Electronic Structure

Dimensionality of mobile electrons at x-ray-irradiated LaAlO₃/SrTiO₃ interfaces

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Abstract

Electronic structure of LaAlO₃/SrTiO₃ (LAO/STO) samples, grown at low oxygen pressure and post-annealed ex situ, was investigated by soft-x-ray ARPES focusing on the Fermi momentum \(k_F\) of the mobile electron system (MES). X-ray irradiation of these samples at temperatures below 100 K creates oxygen vacancies (VOs) injecting Ti \(t_{2g}\) electrons into the MES. At this temperature the oxygen out-diffusion is suppressed, and the VOs should appear mostly in the top STO layer. The x-ray generated MES demonstrates, however, a pronounced three-dimensional (3D) behavior as evidenced by variations of its experimental \(k_F\) over different Brillouin zones. Identical to bare STO, this behavior indicates an unexpectedly large extension of the x-ray generated MES into the STO depth. The intrinsic MES in the standard LAO/STO samples annealed in situ, in contrast, demonstrates purely two-dimensional (2D) behaviour. The relevance of our ARPES data analysis is supported by model calculations to compare the intensity vs gradient methods of the \(k_F\) determination as a function of the energy resolution ratio to the bandwidth. Based on self-interaction-corrected DFT calculations of the MES induced by VOs at the interface and in STO bulk, we discuss possible scenarios of the puzzling 3D-ity. It may involve either a dense ladder of quantum-well states formed in a long-range interfacial potential or, more likely, x-ray-induced bulk metallicity in STO accessed in the ARPES experiment through a short-range interfacial barrier. The mechanism of this metallicity may involve remnant VOs and photocconductivity-induced metallic states in the STO bulk, and even more exotic mechanisms such as x-ray induced formation of Frenkel pairs.

1. Introduction

Transition-metal oxides (TMOs) presently play one of the forefront roles in theoretical and experimental condensed matter research (for entries see [1]). An involved interplay between the spin, charge, orbital and lattice degrees of freedom in these materials results in a wealth of phenomena interesting from the fundamental point of view and bearing potential for technological applications. These include rich electronic and magnetic phase diagrams, metal–insulator transitions, colossal magnetoresistance, ferroelectricity, multiferroicity, high-\(T_c\) superconductivity, etc. Interfaces and heterostructures of TMOs can add another dimension to their fascinating properties, giving rise to physical phenomena which cannot be anticipated from the properties of individual constituents, with the new functionalities having great promise for future device applications (see, for example, the reviews [2, 3]).

The interface between LaAlO₃ (LAO) and SrTiO₃ (STO) is a paradigm example of new functionalities that can be formed by interfacing TMOs. Although bulk LAO and STO are both band insulators, their inter-
face spontaneously forms a mobile electron system (MES) [2, 3]. Its high electron mobility co-exists with superconductivity, ferromagnetism, large magnetoresistance and other non-trivial phenomena which can in addition be tuned with field effect. The MES electrons are localized in the interfacial quantum well (QW) on the STO side and populate the Ti $t_{2g}$-derived in-plane $d_{xy}$ states and out-of-plane $d_{z^2}$ states [4–6]. The latter, furthermore, can be manipulated through artificial quantum confinement in thin STO layers [7]. Whereas in stoichiometric LAO/STO the MES is localized within a few layers from the interface and is therefore purely 2D, oxygen deficiency of STO can extend the MES to a much larger depth of more than 1000 Å, resulting in its essentially 3D character [8–10]. Two phenomena bearing key importance for virtually all physical properties of LAO/STO are (1) polaronic nature of the charge carriers, where strong electron–phonon coupling to the LO phonon mode reduces their low-temperature mobility by a factor of about 2.5, and coupling to soft phonon modes dramatically reduces mobility with temperature [11, 12], and (2) electronic phase separation where the conducting MES puddles are embedded in the insulating host phase [12–14].

