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Magnetism, spin texture and in-gap states: Atomic specialization at the surface of oxygen-deficient SrTiO₃

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Motivated by recent spin- and angular-resolved photoemission measurements (SARPES) of the two-dimensional electronic states confined near the (001) surface of oxygen-deficient SrTiO₃, we explore their spin structure by means of *ab initio* density functional theory (DFT) calculations of slabs. Relativistic non-magnetic DFT calculations display Rashba-like spin winding with a splitting of a few meV and when surface magnetism on the Ti ions is included, bands become spin-split with an energy difference ~ 100 meV at the Γ point, consistent with SARPES findings. While magnetism tends to suppress the effects of the relativistic Rashba interaction, signatures of it are still clearly visible in terms of complex spin textures. Furthermore, we observe an *atomic specialization* phenomenon, namely two types of electronic contributions; one is from Ti atoms neighboring the oxygen vacancies that acquire rather large magnetic moments and mostly create in-gap states; another comes from the partly polarized t_{2g} itinerant electron system and are responsible for the Rashba spin winding and the spin splitting at the Fermi surface.

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Introduction. - Transition metal oxides constitute a major topic of interest in the scientific community as these materials are endowed with a broad range of significant functionalities, ranging from ferroelectricity to metalinsulator transitions as well as from magnetism to superconductivity. Many of these compounds exhibit structural instabilities, strong electronic correlations and complex phase diagrams with competing ground states. Artificial structures of transition metal oxides therefore seem ideal to explore interfacial effects that could possibly lead to new phases. In this respect, the observation a decade ago [1, 2] of a two-dimensional electronic system (2DES) at the interface between the wide band-gap insulators $LaAlO_3$ (LAO) and $SrTiO_3$ (STO) has attracted a considerable amount of attention. It was found that the 2DES hosts gate-tunable insulator to metal, insulator to superconductor transitions, magnetism [3] and a large interfacial spin-orbit effect [4]. The mechanisms responsible for these special properties are still under debate. Questions include the intrinsic (i.e. electronic reconstruction) versus extrinsic (e.g. oxygen vacancies) mechanism [5–7] responsible for the formation of the 2DES, and also the role and spatial distribution of the various d orbitals which could contribute selectively to a specific charge or spin property.

Angular-resolved photoemission (ARPES) measurements revealed the existence of 2DES with similar features to those seen at the LAO-STO interface at *bare surfaces* of several insulating perovskite oxide crystals, and among them (001) oriented STO [8, 9]. In this case, the carriers originate from oxygen vacancies. These vacancies are likely created during the sample preparation process and also when the sample is illuminated during the measurement [10]. Spin resolved ARPES (SARPES) of the 2DES at the (001) oriented STO surface [11] have highlighted the existence of sizable Rashba-like spin textures along with a large energy splitting that has been interpreted as a signature of ferromagnetism. Their simultaneous occurrence is puzzling since the two effects *a priori* compete with each other and estimates of the respective energy scales give about 100 meV for magnetism and a few meV for surface spin-orbit coupling.

A number of density functional theory (DFT) studies on oxygen-deficient bulk STO [12–14] found magnetism and in-gap bound states below the conduction band. The reported location of the bound states and the size of the spin polarization depend strongly on the computational details. Recent non-magnetic DFT calculations for slab geometries [15, 16] found the formation of a 2DES at the surface, in addition to in-gap states. Effects of spin orbit coupling [17, 18] and correlation [19] have also been studied at the model level.

In the present work, we investigate via first principles DFT the spin textures and magnetism of the 2DES states at the (001) oriented surface of oxygen-deficient STO. As the preexisting literature on this problem is already large, let us emphasize that the new results reported here are relevant for the interpretation of the following experimental facts: (i) spin-ARPES appears to highlight the occurrence of both Rashba-like textures and ferromagnetism [11]; and (ii) ARPES spectra pertaining to 2DES at interfaces and also at surfaces all seem to appear con-

comitant with a universal non-dispersive feature at an energy of about 1.3 eV below the Fermi energy [20–22]. Our DFT calculations are accompanied by tight-binding model considerations.



