Magnetism of (LaCoO$_3$)$_n$+(LaTiO$_3$)$_n$ superlattices (n=1,2)
Alex Taekyung Lee and Sohrab Ismail-Beigi
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Magnetism of \((\text{LaCoO}_3)_n+(\text{LaTiO}_3)_n\) superlattices with \(n = 1, 2\)

Alex Taekyung Lee and Sohrab Ismail-Beigi

Department of Applied Physics, Yale University

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LaCoO

is a rare earth and where \(A\) is a rare earth and \(B\) is a transition metal) display fascinating physical properties, such as high-temperature superconductivity in the cuprates or colossal magnetoresistance in the manganites [1, 2]. The electronic and structural properties of perovskites and intertwined lead to complex behaviors due to the interplay of charge, magnetic, structural, and orbital degrees of freedom [3]. Creating heterostructures of transition metal perovskites enlarges the playground for manipulation of the local structure, charge state and electronic properties of these materials. In this work, we focus on the electronic states of cobalt-containing perovskites.

Bulk LaCoO

has attracted much attention because of the complex electronic and magnetic structure of the Co

ions it contains. LaCoO

is a nonmagnetic insulator at the lowest temperatures [4, 5] when the Co

adopt a low-spin (LS) state \(t^6_{2g}, S = 0\). It becomes a paramagnetic insulator for temperatures \(100 \, K < t < 500 \, K\) and then becomes metallic above \(500 \, K\). However, the spin state of Co in the thermally excited material is still in debate. The magnetic transition near \(100 \, K\) is considered as a transition from LS to high-spin (HS) \(t^5_{2g}e^2_{g}, S = 2\) [6, 7] or to intermediate spin (IS) \(t^5_{2g}e^3_{g}, S = 1\) states [5, 8–11], while IS and HS are mixed as temperature increases [10, 11].

First principles density functional theory (DFT) calculations based on DFT+\(U\) theory [12] for LaCoO

confirm that the LS state is the ground state [13–15], while the IS state is more stable than HS [13]. On the other hand, several theoretical studies suggested that the excited state at \(100 \, K < t < 500 \, K\) can be a mixture of HS and LS, which we denote as HS/LS below. Using the experimental crystal structures as a function of temperature, GGA+\(U\) calculations suggested that HS/LS is stabilized for the structures above \(200 \, K\) [15]. Unrestricted Hartree-Fock calculations also proposed that the non-magnetic to paramagnetic transition is due to the LS state to LS/HS state [16], and effective Hamiltonian calculations suggested a similar scenario [17, 18]. Recent DFT+dynamical mean-field theory calculations showed that LaCoO

has large charge fluctuations: these calculations show that it is not possible to explain the spin state with a single multiplet at any temperature, although they find that at the onset of the spin-spin transition HS multiplets are excited with the IS multiplets being excited later around the onset of the metal-insulator transition [19].

On the other hand, the electronic and magnetic structures of Co

in bulk CoO does not show such complexity or ambiguity. At low temperatures, CoO is an insulator with energy gap \(E_g = 2.4 \, eV\) [20]. The Co

cations have a HS state \(t^5_{2g}e^2_{g}\) with \(S = 3/2\), and antiferromagnetically order at \(T_N = 289 \, K\) with a magnetic ordering wave vector of \(q_{cc} = (\frac{1}{2}, \frac{1}{2}, \frac{1}{2})\) [21, 22]. Unlike LaCoO

, Co

in CoO does not show LS or IS states nor a spin-state transition.

An intriguing set of questions arise as to what happens if the Co

ions are realized in a perovskite heterostructure containing LaCoO

. Is the difference between the electronic behavior of LaCoO

and CoO only due to charge state of the Co cations? To what extent can the Co electronic properties be controlled or manipulated by changing the local lattice structure surrounding the Co cation? And how can one make strong modifications to the local structure? One way to control the charge state of transition metal cations and the local lattice structure is via heterostructuring. For example, charge transfer can be triggered from Ti to Ni in LaNiO

+LaTiO

superlattices [23, 24] which results in a Ni

charge state with large \(c_{g}\) orbital polarization, which is in strong contrast to the orbitally unpolarized Ni

in bulk LaNiO

. Recent work on LaCoO

+LaTiO

superlattices also shows charge transfer and robust orbital polarization [25].

In this work, we study the electronic and magnetic properties of Co in \((\text{LaTiO}_3)_1+(\text{LaCoO}_3)_{1}\) and \((\text{LaTiO}_3)_2+(\text{LaCoO}_3)_{2}\) superlattices using DFT+\(U\) theory. We explore carefully the effect of the Hubbard \(U\) parameter, account for the effects of different exchange-correlation approximations, examine the effect of strain.
from the substrate, as well as the effects of doping. We explain the origin of the magnetic stabilities for different spin states and magnetic orderings by using a simple band interaction picture. Interestingly, from the analysis of the magnetic stabilities, we are able to show that ferromagnetic spin order on for the Co\(^{2+}\) cations can be stabilized in the (LaTiO\(_3\))\(_1\)+(LaCoO\(_3\))\(_1\) superlattice by hole doping which is an unexpected and unique finding for Co\(^{2+}\) as detailed below (Summary and outlook section). We note that recent experiments [25] have only studied the (LaTiO\(_3\))\(_2\)+(LaCoO\(_3\))\(_2\) superlattice so our results for the (LaTiO\(_3\))\(_1\)+(LaCoO\(_3\))\(_1\) superlattice present exciting predictions for experiments.

This paper is organized as follows. Sec. II describes our computational approach. Sec. III describes the key predicted superlattice properties: structure, electron transfer, Co spin states, and magnetic properties as a function of strain and doping. Sec. IV A is devoted to analyzing and understanding computed properties: energetics of HS and LS states as a function of \(U_{\text{Co}}\), the origin of the \(U_{\text{Co}}\) dependences, stabilities of various magnetic orderings as a function of \(U_{\text{Co}}\), a simple band interaction model for understanding the results, further strain effects, the effects of different exchange-correlation approximations, the cause of the stabilization of ferromagnetism in hole-doped (LaCoO\(_3\))\(_1\)+(LaTiO\(_3\))\(_1\) superlattices. In Sec. IV B, we apply the above analysis of electronic and magnetic properties to the (LaCoO\(_3\))\(_2\)+(LaTiO\(_3\))\(_2\) superlattice and describe the importance of out-of-plane Co–Co magnetic interaction. We then summarize the work and provide an outlook.