X-ray irradiation can dramatically change electronic and magnetic properties of oxide materials, which is typically connected with creation of oxygen vacancies (VOs) [14–22]. In a simplified picture of oxygen-deficient (OD) STO, each VOs releases two electrons from the neighbouring Ti atom. One of them joins the MES formed by delocalized quasiparticles, which are Ti $t_{2g}$ derived, weakly correlated, non-magnetic and form large polarons [11]. The other electron stays near the Ti ion to form localized in-gap state (IGSs) at binding energy $E_{0} \sim \sim −1.3$ eV, which are Ti $e_g$ derived, strongly correlated, magnetic and are often viewed as small polarons. In a more elaborate picture, the electron distribution between the MES and IGSs depends on particular configurations of VOs [23] which tend to form ferromagnetic clusters [14]. The VOs have a dramatic effect on the transport properties of STO, with their concentration of only 0.03% transforming STO into metal [24]. This picture of dichotomic electron system in the bulk as well at the surfaces and interfaces of OD-STO can be described within the combination of density functional theory with explicit electron-correlation schemes, such as e.g. dynamical mean-field theory (DMFT) (see, for example, references [21, 25–29]). Recently, it has been experimentally confirmed by resonant photoemission (ResPE) [22]. The coexistence of the two radically different MES and IGS electron subsystems hugely enriches the physics of the OD-STO systems compared to the stoichiometric ones, critically affecting the conductivity, electronic [12–14] and magnetic [30, 31] phase separation.

Here, we use soft-x-ray angle-resolved photoelectron spectroscopy (ARPES) to explore the dimensionality of the MES generated by x-ray irradiation of LAO/STO heterostructures grown in oxygen-deficient conditions and post-annealed (PA-LAO/STO). Our analysis focuses on variations of the experimental $k_F$ across the Brillouin zones (BZs), with the $k_F$ values determined from the gradient of bandwidth-integrated spectral intensity. Although the VOs generated by x-rays at our low temperature of 12 K should be located in the top STO layer, we observe an unexpected 3D-ity of x-ray generated MES, indicating its large extension into the STO depth. This behaviour, identical from the LAO/STO samples grown and annealed under the standard protocol [11]. Their difference most important in the context of our ARPES study is that under x-ray irradiation at temperatures below $\sim 100$ K the post-annealed samples readily build up VOs in STO, which is associated with blue-light fluorescence [34, 35]. In the present experiment the samples were irradiated at 12 K. In the ARPES spectra acquired in parallel with the irradiation, build up of VOs was evidenced by gradual development of the initially absent IGS peak in the band gap and Ti$^{2+}$ component of the Ti $2p$ core levels, which are characteristic signatures of the VOs (for detailed analysis of the time evolution see [14]).

The exact mechanism of irradiation-induced creation of VOs at the LAO/STO interfaces and even bare STO surfaces is not yet quite understood. There are a few intriguing observations: (1) the x-ray generated VOs and corresponding MES can be observed only at low sample temperatures below $\sim 100$ K. Once created, they stay stable without further irradiation for at least tens of hours. Upon increase of temperature well above $\sim 100$ K, however, they quench and cannot develop even at relatively high synchrotron-radiation power densities (see below). This fact excludes that the classical thermally-activated diffusion of oxygen atoms out of STO

2. Sample preparation

Our LAO/STO samples with a LAO layer of the critical thickness 4 u.c. on top of TiO$_2$-terminated STO(100) were grown with pulsed laser deposition. We followed a non-standard growth protocol, where the STO substrate was annealed at 500 °C in vacuum, the O$_2$ pressure during the LAO deposition at 720 °C reduced to $8 \times 10^{-5}$ mbar, and the post-growth annealing for 12 h performed ex situ at a temperature of 500 °C [14, 22]. Although such samples are finally (nearly) stoichiometric on the oxygen elemental ratio, at least to Ti as evidenced by Ti 2p core-level spectra discussed below, their transport properties [32, 33] are quite different from the LAO/STO samples grown and annealed under the standard protocol [11]. Their difference most important in the context of our ARPES study is that under x-ray irradiation at temperatures below $\sim 100$ K the post-annealed samples readily build up VOs in STO, which is associated with blue-light fluorescence [34, 35]. In the present experiment the samples were irradiated at 12 K. In the ARPES spectra acquired in parallel with the irradiation, build up of VOs was evidenced by gradual development of the initially absent IGS peak in the band gap and Ti$^{2+}$ component of the Ti $2p$ core levels, which are characteristic signatures of the VOs (for detailed analysis of the time evolution see [14]).
be involved in the generation of the V$_{0}$S, because in this case the temperature dependence would be opposite. Instead, their creation and/or stabilization might in some way be linked to the cubic to tetragonal phase transition and concomitant creation of domains in STO below 105 K [36, 37]. The electric field in STO due to the interfacial band bending, separating the photoexcited electrons and holes [38] as well as the repulsion of the mobile photoexcited electrons from the localized ones staying at the V$_{0}$S [39] may also contribute to the stability of the x-ray generated V$_{0}$S and MES; (2) the V$_{0}$S can be quenched by exposure of the LAO/STO samples to x-ray irradiation in O$_{2}$ pressure of about 10$^{-7}$ mbar on a time scale of seconds, which is much faster than the generation of V$_{0}$S on a time scale of tens of minutes. The high efficiency of this reaction can be explained by that x-ray irradiation cracks the physisorbed O$_{2}$ molecules into atomic oxygen [19, 20]. We note that whereas for STO surfaces only absorption of O$_{2}$ molecules is sufficient to quench the V$_{0}$S [17], in our case of the buried LAO/STO interface their x-ray cracking is necessary. This observation can be explained, tentatively, by that oxygen would effectively penetrate through the LAO layer only in its atomic form; (3) the V$_{0}$-generation rate strongly depends on the sample growth protocol, including temperature and O$_{2}$ pressure during the substrate annealing and the LAO deposition, and in situ vs ex situ annealing. Remarkably, the LAO/STO samples grown under the standard protocol (ST-LAO/STO), including substrate annealing in O$_{2}$, high O$_{2}$ pressure during the LAO deposition and in situ annealing, are fairly immune to x-ray irradiation [11]. The susceptibility of the post-annealed samples to x-ray irradiation may be connected with a certain amount of structural defects remaining in them after the post-annealing despite their complete or nearly complete stoichiometric recovery. At the moment, it is difficult to reconcile all these observations in one single mechanism of creation and stabilization of the V$_{0}$S, and this is not a prime objective of this study.