FIG. 1. (a) SrO-terminated $2 \times 2 \times 4$ slab with one oxygen vacancy at the topmost level. O_{α} denotes the position where a second vacancy is introduced for divacancy $2 \times 2 \times 4$ calculations (see text). (b) TiO₂-terminated $3 \times 3 \times 4$ slab with two vertically positioned oxygen vacancies. Other slab geometries are included in the Supp. Inf.

Methods.- In order to extract the basic trends in the electronic properties we considered three representative slabs of (001) oriented STO with different oxygen vacancy concentrations and slab terminations in our DFT calculations; a single vacancy at the topmost level of a SrO-terminated $2 \times 2 \times 4$ slab (Fig. 1 (a)), a vertical divacancy at the topmost level of a SrO-terminated $2 \times 2 \times 4$ slab (Fig. 1 (a)), and a horizontal divacancy located at the topmost level of a TiO₂-terminated $3 \times 3 \times 4$ slab (Fig. 1 (b), [23]). These slabs are chosen as test cases for investigating the effects of surface reconstruction, in-gap states, Rashba spin-orbit coupling and magnetism. We have made sure with a larger set of slabs (not shown) that all observations discussed below do not crucially depend on TiO_2 versus SrO termination. In all slabs we included a vacuum layer of at least 20 Å to avoid any spurious interactions between the periodic images. To our knowledge, no measurements of vacancy densities have been performed so far; therefore we have chosen our test slabs, such that the surface vacancy induced carrier densities are comparable to the experimental observations of $n_{2D} \approx 0.25 - 0.33 \text{ e}^{-}/\text{a}^{2}$ (see f.i. [8, 9, 24]). Every vacancy nominally releases two electrons. Due to the in-gap state not all the additional electrons are contributing to the 2DES, so that we can estimate a required density of about one vacancy per 4 unit cells. Our mono- and divacancy $2 \times 2 \times 4$ and $3 \times 3 \times 4$ slabs are therefore compatible with the experimental estimates.

In order to account for possible surface reconstructions, the internal coordinates of the slabs were relaxed with the projector-augmented wave basis [25] as implemented in VASP [26, 27]. We used the generalizedgradient approximation (GGA) [28] in the Dudarev [29] GGA+U scheme as described in Ref. 15. The electronic structure was analyzed with the all-electron full-potential local orbital (FPLO) [30] method and GGA+U functional [31]. Checks were also performed with the linearized augmented plane wave method as implemented in Wien2k [32]. Spin textures for the various slabs were obtained from full relativistic calculations in FPLO (GGA+SO+U) with a newly implemented subroutine.

Results and discussion.- We start with the analysis of spin textures in the absence of magnetism. In Figs. 2(a) and 2(d) we show the spin textures (projections of the spin polarization vectors onto the xy plane) at the Fermi surface $(k_z = 0)$ obtained for non-magnetic ground states in GGA+SO+U calculations for the monovacancy $2 \times 2 \times 4$ and the divacancy $3 \times 3 \times 4$ slabs, respectively (results for the divacancy $2 \times 2 \times 4$ are shown in the Suppl. Mat.). We used typical values for the parameters $(U = 5 \text{ eV} \text{ and } J_{\text{H}} = 0.64 \text{ eV} \text{ on Ti } 3d \text{ orbitals [33]})$. In all cases, every two bands show a small energy splitting of a few meV due to the spin-orbit interaction. Spins at the Fermi surface $(k_z = 0)$ are fully polarized on the xyplane, oppositely oriented in the split bands. On some of the bands the spins are pointing clock- and anti-clockwise around Γ , which is a clear signature of the relativistic Rashba effect. On other bands we additionally notice a texture of rotating spins, born out of the more complex interplay between spin and orbital degrees of freedom [10, 34]. The rather small size of the spin-splitting contrasts with the large value reported in recent SARPES experiments [11].

Next, we consider solutions with ferromagnetic order and spin-orbit interactions. The ferromagnetic solution is indeed the groundstate of the systems we consider [35]. Fig. 2 (b) displays the spin-projected bandstructure obtained from spin-polarized GGA+SO+U for the monovacancy $2 \times 2 \times 4$ slab (similarly, results for the divacancy $2 \times 2 \times 4$ slab are shown in the Suppl. Inf.). We adopt the magnetic moment quantization axis along z but below we also discuss the case of a quantization along x. The size of the magnetic splitting can be inferred from the black arrows connecting the majority and minority spin bands. For a comparison to the experiment we have to consider the splitting of the light bands of d_{xy} character, as heavy bands have been silenced in the measurements [11]. The energy separation at the Γ point of the two spin-split d_{xy} bands originating from Ti_d (Fig. 1(a)) is of the order of 60 meV and in qualitative agreement to the experimental data [11] ($\Delta E \approx 100 \text{ meV}$).