II. Computational Details

We use density functional theory (DFT) with the projector augmented wave (PAW) method [26] and the revised version of the generalized gradient approximation (GGA) proposed by Perdew et al. (PBEsol) [27] as implemented in the VASP software [28]. We also have done selective tests using the local density approximation (LDA) [29, 30]. In all cases, the spin-dependent version of the exchange correlation functional is employed. A plane wave basis with a kinetic energy cutoff of 500 eV is used. We use \(\Gamma\)-centered \(k\)-point meshes of size \(9\times9\times7\) and \(9\times9\times5\) for (LaCoO\(_3\))\(_1\)+(LaTiO\(_3\))\(_1\) and (LaCoO\(_3\))\(_2\)+(LaTiO\(_3\))\(_2\), respectively. The sizes of the supercells for (LaCoO\(_3\))\(_1\)+(LaTiO\(_3\))\(_1\) and (LaCoO\(_3\))\(_2\)+(LaTiO\(_3\))\(_2\) are 20 and 40 atoms, respectively, corresponding to \(\sqrt{2} \times \sqrt{2} \times \sqrt{2} \) (i.e., \(c(2 \times 2)\)) in-plane unit cells. These unit cells have two Co atoms in each CoO\(_2\) plane, so that the (LaCoO\(_3\))\(_1\)+(LaTiO\(_3\))\(_1\) and (LaCoO\(_3\))\(_2\)+(LaTiO\(_3\))\(_2\) each have 2 and 4 Co atoms in their unit cells, respectively. Atomic positions within the unit cells were relaxed until the residual forces were less than 0.01 eV/Å in all axial directions, and the stress is less than 0.5kB along the \(z\) axis. The GGA+\(U\) scheme within the rotationally invariant formalism together with the fully localized limit double-counting formula [12] is used to study the effect of electron interactions within the 3\(d\) orbital manifolds. The electronic and structural properties critically depend on \(U_{\text{Co}}\), and therefore we carefully explore a range of values. We also explore how the results depend on \(U_{\text{Tl}}\), which plays a secondary role in the physics of these materials. We do not employ an on-site exchange interaction \(J\) for any species, as the exchange interaction is already accounted for within the spin-dependent DFT exchange-correlation potential [31, 32]. We used Maximally Localized Wannier Functions (MLWF) as implemented by the Wannier90 software [33] to obtain a localized tight-binding representation describing Co 3\(d\)-derived bands. Finally, we point the reader to Appendix A for our results on the \(U\) dependence of the properties of bulk LaCoO\(_3\), CoO, and LaTiO\(_3\) which help us find proper ranges and reasonable values for the \(U\) parameters.

III. Key results

A. (LaCoO\(_3\))\(_1\)+(LaTiO\(_3\))\(_1\) superlattice

In this subsection, we discuss our key findings for the atomic, electronic, and magnetic properties of (LaCoO\(_3\))\(_1\)+(LaTiO\(_3\))\(_1\) superlattices.
We find that \( R \) and choose the most stable one. Since bulk LCO has the not known experimentally (due to thinness of the samples) and is resulting in Jahn-Teller-like distortions, tilting of oxygen octahedra is also important in determining the electronic structure of transition metal oxides. Unlike bulk oxides, the tilt pattern of the superlattice is harder to observe \( a \) priori, so we consider several tilt patterns and choose the most stable one. Since bulk LCO has the \( R3c \) crystal structure with \( a^{-}a^{-}a^{-} \) octahedral tilts and bulk LaTiO\(_3\) has the \( Pbnm \) structure with \( a^{-}a^{-}b^{+} \) tilts, we examine three cases: \( a^{0}a^{0}a^{0} \), \( a^{-}a^{-}a^{-} \) and \( a^{-}a^{-}b^{+} \). We find that \( a^{-}a^{-}b^{+} \) is always the most stable, so we only examine this tilt pattern below unless otherwise specified.

The structures are relaxed along the \( c \) axis, which is normal to the interface, while in-plane lattice parameters \( a \) and \( b \) are fixed and set equal \( a = b \) to simulate epitaxial strain from a substrate. Three in-plane lattice parameter values are used below: 3.663 Å, 3.784 Å, and 3.905 Å. The two values 3.784 Å and 3.905 Å are the experimental lattice parameters of LaAlO\(_3\) (LAO) and SrTiO\(_3\) (STO) substrates, respectively. To study the effect of the compressive strain on LCO, we use 3.663 Å which is smaller than LCO lattice. We note that the optimized lattice parameter of bulk LCO within our GGA+\( U \) calculations (with \( U_{Co} = 3 \) eV) is 3.805 Å, which is close to the LAO lattice constant.

2. Electron transfer from Ti to Co

Next, we discuss the charge states of transition metals and the electron transfer. Given the difference in electronegativity between Ti and Co [34, 35], electron transfer is expected from Ti \( d \) to Co \( d \) across the interface. Projected densities of states (see Fig. 2) within GGA+\( U \) show that the Ti \( d \) bands are unoccupied, which corresponds to Ti\(^{4+}\), and describe the conduction band minimum. On the other hand, Co has 2+ charge state with \( d^{7} \) electrons, indicating full charge transfer (formal charge).

In order to further examine the nature of the charge transfer, we computed the mean energy of the 3d orbitals of Ti and Co (by using the projected density of states as weight) to learn about the electronic potential landscape across the interface [36]. For (LCO\(_3\))\(_1\)+(LTO)\(_1\) and (LCO\(_3\))\(_2\)+(LTO)\(_2\), the Ti 3d atomic orbitals are higher than the Co 3d by 5.2 and 1.9 eV, respectively.

Due to the charge transfer, we have Ti\(^{4+}\) and Co\(^{2+}\), and we expect local electric fields pointing from the TiO\(_2\) layers to the CoO\(_2\) layers. This can be confirmed by looking at the positions of the positively charged La\(^{3+}\) cations and negatively charged apical oxygen anions (O\(^{2-}\)): the displacements are visible to the eye in Fig. 1. For example, for (LCO\(_2\))\(_2\)+(LTO)\(_2\) on an STO substrate, the out of plane Co–O bond length between the CoO\(_2\) planes (denoted as a bulk-like Co–O bond in Fig. 1) is 2.02 Å, while the bond length between the CoO\(_2\) and TiO\(_2\) planes (denoted as an interfacial Co–O bond in Fig. 1) is 2.28 Å, indicating that O\(^{2-}\) is shifted to the opposite direction of the local electric field.

In (LTO)\(_1\)+(LCO)\(_1\), the CoO\(_6\) octahedron faces interfaces on both sides. A simpler theoretical view of the structure, unencumbered by octahedral tilting, is af-
forced by considering \((\text{LCO})_1+\text{(LTO)}_1\) with \(d^0 a^6 d^0\) tilt within GGA+U whose structure is shown in Fig. 3. Here, the distance between the \(\text{La}^{3+}\) plane and the \(\text{CoO}_2\) plane is 1.75 Å, while the distance between \(\text{La}^{3+}\) and \(\text{TiO}_2\) plane is 2.10 Å.