3. Experimental and theoretical methods

Our SX-ARPES experiments used ResPE at the Ti L-edge in order to boost the signal from the Ti derived electron states at the buried LAO/STO interface. The measurements were performed at the soft-x-ray ARPES endstation [40] of the Advanced Resonant Spectroscopies (ADRESS) beamline [41] at the Swiss Light Source, Paul Scherrer Institute, Switzerland. The experimental geometry is sketched in figure 1(a). A photon flux of about 10$^{15}$ photons s$^{-1}$ was focused in a spot size of 30 × 75 μm$^2$ on the sample surface, and the combined (beamline and analyzer) energy resolution (∆E) was ~50 meV full width at half maximum (FWHM). The sample was kept at 12 K in order to suppress smearing of the k-dispersive spectral fraction [42] and, most importantly, allow a build-up of the V$_{0}$S under x-ray irradiation (see above). The x-ray absorption spectroscopy (XAS) data were measured in the total electron yield (TEY). All ARPES and XAS data presented below are acquired at saturation after more than 2 h of irradiation time. The XAS and angle-integrated photoemission data were measured with circularly polarized incident x-rays. The in-plane Fermi surface (FS) maps and ARPES band dispersion images presented below were measured with s-polarized x-rays as shown in figure 1(a), and the out-of-plane FS map with p-polarized x-rays.

Our electronic structure calculations for OD LAO/STO interfaces addressed possible MES contributions from V$_{0}$S in the top TiO$_{2}$ layer as well as in deeper-lying TiO$_{2}$ layers on equal footing as embedded in a LAO/STO supercell. The calculations utilized the STO-bulk lattice constant of a = 3.905 Å, and were structurally relaxed within the generalized-gradient approximation of Kohn–Sham’s density-functional theory (DFT) utilizing a mixed-basis pseudopotential code. The final electronic structure was determined with a DFT+self-interaction-correction (SIC) scheme, where explicit Coulomb interactions on oxygen sites are described within SIC, and are incorporated in the O pseudopotential [43]. The SIC was applied to the O 2s and the O 2p orbitals via weight factors w$_{p}$ (see reference [43] for more details). While the O 2s orbital is by default fully corrected with w$_{p}$ = 1.0, the common choice w$_{p}$ = 0.8 was used for O 2p orbitals. Since the Ti t$_{2g}$ states, mostly relevant for the MES, are only weakly occupied in LAO/STO, neglecting the explicit Coulomb interactions on Ti when focussing on the MES proves as an adequate approximation for the very large supercells. Note that when applied to bulk STO, the present DFT+SIC treatment results in a reasonable band gap of 3 eV, curing the notorious underestimation in standard exchange–correlation functionals.