In order to identify the microscopic origin of the peculiar electronic and magnetic features described above we plot in Fig. 3 the layer- and orbital-resolved GGA+SO+U bandstructure near E_F for the monovacancy $2 \times 2 \times 4$ slab and in Fig. 4 (a) we display the Ti $t_{2g} - e_g$ resolved magnetic moments as a function of the distance between Ti and O vacancy. We find that (i) the magnetic splitting of the light d_{xy} bands at the Fermi



FIG. 2. Spin-textures and spin-polarized bandstructures for the monovacancy $2 \times 2 \times 4$ slab (a)-(c) and the divacancy $3 \times 3 \times 4$ slab (d)-(f). (a) and (d) are nonmagnetic GGA+SO+U calculations with U = 5 eV, and (b), (c), (e), (f) are ferromagnetic GGA+SO+U calculations with $m \parallel \hat{z}$. Note that the large magnetic moments along \hat{z} can be inferred from the exchange splitting in (b) and (e); the shown spin textures are projections on the xy plane. Reciprocal space units are $2\pi/(2a) = 0.805$ Å⁻¹ for (a) and (c), $2\pi/(3a) = 0.536$ Å⁻¹ for (d) and (f), where a is the bulk STO lattice parameter. The different vector scales are chosen to enhance the visibility of the spin windings.



FIG. 3. Ferromagnetic GGA+SO+U Ti 3d bandstructure for the monovacancy $2 \times 2 \times 4$ slab with $m \parallel \hat{z}$. (a) Layer resolved. (b) Orbitally resolved. Note, that the thickness of lines is proportional to the strength of the 3d character on the bands (the heavy band at -0.4 eV is strongly hybridized with Ti 4s and 4p orbitals, see also the main text).

level is caused by itinerant electrons belonging to Ti located not in the immediate vicinity of the oxygen vacancy (e.g. Ti_d in Fig. 1 (a)) with Ti 3d magnetic moments of the order of 0.1 μ_B (see Fig. 4 (a)). (ii) Ti atoms neighboring the oxygen vacancy in the uppermost layer (Ti_a, Ti_b, Ti_c in Fig. 1(a)) have the largest magnetic moment (Fig. 4 (a)) and are mostly responsible for the heavy bands (Ti_a) and occupied states at higher binding energies (Ti_b, Ti_c, see Fig. 3). In particular, we observe an in-gap band of Ti e_g (d_{z²}) character hybridizing with Ti 4s and 4p corresponding to Ti_a. It sits at -0.4 eV in the single vacancy case and is shifted to about -1 eV in the 2×2×4 divacancy case (see Suppl. Inf.). The position of this band depends on the parameters U and $J_{\rm H}$ chosen for the GGA+U calculations and on the concentration and position of vacancies in the slab (see Suppl. Inf. and Ref. 16).

Interestingly, already with this minimal slab, we find a phenomenon of *atomic specialization*, i.e. there are two types of electronic contributions to magnetism: one from Ti atoms neighboring the oxygen vacancy that acquire rather large magnetic moments and are mostly located below the Fermi surface inducing in-gap states; and an-



FIG. 4. $t_{2g}-e_g$ resolved magnetic moments in GGA+SO+U=5 eV for the (a) monovacancy $2 \times 2 \times 4$ and (b) divacancy $3 \times 3 \times 4$ slab in the ferromagnetic $m \parallel \hat{z}$ setup. Note, that the moments shown do not contain hybridization contributions.

other, from those Ti atoms lying further away from the oxygen vacancies, that correspond to polarized t_{2g} itinerant electrons with small magnetic moments, which are responsible for the Rashba spin winding and the spin splitting at the Fermi surface. These remarkable effects will be even more pronounced in larger slabs.