3. Spin states of \(\text{Co}^{2+}\)

The HS state \((t^2_g e^2_g)\) with \(S = 3/2\) is depicted in Figs. 4(a) and (b): the spin-up Co 3d bands are fully occupied, while spin-down bands have two electrons in the \(t_{2g}\) channel. Due to the non-zero \(U_{\text{Co}}\), the spin-down \(t_{2g}\) splits into two degenerate bands \((d_{xz}, d_{yz})\) under the Fermi level and one empty single band \((d_{xy})\) above the Fermi level, resulting in a Mott-like insulator. We find that \(U_{\text{Co}} > 0\) is necessary for opening an energy gap for the HS state. In addition, the HS state is not even metastable at \(U_{\text{Co}} = 0\): \(U_{\text{Co}} > 2\) eV is needed (with \(U_{\text{Ti}} = 3\) eV) to obtain a gapped and stable HS state. Regarding orbital polarization, the degeneracy of the \(t_{2g}\) manifold is already removed at \(U_{\text{Co}} = 0\) because of the broken structural symmetry due to interface. This splitting between \(d_{xz}/d_{yz}\) and \(d_{xy}\) is further enhanced by imposed epitaxial strain and also enlarged by non-zero \(U_{\text{Co}}\).

The LS state \((t^6_e g^2)\) has \(S = 1/2\) and is presented in Figs. 4(c) and (d): the \(t_{2g}\) bands are fully occupied, while the one remaining electron is in the \(e_g\) channel. The degeneracy of the \(e_g\) manifold is already broken at \(U_{\text{Co}} = 0\) due to the interface formation and epitaxial strain. Due to the large degeneracy splitting of the \(e_g\) bands, only the spin-up \(d_{x^2-y^2}\) and spin-down \(e_g\) bands are empty. As a result, the LS state is already insulating at \(U_{\text{Co}} = 0\) and has strong \(e_g\) orbital polarization; \(U_{\text{Co}} > 0\) enhances the orbital polarization and the energy gap.

4. Magnetic stability and effect of strain

There are two distinct Co atoms in our \((\text{LTO})_1+\text{(LCO)}_1\) unit cell. Each Co can be HS or LS as well as spin up or spin down polarized. Therefore, there are 6 different spin configurations: ferromagnetic with two high-spin Co (HS FM), antiferromagnetic with two high-spin Co (HS AFM), ferromagnetic with one high-spin and one low-spin Co (HS/LS FM), ferromagnetic with one high-spin and one low-spin Co (HS/LS FIM), ferromagnetic with two low-spin Co (LS FM), and antiferromagnetic with two low-spin Co (LS AFM).

As summarized by Fig. 5, HS AFM is most stable over a wide range of \(U_{\text{Co}}:\ 3.5 < U \leq 5\) eV for \(a = 3.784\) Å, and \(2.5 < U < 5\) eV for \(a = 3.784\) and \(a = 3.905\) Å. For a fixed Co spin state, HS AFM is more stable than HS FM, HS/LS FM is more stable than HS/LS FIM, and LS FM is more stable than LS AFM.

Interestingly, as shown in Fig. 5, the LS phase is stabilized by compressive strain whereas the HS phase is stabilized by tensile strain. In addition, both the HS FM and HS AFM phases are stabilized as \(U_{\text{Co}}\) increases. Certain combinations of strain and \(U_{\text{Co}}\) can drive magnetic transitions for the \(\text{Co}^{2+}\): LS FM can be stabilized over HS AFM for compressive strain and low \(U_{\text{Co}}\) values, and an intermediate spin (HS/LS combination) state can become the ground state (e.g., \(U_{\text{Co}} = 2.5\) eV and \(a = 3.663\) Å). However, over broad ranges of parameters, it is the HS AFM that is the ground state. To the best of our knowledge, HS to LS or FM to AFM transitions have not been suggested or observed for \(\text{Co}^{2+}\) in bulk CoO, and this makes the LCO+LTO superlattice a potential test bed for modifying the spin state and magnetic interactions of \(\text{Co}^{2+}\) cations. We analyze the origin of the magnetic stability (Sec. IV A 3) and effect of strain (Sec. IV A 4) below.

5. Stabilization of ferromagnetism with hole doping

Electron or hole doping is a general and powerful way to change the magnetic stability of transition metal oxides such as cuprates [37, 38] and manganites [39, 40]. For Co, the situation is more complex as a variety of spin states are possible, requiring a detailed analysis.

By manually changing total number of electrons in the supercell, both the spin state and the magnetic ordering can be changed via electron or hole doping in \((\text{LTO})_1+\text{(LCO)}_1\) superlattices. In Fig. 6, we present the energies of six different phases as a function of electron/hole doping. The spin of each Co cation is in fact different since the added electron or hole resides on one
FIG. 5: $U_{Co}$ dependence of the energies of different magnetic states of (LTO)$_1$+(LCO)$_1$ within GGA+$U$, for different in-plane lattice parameters (a) 3.663 Å, (b) 3.784 Å, and (c) 3.905 Å. Energies of LS FM phase is set to zero.

of the Co ions. Thus, what is referred to as an AFM alignment of Co spins is in fact a ferrimagnetic ordering, but for simplicity we refer to such phases by their parent undoped phase in Fig. 6.

The figure shows that the relative energetic stabilities are essentially unchanged by electron doping: the HS/LS FM phase is most stable at $a = 3.663$ Å (Fig. 6(a)), and the HS AFM phase is most stable at $a = 3.784$ Å and $a = 3.905$ Å (Figs. 6(b,c)) for all considered electron doping values. On the other hand, the relative magnetic stabilities are changed dramatically by hole doping: the energies of the hole-doped HS phases show large changes (red and blue curves in Fig. 6). The ferromagnetic HS FM phase becomes the ground state if the hole doping per Co is larger than ≈0.2, ≈0.13, and ≈0.15 for $a = 3.663, 3.784,$ and $3.905$ Å, respectively. This pre-
diction is particularly interesting, because hole doping via Sr substitution on La sites should be much easier to achieve in LTO-based systems than electron doping. We analyze the doping effect in detail below (Sec. IV A 5) and show that the FM stabilization is driven by the double exchange mechanism. We note that the presence of an AFM to FM transition with hoping is not obvious a priori e.g., Fig. 6(a) shows that the LS state lacks any such transition while the HS state has a transition as a function of hole doping. To the best of our knowledge, magnetic systems comprised of Co²⁺ cations do not show ferromagnetism nor are they expected to display double exchange physics. We look forward to experimental verification of this prediction.