4. Electronic structure overview

Figure 1(b) shows the Ti 2p$_{3/2}$ core-level spectrum of our PA-LAO/STO samples taken at $h\nu = 1000$ eV, which is decomposed into the Ti$^{4+}$ and Ti$^{3+}$ components (linear background removed). The latter is characteristic of the V$_{0}$S. (c) shows XAS spectrum through the Ti L$_{3}$- and L$_{2}$-edges (excited from the 2p$_{3/2}$ and 2p$_{1/2}$ core levels, respectively). The pairs of salient XAS peaks at each edge are formed by the Ti4+ e$_{g}$ states. The corresponding map of ResPE intensity as a function of E$_{F}$ and excitation energy $h\nu$, identifying the Ti-derived electron states [44–48] is displayed in figure 1(d). Above the broad valence band composed from the O 2p
states in STO and LAO, the map shows the resonating broad IGS peak at $E_B \sim -1.3$ eV which is a hallmark of the VO$_5$S. Recent analysis of its resonant behavior [22] confirms that the IGS-subsystem is derived from $e_g$ states of the Ti$_{3}^{+}$ ions pushed down in energy by strong electron correlations [25, 27, 28]. The narrow resonating peak at the Fermi energy ($E_F$) identifies the $t_{2g}$-derived MES. Its intensity maximum is shifted from the Ti$_{4}^{+}$ $t_{2g}$ to Ti$_{4}^{+}$ $e_g$ peak in the XAS spectrum due to remnant $k$-conservation between the intermediate and final states coupled in the ResPE process [22].

We note that Ti$_{3}^{+}$ $e_g$ and $t_{2g}$ signals in the XAS spectrum in figure 1(c), corresponding to the Ti atoms hosting the VO$_5$S, are almost invisible. At the same time, the IGS signal in the ResPE map (d), corresponding to the Ti$_{3}^{+}$ $e_g$ states of these atoms, is profound. This observation shows that most VO$_5$S are located in vicinity of the top STO layer, because the width of this region is much smaller than the probing depth of the TEY-XAS measurements of the order of 50 Å, and comparable with that of SX-ARPES measurements of the order of 11 Å at our excitation energy [20].

The image (e) in figure 1 shows the ARPES intensity measured along the $\Gamma X$ direction with $s$-polarized incident x-rays. With the experimental geometry shown in figure 1(a), this polarization selects predominantly the antisymmetric $d_{xy}$- and $d_{yz}$-derived electron states [6, 11, 22]. Furthermore, the choice of $h\nu = 466.4$ eV at the $L_2$ edge enhanced the $d_{yz}$ states in comparison with the $d_{xy}$ ones [22]. The ARPES data is overlaid with a sketch of the $d_{xy}$ and $d_{yz}$ dispersions. We note pronounced spectral intensity variations along the bands, caused by energy- and $k$-dependent photoemission matrix elements. Particularly remarkable is the spectral intensity asymmetry relative to the $\Gamma$ points, absent in VUV-ARPES on bare STO surfaces [18, 49]. With $s$-polarized x-rays in our experimental geometry, this phenomenon is beyond the conventional dipole approximation for the photoemission process (because the angle between the photoemission direction and electric vector of the light is the same) and should be attributed to the photon momentum that is $\sim 0.24$ Å$^{-1}$ for our $h\nu$ in the soft-x-ray energy range. We observe that for PA-LAO/STO the lowest $d_{xy}$ band is much smeared compared to STO, and manifests itself as intensity hot spots where it intersects the $d_{yz}$ band and hybridizes with it because of the symmetry breaking caused by the tetragonal lattice distortion and spin–orbit coupling. In view of the location of these $d_{xy}$ states in the top TiO$_2$ layer, the observed smearing should be connected with a disorder induced in this layer by the LAO overlayer and x-ray generated VO$_5$S. As we will see below, the absence of high-order $d_{xy}$
bands in PA-LAO/STO is consistent with the steeper electrostatic band-bending potential \( V(z) \) compared to bare STO caused by the additional electron density injected by the V\(_{\text{Os}}\). We note that the band order and band dispersions observed in PA-LAO/STO (and STO) and ST-LAO/STO samples \([6, 11]\) demonstrate the same \( d_{xy} < d_{x^2−y^2} \) band order. However, the observed band filling in PA-LAO/STO (STO) is somewhat larger than in ST-LAO/STO. Furthermore, the polaronic-hump weight and thus renormalization of the effective mass \( m^* \) in PA-LAO/STO (STO) reduce to \(~1.5\) compared to \(~2.5\) in ST-LAO/STO [11]. Importantly, as we will see below, these materials demonstrate different dimensionality of the MES. Finally, figure 1(f) shows the experimental FS of PA-LAO/STO where we observe the circular \( d_{xy} \) and elliptical \( d_{yz} \) sheets.

