly from 2.04 Å in the 2:2 system to 2.08 Å in the 8:2 system. It is found that all Ti atoms shift systematically off the center of their O octahedron towards the p-IF by up to 0.1 Å (in all cases more at the p-IF than at the n-IF), implying that a permanent dipole is created, while bulk SrTiO₃ is not ferroelectric. The off-centering is generally enhanced at the interfaces when *n* increases and reduced when *m* increases. It is known that the properties of SrTiO₃ are very sensitive to dopants and external perturbations³⁰ so that it cannot surprise that the superlattices react similarly³¹. We also note that we obtain for bulk SrTiO₃ a ferroelectric distortion of 0.03 Å, which is a known artefact of the employed methodology³². However, this effect is significantly smaller than the off-centerings described above and therefore does not affect our conclusions.

Bulk LaMnO₃ is an A-type antiferromagnetic (AFM) Mott insulator, due to a combination of superexchange and Zener double exchange³³, whereas SrTiO₃ is a non-magnetic insulator (d^0 configuration of Ti). Figure 3 shows the total densities of states obtained for the 2:2, 2:8, and 8:2 systems. The general



Figure 5. Projected densities of states of the Ti atom next to the n-IF for thicknesses n:m = 2:2, 2:4, 2:6, 2:8, 4:2, 6:2, and 8:2.

shape of the curves is similar, showing half-metallicity in each case. From -7 to -2 eV we find mainly Mn *d*, Ti *d*, and O *p* states, from -2 to 2 eV Mn *d* and O *p* states, and above 2 eV Ti *d*, La *f*, Sr *d*, and O *p* states. Figure 4 demonstrates systematic changes for varying *n* and *m*. The density of states at the Fermi energy remains similar for growing *m* but increases significantly for growing *n* due to the fact that these states mainly belong to the LaMnO₃ layer. Importantly, for increasing *n* as well as *m*, the minority spin band gap is reduced substantially. Indeed, the band onsets in the 2:8 and 8:2 systems are close to the Fermi energy so that the half-metallicity will most likely vanish for *n*, *m* > 8.

The reduction of the minority spin band gap is explained by the projected densities of states of the Ti atoms at the n-IF in Fig. 5. The Ti 3*d* orbitals, being split into d_{xy} , d_{yz} , d_{zz} (t_{2g}) and $d_{x^2-y^2}$, $d_{3z^2-r^2}$ (e_g) states due to the octahedral crystal field, systematically shift to lower energy for increasing *n* and *m*, which is consistent with previous experimental observations for LaAlO₃/SrTiO₃ interfaces³⁴. The energetic lowering is strongest for the d_{xy} states, which thus govern the reduction of the minority spin band gap. The orbital ordering seen in Fig. 5 decreases for increasing *m*, while it remains similar for increasing *n*. In addition, it is always more pronounced at the n-IF than at the p-IF, and almost lost at the p-IF and in the bulk-like regions of the 2:8 system. Figure 6 gives projected densities of states of Ti atoms in layers with increasing distance from the 2:8 n-IF. As to be expected, away from the n-IF the states shift to higher energy. Projected densities of states of the Mn atoms at the p-IF are shown in Fig. 7. Note that the Mn $d_{x^2-y^2}$ and $d_{3z^2-r^2}$ orbitals contribute at the Fermi energy in the majority spin channel, though differently for the different systems. Since bulk LaMnO₃ forced into C_{2h} symmetry becomes metallic, the conducting



Figure 6. Projected densities of states of the Ti atoms in layers with increasing distance from the 2:8 n-IF.

Mn states are a result of the in-plane strain. The metallicity at the p-IF is substantially reduced for increasing *n*, especially in the 8:2 system, as the LaMnO₃ layer becomes more bulk-like. Figure 8 gives projected densities of states of the Mn atoms in layers with increasing distance from the 8:2 p-IF. The distinct difference between the $d_{x^2-y^2}$ and $d_{3z^2-r^2}$ states is reduced when the distance increases.

In order to assess the magnetic ground state, we study the total energies for FM and A-type AFM spin ordering for strained bulk LaMnO₃ and the superlattices. Other spin orderings (such as C-type and G-type AFM) have much higher energies, as we have checked for the 2:2 system. FM spin ordering is found to be favorable, in agreement with previous experiments^{20,23,34} and calculations³⁵ as well as reports on related perovskite oxide interfaces³⁶⁻³⁸. The energy difference (per formula unit) between FM and A-type AFM spin ordering is 0.75 eV for strained bulk LaMnO₃, whereas in the case of the superlattices it slightly decreases from 0.27 eV to 0.20 eV for increasing *m* and remarkably increases from 0.27 eV to 0.85 eV for increasing *n*, which shows that it can be attributed to the LaMnO³ layer (the Ti magnetic moments are very small). Because of this, we normalize our results in the following with respect to the number of Mn atoms. The energy difference steadily decreases for increasing *n* as well as *m* though the FM ordering, which is due to Zener double exchange between partially filled Mn e_g states³⁹, remains favorable. For x = 0.3 FM spin ordering is also established in La_{1-x}Ca_xMnO₃, see the neutron diffraction experiments in Ref. 40. Moreover, X-ray magnetic circular dichroism and absorption spectroscopy on



Figure 7. Projected densities of states of the Mn atom next to the p-IF for thicknesses n:m = 2:2, 2:8, and 8:2.

LaAlO₃/SrTiO₃ have given direct evidence of a FM interface state, originating from the Ti d_{xy} orbitals³⁴. Interestingly, induced Ti magnetic moments of 0.02 to 0.04 μ_B are observed at both interfaces in our systems, see Table 2, whereas far from the interfaces they are negligible (<0.01 μ_B). For increasing *m* they increase (decrease) at the n-IF (p-IF), while for increasing *n* we obtain higher magnetic moments at the p-IF (~0.04 μ_B) than at the n-IF (~0.02 μ_B). We note that even small Ti magnetic moments at perovskite oxide interfaces can have important consequences for the magnetism^{34,41}.