B. Key results for (LaCoO₃)₂+(LaTiO₃)₂ superlattice

Our computational supercells now contain 4 Co and 4 Ti. Since there is a great deal of shared physics between the 1+1 and 2+2 superlattices, we will briefly describe the main physical properties of the 2+2 superlattices while highlighting key differences: our overall approach and methods of analysis are identical for the two superlattices. The Co²⁺ cations in the (LCO)₂+(LTO)₂ superlattice can be HS or LS, similar to (LCO)₁+(LTO)₁, so that there are 5 possible configurations HS/LS ordering in the superlattice as presented in top row of Fig. 7: all HS order (shortened to HS), A-type HS/LS order with ordering wave vector \( \mathbf{q}_{sc} = (0, 0, 1) \) (A-HS/LS), C-type HS/LS order with \( \mathbf{q}_{sc} = (1, 1, 0) \) (C-HS/LS), G-type HS/LS order with \( \mathbf{q}_{sc} = (1, 1, 1) \) (G-HS/LS), and all LS order (LS). Separately, there are 4 possible orderings patterns for the Co magnetic moments: FM, A-type AFM, C-type AFM, and G-type AFM (see second row of Fig. 7). In total, we investigate all \( 5 \times 4 = 20 \) distinct configurations.

We summarize the \( U_{Co} \)-dependent properties of the most relevant low-energy configurations in Fig. 8. For most choices of parameters, our full set data show that the HS G-type AFM phase is the ground state, except for the case of high compressive strain and small \( U_{Co} (a=3.663 \text{ Å and } U_{Co}=2.5 \text{ eV}) \) where we find that G-HS/LS A-AFM phase is more stable. To avoid excessive clutter, Fig. 8(a) only shows data for G-type AFM magnetic ordering. In general, the energies of the A-HS/LS, C-HS/LS, G-HS/LS phases lie between those of the LS and HS phases. For a fixed magnetic pattern (Fig. 8(a,c,e)), the energy difference \( \Delta E \) between LS and HS phases is enhanced as \( U_{Co} \) increases or as the in-plane lattice parameter \( a \) increases, both of which mirror the behavior of \( (LTO)₁+(LCO)₁ \).

In Figs. 8(b,d,f), we compare the energies of different magnetic orderings as a function of \( U_{Co} \) when all then Co are in the HS state. The energies in this case obey the relations

\[ E_{FM}[HS] > E_{A-AFM}[HS] > E_{C-AFM}[HS] > E_{G-AFM}[HS], \]

and, interestingly, this order holds for all \( U_{Co} \) and strain ranges considered. The origin of the magnetic stabilities are described in detail below (Sec. IV B)

IV. Analysis of spin states and magnetic stability

In this section, we provide a detailed analysis of the electronic and magnetic properties of the \( (LaCoO₃)₁+(LaTiO₃)₁ \) and \( (LaCoO₃)₂+(LaTiO₃)₂ \) superlattices. Each subsection deals with a particular physical effect or provides a microscopic model explaining the computed behaviors.

A. Analysis of \( (LaCoO₃)₁+(LaTiO₃)₁ \)

As discussed in Sec. III A 4, there are two distinct Co atoms in our supercell of \( (LTO)₁+(LCO)₁ \). Since two Co atoms can be HS or LS separately, and also can be spin-up or spin-down separately, there are 6 different spin configurations as mentioned: ferromagnetic with two high-spin Co (HS FM), antiferromagnetic with two high-spin Co (HS AFM), ferromagnetic with one high-spin and one low-spin Co (HS/LS FM), ferrimagnetic with one high-spin and one low-spin Co (HS/LS FIM), ferromagnetic with two low-spin Co (LS FM), and antiferromagnetic with two low-spin Co (LS AFM). Among them, HS AFM is the most stable over a wide range of parameters, as shown in Fig. 9(a). For fixed spin states, HS AFM is more stable than HS FM (Fig. 10(a)), HS/LS FM is more stable than HS/LS FIM (Fig. 10(c)), and LS FM is more stable than LS AFM (Fig. 10(e)).

1. HS vs LS with fixed magnetic order: \( U_{Co} \) dependence

The above energy differences strongly depend on \( U_{Co} \). Specifically, when the magnetic ordering is fixed to AFM, the energy difference between different spin states of Co increases monotonically as a function of \( U_{Co} \) as shown in Fig. 9(a). To explain this strong dependence, we analyze the difference \( E[HS \text{ AFM}]-E[LS \text{ AFM}] \) by rewriting the DFT+\( U \) total energy as a spectral decomposition \[ E = E_{DFT+U} = E_{DFT} + E_{fill} + E_{ord} \]

where

\[ E_{fill} = U(2l + 1)\mu(1 - \mu), \quad E_{ord} = -U(2l + 1)\sigma^2. \]

Here \( l = 2 \) is the angular momentum of the atomic-like states for the DFT+\( U \) treatment, and \( \mu \) and \( \sigma \) are the mean value and standard deviation of the eigenvalues
FIG. 7: Five possible HS/LS orderings and four possible magnetic orderings of the \((\text{LaCoO}_3)_{2}+(\text{LaTiO}_3)_{2}\) superlattice.

FIG. 8: \(U\) dependence of energies for different spin configurations of \((\text{LTO})_2+(\text{LCO})_2\) with different in-plane lattice parameters. Energy of LS FM is set to be zero.

of the single particle density matrix of the manifold of atomic-like orbitals (here Co 3d states). Within this spectral decomposition, the combined interaction and double counting energy simply depends on the two numbers \(\mu\) and \(\sigma\) extracted from the local single-particle density ma-

FIG. 9: \(U_{\text{Co}}\) dependence of the total energies for AFM states of \((\text{LTO})_{1}+(\text{LCO})_{1}\) superlattices. (a) Total energies per Co of different magnetic configurations (the total energy of the LS state is chosen as the zero of energy). (b) Spectral decomposition of the \(+U\) energy contribution (see text).

We define \(\Delta E_{\text{fill}}^{\text{AFM}}\) and \(\Delta E_{\text{ord}}^{\text{AFM}}\) via

\[
\Delta E_{\text{fill}}^{\text{AFM}} = E_{\text{fill}}^{\text{AFM}}[\text{HS}] - E_{\text{fill}}^{\text{AFM}}[\text{LS}]
\]

\[
\Delta E_{\text{ord}}^{\text{AFM}} = E_{\text{ord}}^{\text{AFM}}[\text{HS}] - E_{\text{ord}}^{\text{AFM}}[\text{LS}]
\]

and plot them versus \(U_{\text{Co}}\) in Fig. 9(b). While the change of \(\Delta E_{\text{fill}}^{\text{AFM}}\) as a function of \(U_{\text{Co}}\) is relatively weak, \(\Delta E_{\text{ord}}^{\text{AFM}}\) is the dominating contribution, similar to prior studies [41, 42]. Therefore, the \(U_{\text{Co}}\) behavior is dictated largely by the difference in \(\sigma^2\) of the two respective spin configurations. In other words, the orbital occupancy differentiation (i.e., orbital ordering) is more enhanced in the HS state than in the LS state as \(U_{\text{Co}}\) increases, and this enlarges the energy difference between them.