We observe here the same phenomenon of atomic specialization as in the smaller $2 \times 2 \times 4$ slab; Ti atoms neighboring the oxygen vacancies (Ti_a and Ti_b in Fig. 1 (b)) acquire large magnetic moments (see Fig. 4 (b)) and are responsible for the in-gap states located at higher binding energies; while Ti atoms lying further away from the oxygen vacancy (Ti_c and beyond) contribute to the 2DES with itinerant electrons carrying small magnetic moments, which are responsible for the Rashba spin winding and the spin splitting at E_F [36].

Further insight into the atomic dichotomy can be gained through tight-binding cluster diagonalization (see Suppl. Inf.) of structures with various configurations of vacancies. In order to monitor the formation of the 2DES conduction band states and of the in-gap states, we adiabatically turn on the energy contributions that represent the effect of introducing a vacancy into the cluster. Features seen in DFT are qualitatively reproduced.

Based on our above discussion of each individual effect, we may infer that spin textures and spin splitting compete with each other in the t_{2g} bands. Fig. 2 (c) displays the spin texture at $k_z = 0$ obtained from spin-polarized GGA+SO+U calculations for the monovacancy $2 \times 2 \times 4$ slab. The spin texture shows signs of the Rashba winding but it is less pronounced than in the nonmagnetic case. Taking a closer look at the inner pockets in Fig. 2 (c) (blue and red circles centered at Γ) corresponding to the spin up and spin down projections (compare Fig. 2 (b)), we observe a significant Fermi momentum shift of the bands (~ 0.1Å⁻¹), which is of the same order of magnitude as the one observed in SARPES experiments [11]. As can be inferred from the now very small in-plane spin component ($P_{\parallel} \leq 0.04$), ferromagnetism is dominating the arrangement of the spins but the spin winding is still visible. The same features are observed for the divacancy $3\times3\times4$ slab (Fig. 2 (f)), confirming the general validity of the results. We have also checked the robustness of the results with respect to the choice of U in the GGA+U

tions (see Suppl. Inf.). Scanning superconducting quantum interference device measurements [3] on LAO/STO interfaces observed a preference of in-plane magnetic moments, which are ascribed to shape anisotropy. As this is also expected for pure STO surfaces, we additionally performed calculations with the magnetization axis along x; in this case spins are aligned in plane and the Rashba interaction is unable to rotate the spins to achieve a sign change for opposite k points. However, a canting of the spin polarization vectors away from the magnetic axis originating from the Rashba coupling can be still identified, in agreement with previous theoretical considerations [37]. Our results would be compatible with SARPES spectra which see both an almost pure Rashba spin texture and a full in-plane polarization if we assume that the measurements detect a signal from a large number of ferromagnetic domains at once. In this case the effect of magnetism on the alignment of the spin polarization vectors is averaged out and the measured spin texture is completely determined by spin-orbit coupling effects, restoring the sign change for opposite k points and implying that the main features are independent of the magnetic quantization axis.

functional and find that the variation of U introduces only quantitative changes in the ferromagnetic calcula-

Conclusions.- By performing full relativistic nonmagnetic and magnetic density functional theory calculations in the framework of GGA+SO+U on representative oxygen deficient $SrTiO_3$ slabs, we find the magnetic state to be the ground state and we observe clear signatures of *atomic specialization* of the electronic and magnetic contributions. Ti atoms neighboring the oxygen vacancies create e_q localized wavefunctions with large magnetic moments and are responsible for the presence of in-gap states at energies around -0.5 to -1 eV. The position of the in-gap states is influenced by the slab termination, the depth of the oxygen vacancy below the surface and by possible oxygen clustering. On the other hand, Ti atoms lying further away from the oxygen vacancy contribute with polarized t_{2q} it inerant electrons to the conducting 2DES and are responsible for the Rashba spin winding and the spin splitting at the Fermi surface observed in SARPES. Our calculations show that magnetism masks the Rashba effect by increasing the spin splitting of the t_{2a} orbitals and by modifying the individual spin orientation but it does not eliminate spin winding. Considering that an averaging of inhomogeneities near the surface of the measured sample is to be expected (e.g. a sea of 2DES interspersed with islands of magnetism perhaps mirroring a mixture of TiO_2 and SrO in the termination layer), SARPES measurements are explained by our calculations as the combined effect of Rashba and magnetism.

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