We find Mn magnetic moments of 3.72 to $4.03 \mu_B$, with larger values at the n-IF than at the p-IF in all cases. When *m* increases they increase (decrease) at the n-IF (p-IF), while the changes are not systematic under variation of *n*. The highest value appears in the 4:4 system at the n-IF. Concerning the charges in the atomic orbitals, we find small but finite differences from the corresponding bulk values at both interfaces, see Table 2. A detailed analysis shows that for increasing *n* (*m*) the Mn atoms lose more (less) charge at the p-IF. Moreover, the Ti (Mn) atoms gain (lose) charge at the n-IF (p-IF) in all cases, while there is no clear trend at the p-IF (n-IF) because of the larger distances of the atoms from to the interface plane. The observed small charge transfers give rise to the mentioned Ti magnetic moments but cannot explain the ferroelectric distortions in the SrTiO₃ layer, which thus are likely direct consequences of the interface interaction.

Discussion

The structural, electronic, and magnetic properties of $(LaMnO_3)_n/(SrTiO_3)_m$ superlattices have been investigated and compared for various thicknesses *n*:*m*. Both *n* and *m* are found to strongly affect the material properties. In general, the SrTiO₃ layers show heavy distortions of the O octahedra, which are not present in bulk SrTiO₃, consistent with recent experimental findings⁴². Since LaMnO₃ is polar along the [001] direction (with alternating (LaO)⁺ and (MnO₂)⁻ layers) and SrTiO₃ is non-polar, a compensation mechanism is required. As the strained LaMnO₃ is metallic, we could expect accumulation of charge carriers at the interfaces. However, this effect is much too small according to the charge deviations reported in Table 2 so that a second mechanism must play a role. The projected densities of states of the Ti atoms in layers of increasing distance from the 2:8 n-IF in Fig. 6 show an almost continuous shift of all curves to higher energy. This reflects the built up of an electric field, which in turn induces ferroelectric distortions in the SrTiO₃ region, as discussed before. The ferroelectric distortions constitute the main compensation mechanism in the present superlattice. Similar distortions of the TiO₆ octahedra have been reported in Ref. 28.

All studied systems exhibit half-metallicity, though our results indicate that this property will vanish for n, m > 8. FM spin ordering is always energetically favorable over AFM spin ordering but the preference becomes smaller for increasing n, since the A-type AFM ordering of bulk LaMnO₃ starts to dominate. The Ti magnetic moments observed at the interfaces are small. The authors of Ref. 43 have found ferromagnetic and antiferromagnetic alignment of the Mn and Ti spins for n:m ratios of 3:2 and 17:2, respectively, and have argued that Mn-O-Ti superexchange therefore must play a role. In general, defects and cation intermixing are important for the stabilization of perovskite superlattices. For instance,





	magnetic moment				charge deviation				
n:m	Mn <i>n</i> -IF	Mn <i>p</i> -IF	Ti n-IF	Ti p-IF	Mn <i>n</i> -IF	Mn <i>p</i> -IF	Ti n-IF	Ti p-IF	
2:2	3.87	3.86	0.01	0.03	+0.01	-0.05	+0.02	+0.01	
2:4	3.95	3.76	0.02	0.02	+0.03	-0.02	+0.02	+0.01	
2:6	3.96	3.74	0.02	0.02	+0.03	-0.02	+0.02	+0.01	
2:8	3.97	3.72	0.03	0.02	+0.04	-0.01	+0.02	-0.01	
4:4	4.03	3.78	0.02	0.03	+0.03	-0.04	+0.02	0.00	
4:2	3.99	3.85	0.02	0.03	0.00	-0.04	+0.01	+0.01	
6:2	4.01	3.88	0.02	0.03	-0.01	-0.07	+0.06	-0.03	
8:2	3.97	3.89	0.02	0.04	+0.01	-0.07	+0.04	+0.03	

Table 2. Magnetic moments (in μ_B) and charge deviations with respect to the bulk (in electrons) of the interfacial Mn and Ti atoms for different *n:m*. Positive (negative) values represent gain (loss) of charge.

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the high electrical conductivity of the LaAlO₃/SrTiO₃ interface is related to the formation of O vacancies during the deposition process due to the extra charges introduced into the system⁴⁴. In addition, Sr/La intermixing leads to a metallic interface between the insulators LaTiO₃ and SrTiO₃, because mixed valent Ti states are created⁴⁵. We expect that these factors are less important for the LaMnO₃/SrTiO₃ interface, as it was found to be rather sharp⁴². We find a systematic reduction of the minority spin band gaps with increasing *n* and *m*, which originates mainly from an energetic downshift of the Ti d_{xy} states.

Methods

The atomic sphere radii are chosen such that overlap is avoided during the ionic relaxation, for which we set the energy tolerance to 10^{-3} eV, employ a $4 \times 4 \times 1$ k-mesh, a Gaussian smearing of 0.05 eV, and an energy cutoff of 500 eV. To obtain accurate electronic states, we set the energy tolerance to 10^{-5} eV and adopt dense k-meshes of $8 \times 8 \times 3$ for n + m = 4, $8 \times 8 \times 2$ for n + m = 6, and $8 \times 8 \times 1$ for n + m = 8 and 10.

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Author Contributions

F.C. and U.S. designed the study. J.J. performed the calculations. All authors contributed to the interpretation of the results and the writing of the manuscript.

Additional Information

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