2. FM vs AFM for fixed Co spin state: \(U_{\text{Co}}\) dependence

Now we consider the energy difference between two magnetic configurations for fixed Co spin state.
Figs. 10(c,d), the situation is very similar to that in the way as above: e.g., the difference between FM and AFM decreases as a function of \(U_{Co}\), and relatively small in size while \(\Delta E_{\text{ord}}[\text{HS}]\) dominates the behavior in that its increase with increasing \(U_{Co}\) explains the positive slope of \(\Delta E[\text{HS}]\) or \(\Delta E[\text{LS}]\).

We also considered the intermediate spin HS/LS FM and AFM (the latter is in fact ferrimagnetic as the moments are unequal) configurations where the unit cell contains one LS and one HS Co cation. As presented in Fig. 10(e), \(\Delta E[\text{HS/LS}]\) varies by about 10 meV as \(U_{Co}\) changes from 2.5 to 5 eV. \(\Delta (E_{\text{fill}} + E_{\text{ord}})[\text{HS/LS}]\) for the HS Co and the LS Co (see Fig. 10(f)) show opposing behaviors versus \(U_{Co}\), so their sum is relatively constant. As a result, the averaged value of \(\Delta (E_{\text{fill}} + E_{\text{ord}})\) at \(U = 2.5\) eV and \(U = 5\) eV are only differ by 12 meV.

3. Origin of the magnetic stabilities

The above numerical observations of the relative magnetic stabilities require explanations. Here, we will explain the origin of the magnetic energetics of \((\text{LTO})_1 + (\text{LCO})_1\) using an energy band picture (i.e., molecular orbital theory) for three different spin configurations: HS, HS/LS, and LS.

Fig. 11 shows the schematic energy diagram for two in-plane HS Co neighbors with FM and AFM relative spin alignment. The interaction lines in this diagram do not indicate direct Co 3d-Co 3d hybridization but instead the interaction mediated through the intervening O 2p states. The energetic position of the atomic-like energy levels are drawn based on analysis of projected densities of states diagrams such as Fig. 4.

The spin-up majority bands of HS FM are fully occupied, thus the Co-Co interactions do not lead to energetic stabilization as per Fig. 11(a) (i.e., both bonding and anti-bonding states are equally occupied). We note that our Co cations have strong orbital degeneracy breaking whereby the \(t_{2g}\) \(d_{yz}/d_{zx}\) bands are lower in energy than the \(d_{xy}\) band is empty: in the minority-spin channel, the \(d_{yz}/d_{zx}\) bands are filled while the \(d_{xy}\) band is empty. As discussed in Sec. IV A, this splitting results a Mott-like insulating state. In the HS FM case, the minority-spin bands also do not lead to energetic stabilization: the \(d_{yz}(d_{zx})\) spin-down on one Co only interact with the \(d_{yz}(d_{zx})\) spin-down states on the neighboring Co, and as both are filled, there is no net lowering of energy as the bonding and anti-bonding states are again fully occupied (Fig. 11(b)). Thus the energy lowering due to Co-Co interactions for HS FM is zero, \(\Delta E^{FM}[\text{HS}] = 0\).

On the other hand, HS AFM Co-Co superexchange interactions lead to energetic stabilization (Figs. 11(c,d)). While one Co has 5 spin-up and 2 spin-down electrons, the other Co has 2 spin-up and 5 spin-down electrons: \(d_{z^2}\), \(d_{yz}\), and \(d_{zx}\) are fully occupied for both Co ions leading to no energy lowering; but the electrons in the majority-spin \(d_{x^2-y^2}\) and \(d_{xy}\) states interact with empty counterparts on the other Co which leads to energy lowering (only bonding states become filled) as shown in Figs. 11(c,d) and (e), (f).

\[
\Delta E_{\text{fill}}[\text{HS}] = E_{\text{fill}}^{FM}[\text{HS}] - E_{\text{fill}}^{FM}[\text{HS}], \quad \Delta E_{\text{ord}}[\text{HS}] = E_{\text{ord}}^{FM}[\text{HS}] - E_{\text{ord}}^{FM}[\text{HS}].
\]
FIG. 11: Schematic molecular orbital view of Co 3d energy level diagrams and in-plane magnetic interactions between two neighboring HS Co$^{2+}$ cations. (a) and (b) show two neighboring Co with FM alignment, while (c) and (d) show two neighboring Co with AFM alignment. Solid horizontal lines indicate energy levels; dashed lines indicate the effect of interactions; arrows indicate electron filling colored by orbital type. $\Delta_{11}$ and $\Delta_{12}$ are exchange splitting for FM and AFM configurations, respectively. $\Delta E$ is the energy lowering due to the magnetic interactions.

Figs. 11(c) and (d). If we define the strength of the Co-Co interaction for one 3d orbital on one Co with the same orbital on the neighbor for the FM case as $-\Delta_{11}$ (see Fig. 11(b)) and for the AFM case as $-\Delta_{12}$ (see Fig. 11(c)), respectively, then the energy lowering of the spin-up channel in the AFM case is

$$\Delta E_{1}^{AFM}[HS] = -\Delta_{12} d_{x^2-y^2} - \Delta_{12} d_{xy},$$

The spin-down channel has same amount of the energy lowering. Therefore, within this model, the energy difference between HS FM and HS AFM can be written as

$$\Delta E[HS] = \Delta E^{AFM}[HS] - \Delta E^{FM}[HS]$$

$$= -2 \left( \Delta_{12} d_{x^2-y^2} + \Delta_{12} d_{xy} + \Delta_{12} d_{z^2} \right),$$

which is negative as the $\Delta_{ij}$ are positive by our convention. Hence, this analysis provides a simple explanation of the stability of HS AFM compared to HS FM.

FIG. 12: Schematic diagram of the in-plane magnetic interaction between HS Co and LS Co.
lowering for FM ordering of HS and LS neighboring Co is
\[ \Delta E_{\text{FM}}^{\text{HS/LS}} = -\Delta_{d_{z^2}}^{11} - \Delta_{d_{xy}}^{12}. \] (7)

The case of ferrimagnetic (FIM) ordering between a HS Co and LS Co is shown in Figs. 12(c,d). For the spin-up channel, the HS \( d_{x^2-y^2} \) and \( d_{z^2} \) are occupied but empty for the LS Co, leading to an energy lowering of \( -\Delta_{d_{x^2-y^2}}^{12} - \Delta_{d_{z^2}}^{12} \); a similar lowering happens for the low-spin channel due to \( d_{xy} \) and \( d_{z^2} \). The total energetic stabilization for this FIM HS–LS case is
\[ \Delta E_{\text{FIM}}^{\text{HS/LS}} = -2\Delta_{d_{z^2}}^{12} - \Delta_{d_{x^2-y^2}}^{12} - \Delta_{d_{xy}}^{12}. \] (8)

Hence, the energy difference between HS/LS FM and HS/LS FIM is thus
\[ \Delta E^{\text{HS/LS}} = \Delta E_{\text{FIM}}^{\text{HS/LS}} - \Delta E_{\text{FIM}}^{\text{HS/LS}} = - \left[ \Delta_{d_{x^2-y^2}}^{11} - \Delta_{d_{xy}}^{12} \right] - \left[ \Delta_{d_{x^2}}^{11} - \Delta_{d_{xy}}^{12} \right] + 2\Delta_{d_{z^2}}^{12}. \] (9)

For the \( d_{x^2-y^2} \) and \( d_{xy} \) contributions, there is some partial cancellation between the \( \Delta_{d_{x^2-y^2}}^{11} \) and \( \Delta_{d_{xy}}^{12} \) terms although these contributions should be net negative since FM interactions \( \Delta^{11} \) take place between degenerate orbitals while the AFM \( \Delta^{12} \) are between strongly exchange-split 3d orbitals leading to weaker interactions (i.e., homopolar versus heteropolar bonding in the language of semiconductors). However, with the addition of the positive contribution \( 2\Delta_{d_{z^2}}^{12} \), the total is expected to be relatively small in magnitude, in agreement with the small energy differences shown in Fig. 10(e). Comparison to data in the figure shows that \( \Delta_{d_{x^2-y^2}}^{11} > \Delta_{d_{xy}}^{12} \) must be true so that FIM is higher in energy than FM.

Understanding the stability of the FM state for two LS Co neighbors, however, is more complicated. The simple band interaction picture cannot explain the data in Fig. 10(e) which show that LS FM is more stable than LS AFM: as per Fig. 13, the energy lowering of the FM configuration is zero while the AFM ordering should be lower in energy by \( -2\Delta_{d_{z^2}}^{12} \). Furthermore, standard double exchange for FM configurations is unlikely to be a viable explanation since the system is not metallic (it has an energy gap) due to the large splitting between \( t_{2g} \) and \( e_g \) bands. Nor can Goodenough-Kanamori (GK) type FM superexchange be the origin of the stability, because GK superexchange is maximum if \( \angle \text{TM–O–TM}=90^\circ \) and cannot be applied for \( \angle \text{TM–O–TM}=180^\circ \) [43–45]. We find that FM is also stable for the insulating undistorted \( a^0b^0d^0 \) phase, where Co–Co angle is 180\(^\circ\).

In the above analysis, we only considered the interaction between same \( d \) orbitals on the neighboring Co, e.g., \( d_{x^2-d_{z^2}} \) or \( d_{xy-d_{xy}} \), since they are usually the dominant ones. The model predicted that the LS AFM state is stabilized by \( -2\Delta_{d_{z^2}}^{12} \); but will be a very small quantity in this case since the exchange splitting of the \( d_{z^2} \) band is
larger than 4 eV (see Fig. 4(b)) and we expect the hopping between the two neighboring \( d_z \) orbitals to be small since they are strongly directional out-of-plane. Hence, it is likely that interactions between different orbitals on the neighboring Co can play a role in this case.

To capture the essence of the problem, we consider the ideal \( a^0 \alpha a^0 \) structure without octahedral tilts so the Co–O–Co bonding is along a straight line. In this case, the only symmetry allowed interactions that can lower the energy for the FM order are \( d_{z^2} - d_{x^2-y^2} \) between neighboring atoms for the majority spin-up channel (Figs. 14(a,b)). For AFM order, the relevant hoppings are \( d_{z^2} - d_z \) (small and already discussed above) and \( d_{z^2} - d_{x^2-y^2} \) in both spin channels as per Figs. 14(c,d).

We now create a more quantitative model by extracting on-site energies and hopping parameters for the LS FM and AFM cases by generating Maximally Localize Wannier Functions (MWLF) [33] that span the energy bands \( \{ \delta E \} \) as obtained from a Maximally Localize Wannier Function analysis.

### TABLE I: Hopping parameters \((t)\) and energy splitting of \( d_{z^2}\) and \( d_{z^2} \) bands \((\delta E)\) as obtained from a Maximally Localize Wannier Function analysis.

<table>
<thead>
<tr>
<th>Order Interaction</th>
<th>( \delta E ) (eV)</th>
<th>( t ) (eV)</th>
<th>( t^2/\delta E ) (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FM ( d_{z^2} - d_{x^2-y^2} )</td>
<td>2.04</td>
<td>-0.120</td>
<td>7.0 \times 10^{-3}</td>
</tr>
<tr>
<td>AFM ( d_{z^2} - d_{z^2} )</td>
<td>3.38</td>
<td>-0.010</td>
<td>2.9 \times 10^{-5}</td>
</tr>
<tr>
<td>AFM ( d_{z^2} - d_{x^2-y^2} )</td>
<td>2.98</td>
<td>-0.116</td>
<td>4.5 \times 10^{-3}</td>
</tr>
</tbody>
</table>

FIG. 15: \( U_{Co} \) dependence of the energies of different magnetic states of \((\text{LTO})_1 + (\text{LCO})_1\) for different in-plane lattice parameters (3.663 Å for (a,b), 3.784 Å for (c,d), and 3.905 Å for (e,f)) and two different exchange-correlation functionals \((a,c,e)\) for GGA+U and \((b,d,f)\) for LDA+U. Energies of LS FM phase is set to zero. The energy of the FM phase is set to zero. (a), (c), (e) are same as Fig. 5.

4. Effect of strain and exchange-correlation functional

Given that the spin, orbital, and lattice degrees of freedom interact strongly in a perovskite transition metal oxygen octahedron, the electronic and magnetic properties of Co in \((\text{LTO})_1 + (\text{LCO})_1\) depend on strain markedly. To study this, we consider the three different in-plane lattice parameters discussed above \((a = 3.663 \text{ Å}, 3.784 \text{ Å}, \) and \(3.905 \text{ Å})\). Within GGA+U for \( U_{Co} = 3 \text{ eV} \), these three lattice parameters corresponds to 3.7% compressive, 0.6% compressive, and 2.6% tensile strain.

In Fig. 15, we summarize the strain-dependent stabilities of different spin states and magnetic orderings of Co within both GGA+U and LDA+U. As discussed in Sec. III A 4, the HS phases are stabilized greatly as the lattice parameter increases (see Figs. 15(a,c,e)): the LS phase is stabilized by compressive strain, whereas the HS phase is stabilized by tensile strain. In addition, similar to the case of \( a = 3.784 \) Å discussed in detail above in Sec. IV A 1, HS FM and HS AFM phases are stabilized as \( U_{Co} \) increases (see Figs. 15(a,c,e)). Various combinations of strain and \( U_{Co} \) can drive magnetic transitions for the Co\(^{2+}\). LS FM can be stabilized over the HS AFM for compressive strain and low \( U_{Co} \) and an intermediate spin (HS/LS combination) state can become the ground state (e.g., \( U_{Co} = 2.5 \text{ eV} \) and \( a = 3.663 \) Å). However, over broad ranges of parameters, it is the HS AFM that is the ground state. To the best of our knowledge, HS to
LS or FM to AFM transitions have not been suggested or observed for Co\textsuperscript{2+} in bulk CoO, and this makes the LCO+LTO superlattice a potential test bed for modifying the spin state and magnetic interactions of Co\textsuperscript{2+} ions.

Analysis of the results of Fig. 15 show that the $U_{\text{Co}}$ dependence of the energy difference between HS and LS phases originates from the orbital occupancy differentiation energy $E_{\text{ord}}$ (Eq. 2) much like what was shown in Fig. 9. On the other hand, we do not find that the relative stabilization of the LS phase for compressive strain is connected to the atomic-like +$U$ contributions to the total energy $E_{\text{tot}}$ nor $E_{\text{hll}}$ but must instead originate from the underlying DFT energy $E_{\text{DFT}}$ that encodes the electronic dispersion and hopping contributions. The idea that the DFT part of the total energy controls the relative stability of the LS phase means that its stability should strongly depend on the choice of DFT functional. This is borne out by Fig. 15 which compares the GGA and LDA functionals. The trends versus $U_{\text{Co}}$ and strain are very similar between the two functionals, but the energy of the HS phases is shifted upwards compared to the LS phases as we switch from GGA to LDA.

\subsection*{5. Effect of doping}

Here we analyze the origin of the ferromagnetic stability with hole doping, which is discussed in Sec. III A 5. As described above, we find that both the spin state and the magnetic ordering can be changed via electron or hole doping in (LTO)$_1$+(LCO)$_1$ superlattices.

To understand these results, we begin with the fact that both the electron-doped and hole-doped HS phases become metallic in our calculations. Doped electrons partially occupy the minority spin-down $d_{xy}$ states, and doped holes resides on majority spin-up $d_{z^2}$ and $d_{x^2-y^2}$ states. Given the metallicity of the hole-doped state, the stability of the FM ordering turns out to be due to the double exchange mechanism.

We use the schematic band interaction diagrams of in Fig. 16 to explain this stabilization mechanism. As per Fig. 16(a), the energy lowering of FM HS ordering is nonzero because of the holes in majority spin-up $d_{z^2}$ and $d_{x^2-y^2}$ bands. Defining the number of holes as $n_{d_{z^2}}^h$ and $n_{d_{x^2-y^2}}^h$, the energy lowering of the FM ordering is

$$\Delta E_{\text{FM}} = -2n_{d_{x^2-y^2}}^h \Delta_{x^2-y^2}^{11}, \quad (10)$$

where the factor of two is due to the holes on both neighboring Co migrating to the shared anti-bonding $d_{x^2-y^2}$ state. The energy lowering of the AFM ordering (strictly speaking, it is ferrimagnetic as explained above) is

$$\Delta E_{\text{AFM}} = -(2 - 2n_{d_{x^2-y^2}}^h) \Delta_{x^2-y^2}^{12} - 2\Delta_{x^2-y^2}^{12} - 2\Delta_{x^2-y^2}^{12}. \quad (11)$$

The energy difference between the two phases is thus

$$\Delta E_{\text{FM}} - \Delta E_{\text{AFM}} = -2n_{d_{x^2-y^2}}^h \left(\Delta_{x^2-y^2}^{11} + 2\Delta_{x^2-y^2}^{12} \right) + 2 \left(\Delta_{x^2-y^2}^{12} + \Delta_{x^2-y^2}^{12} + \Delta_{x^2-y^2}^{12}\right). \quad (12)$$

This energy difference can change sign with increasing hole doping $n_{d_{x^2-y^2}}^h$. Furthermore, since $\Delta_{x^2-y^2}^{11}$ energies are much larger than $\Delta_{x^2-y^2}^{12}$ energies, the value of $n_{d_{x^2-y^2}}^h$ at which it changes sign should be small, in agreement with our numerical findings.
and interestingly this order holds for all $U_{\text{Co}}$ and strain ranges considered.

As described in Fig. 11 and its associated discussion, in-plane magnetic interactions stabilizes AFM spin alignment, thus $E_{\text{AFM}}^{\text{FM}}[\text{HS}]$ and $E_{\text{AFM}}^{\text{AFM}}[\text{HS}]$ are above $E_{\text{AFM}}^{\text{FM}}[\text{HS}]$ and $E_{\text{AFM}}^{\text{AFM}}[\text{HS}]$.

In order to explain why $E_{\text{AFM}}^{\text{FM}}[\text{HS}] > E_{\text{AFM}}^{\text{AFM}}[\text{HS}]$ and $E_{\text{AFM}}^{\text{AFM}}[\text{HS}] > E_{\text{AFM}}^{\text{AFM}}[\text{HS}]$, i.e., the stability AFM order along the out-of-plane direction, we examine the out-of-plane magnetic interaction for the HS phase in Fig. 18. In the out-of-plane direction, only $d_{x^2}$, $d_{z^2}$, and $d_{xy}$ on two neighboring Co will interact significantly. While the FM alignment has no net energy lowering due to out-of-plane interactions (see Figs. 18(a,b)), the interactions between the $d_{x^2}$ on the two Co lowers the energy by $-2\Delta d_{x^2}$ for AFM alignment (Figs. 18(c,d)). The energy difference between the FM and AFM configurations due to out-of-plane interactions is

$$\Delta E[\text{HS}] = \Delta E_{\text{AFM}}[\text{HS}] - \Delta E_{\text{FM}}[\text{HS}] = -2\Delta d_{x^2},$$

so that we expect $E_{\text{AFM}}^{\text{AFM}}[\text{HS}] < E_{\text{AFM}}^{\text{FM}}[\text{HS}]$ and $E_{\text{AFM}}^{\text{AFM}}[\text{HS}] < E_{\text{AFM}}^{\text{AFM}}[\text{HS}]$. 

B. Analysis of (LaTiO$_3$)$_2$+(LaCoO$_3$)$_2$

We now turn to analyze the electronic and magnetic properties of (LaCoO$_3$)$_2$+(LaTiO$_3$)$_2$ superlattices, which are briefly discussed in Sec. III B. As in Sec III B, there is a great deal of shared physics between the 1+1 and 2+2 superlattices, thus we will only describe the main physical properties of the 2+2 superlattices while highlighting key differences: our overall approach and methods of analysis are identical for the two superlattices.

There are 5 possible configurations HS/LS ordering in the (LaCoO$_3$)$_2$+(LaTiO$_3$)$_2$ supercell, as we already discussed in Sec. III B and top row of Fig. 7: all HS order (shortened to HS), A-type HS/LS order (A-HS/LS), C-type HS/LS order (C-HS/LS), G-type HS/LS order (G-HS/LS), and all LS order (LS). Separately, there are 4 possible orderings patterns for the Co magnetic moments: FM, A-type AFM, C-type AFM, and G-type AFM (see second row of Fig. 7). In total, we investigate all $5 \times 4 = 20$ configurations.

As discussed in Sec. III B, HS G-AFM is the ground state for most choices of parameters (see Figs. 8 or 17), and the energies of the A-HS/LS, C-HS/LS, G-HS/LS phases lie between those of the LS and HS phases. For a fixed magnetic pattern (Fig. 8(a,c,e)), the energy difference ($\Delta E$) between LS and HS phases is enhanced as $U_{\text{Co}}$ increases or in-plane lattice parameter $a$ increases which is similar to the behavior of (LTO)$_1$+(LCO)$_1$. Again, like the behavior of the 1+1 superlattices described in Sec. IV A 1, the $U_{\text{Co}}$-dependence of $\Delta E$ for 2+2 superlattices turns out to be explained by the behavior of the ordering energy $\Delta E_{\text{ord}}$ (Eqs. $\text{(1.2)}$) between LS and HS phases.

As mentioned in Sec. III B, the energies different magnetic orderings obey the relations

$$E_{\text{FM}}^{\text{FM}}[\text{HS}] > E_{\text{AFM}}^{\text{AFM}}[\text{HS}] > E_{\text{AFM}}^{\text{AFM}}[\text{HS}] > E_{\text{AFM}}^{\text{AFM}}[\text{HS}],$$

and interestingly this order holds for all $U_{\text{Co}}$ and strain ranges considered.

As described in Fig. 18 and its associated discussion, in-plane magnetic interactions stabilizes AFM spin
V. Summary and outlook

LCO+LTO superlattices show strong electron transfer, significant structural distortion, and robust orbital polarization. For a wide range of computational parameters, we predict high-spin Co and a checkerboard antiferromagnetic (AFM) order on the Co superlattice (2D checkerboard for the 1+1 superlattice and 3D checkerboard, i.e., G-type, for the 2+2 superlattice). A detailed analysis of the electronic structure reveals how superexchange interactions control the stability of the magnetic orders. Finally, hole doping of the superlattices can lead to stabilization of a ferromagnetic ground state of Co$^{2+}$. This is unexpected and interesting especially for future experiments. Generally, materials containing Co$^{2+}$ as the sole magnetic cation, such as CoO or Co$_2$O$_4$ (which also contains non-magnetic Co$^{3+}$ [46]), show antiferromagnetic ordering. A ferrimagnetic compound such as CoFe$_2$O$_4$ contains Fe$^{3+}$ at the A sites and Fe$^{2+}$/Co$^{2+}$ at B sites which are aligned antiferromagnetically [47]. Hence, the ability to create a ferromagnet using high-spin Co$^{2+}$ is a new and, in our mind, surprising prediction.

Experimentally, we hope to see verification of the checkerboard AFM order of these superlattices. Even more exciting will be the experimental realization of ferromagnetism via hole doping through replacement of La by Sr; by careful control of the doping, the FM and AFM order can be made nearly degenerate in energy, potentially leading to interesting magnetic response and new magnetic phases.

Theoretically, more advanced models such as DFT+dynamical mean field theory can be used to verify the predictions made here using GGA+U. Furthermore, understanding the origin of the orbital polarization in these charge-transfer systems is interesting in terms of basic science as well as electronic engineering. Finally, investigation of thicker LCO layers in the superlattice, i.e., (LTO)$_n$+(LCO)$_m$ for $n \geq 3$, should prove interesting as well, since the interaction between Co$^{2+}$ at the interface and Co$^{3+}$ at the bulk-like region may lead to other unexpected physical phenomena.

VI. Acknowledgments

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Here, we aim to obtain optimal $U_{Co}$ value for Co$_2^+$ in the LCO+LTO superlattices by studying bulk LaCoO$_3$ (Co$_3^+$) and CoO (Co$_2^+$). Below, we will show that the experimental nonmagnetic ground state for LaCoO$_3$ is stable with $U_{Co} \leq 2.5$ eV for LaCoO$_3$, while $U_{Co} \sim 4.5$ eV is needed to obtain the experimental band gap of bulk CoO (see Figure 19). We will conclude that $U_{Co} = 3$ eV is a reasonable value.

A. Appendix: $U$ dependence of LaCoO$_3$ & CoO
The total energies of the nonmagnetic insulating phase (NM) for 0 ≤ UCo < 2.5 eV is appropriate to reproduce the nonmagnetic low-temperature ground state in the experiment. Within LDA+U, the S = 0 NM state is the ground state for 0 ≤ UCo ≤ 4.5 eV.

In addition to the energetics of the various spin states, we also consider the band gap (E_g) of bulk LaCoO_3. As summarized in Fig. 19 (c), E_g within GGA+U with U = 2.5 eV and U = 3 eV are 0.51 and 0.67 eV, respectively, which are both close to the theoretical gap of ≈ 0.6 eV [48]. Within LDA+U, U = 3.5 eV gives E_g = 0.65 eV, as shown in Fig. 19 (d).

Considering both the nature of the magnetic ground state and a reasonable value of the energy gap, we conclude that U = 2.5 eV within GGA+U and U = 3.5 eV within LDA+U are reasonable values for studying bulk LaCoO_3. Our U values are comparable to those from previous DFT+U studies: U_{Co}−J_{Co}=2.7, 3.5, and 4.5 eV [15] or LDA+U values of U_{Co} = 7.8 eV and J_{Co} = 0.92 eV [13].

### 2. Bulk CoO

In the LaCoO_3+LaTiO_3 superlattice, the electron transfer from Ti to Co will lead to a 2+ charge state for Co (see the next section). Therefore, we also calculate the U_{Co} dependence of the band gap of bulk CoO which also contains Co^{2+}.

At low temperature, bulk CoO is antiferromagnetic with wave vector q_{205} = (\frac{1}{3}, \frac{2}{3}, \frac{1}{3}) [21, 22], and an insulator with energy gap E_g = 2.4 eV [20]. In Figs. 19 (c) and (d), we plot E_g of CoO as a function of U_{Co}. U_{Co} = 4 eV within GGA+U and U_{Co} ≈ 4.8 eV within LDA+U give the experimental gap. These values are similar to the previous GGA+U results: E_g = 2 eV with U_{Co}−J_{Co}=3.3 eV [49], E_g = 2.8 eV with U_{Co} = 7.1 eV and J_{Co} = 1 eV [49].
3. Choosing $U_{Co}$

Within GGA+$U$, we need $U_{Co} = 2.5$ eV for bulk LaCoO$_3$ to be described well, and $U_{Co} = 4$ eV for bulk CoO. Since we expect our superlattice to contain Co$^{2+}$ which is more similar to CoO, we choose $U_{Co} = 3$ eV unless otherwise specified. Below, we will explore $0 \leq U_{Co} \leq 5$ eV to study the $U$-dependence of the electronic and magnetic properties. We consider $U_{Ti} = 0$, 3, and 5 eV, and use $U_{Ti} = 3$ eV unless specified otherwise. This value of $U_{Ti}$ is obtained by fitting the optical gap of LaTiO$_3$ to be 0.2 eV based on our previous work [50].