5. Dimensionality of the MES

Do the MES electrons in PA-LAO/STO, developing under irradiation, stay confined at the interface and keep their 2D character, or expand into the STO bulk and become 3D? To answer this question, we investigated their ARPES response as a function of out-of-plane momentum \( k_z \) varied in an extended \( hv\)-range above the Ti \( L\)-edge. Because of the much weaker off-resonance response of the MES, such measurements are extremely photon hungry. We used \( p\)-polarization of incident x-rays to switch from the two-band combination of the \( d_{xy} \) and \( d_{yz} \) antisymmetric states to the single-band \( d_{xz} \) state. Figure 2 shows an experimental map of ARPES intensity at \( E_k \) as a function of \( hv \), rendered into the out-of-plane momentum \( k_z \) according to the relation

\[
k_z = \sqrt{\frac{2m(E_k + V_0)}{\hbar^2}} - k_{\parallel} + \frac{p_\gamma}{\sqrt{2m}} ,
\]

where \( m \) is the free-electron mass, \( E_k \) photoelectron kinetic energy, \( V_0 \) the inner potential set to 15 eV, and \( p_\gamma \) the photon momentum correction in our experimental geometry [40].

The map in figure 2 clearly shows FS contours of the \( d_{xz} \)-derived interfacial states whose intensity periodically blows up when \( k_z \) hits the \( \Gamma \)-points of the BZ. However, this pattern alone does not necessarily indicate a 3D-ity of these states, because 2D states formed by the out-of-plane \( d_{xz} \)-orbitals confined in the interfacial QW would also produce periodic intensity oscillations. The only difference is that the out-of-plane dispersion of the 3D states will manifest itself as rounded FS contours in the \((k_x, k_z)\) coordinates, and the absence of it for the 2D states as straight FS contours (for in-depth analysis of ARPES response of 2D states see [50]). In our case, relatively poor statistics and variations of the photoemission matrix element with \( hv \) and \( k_z \) do not allow reliable discrimination between these 2D and 3D patterns of the \( k_z \)-dependent ARPES intensity from the MES. As the acquisition of this dataset has already required more than 5 h, further increase of the acquisition time for these off-resonance measurements is less practical.

In order to confirm the 3D-ity of the x-ray induced MES, we returned to the resonant \( hv = 466.4 \) eV delivering maximal intensity to the \( d_{xy} \) states, and investigated the variation of the apparent \( k_{\parallel} \) as a function of \( k_{\parallel} \) when moving to the next BZ. The measured ARPES image through two BZs is presented in figure 4(a). As in our case the occupied bandwidth \( W \sim 50 \) meV is basically equal to the experimental \( \Delta E \), accurate determination of \( k_{\parallel} \) is not trivial. Our ARPES simulations, described in the appendix, have demonstrated that whereas the conventional method to determine \( k_{\parallel} \) from the peaks of spectral intensity at \( E_k \) (intensity method) stays accurate only as long as \( W \) is much larger than \( \Delta E \), the extremes of the gradient \( dI/dk \) of the bandwidth-integrated spectral intensity (gradient method) yield accurate \( k_{\parallel} \) values even when \( W \) is of the order of or even smaller than \( \Delta E \). The experimental \( dI/dk \) (in our case integrated over an energy window of 100 meV, with the excluded polaronic introducing a negligible correction) is shown in figure 3(b). Relevant for our MES dimensionality analysis is the \( d_{xz} \) band, where the \( dI/dk \) extremes marked \( k^0 \) and \( k^1 \) identify its apparent \( k_{\parallel} \) (for the \( k_{\parallel} \)-determination method see appendix). Remarkably, it reduces from \( k^0 \sim 0.44 \) Å\(^{-1}\) in the first BZ to \( k^1 \sim 0.29 \) Å\(^{-1}\) in the second one (hereinafter the accuracy of all \( k_{\parallel} \) values, including the statistical and instrumental errors, is about \pm 0.025 Å\(^{-1}\)). We note that on the left side of \( \Gamma_6 \) and right side of \( \Gamma_7 \) the spectral intensity from this band vanishes before reaching \( E_F \), presumably due to a sharp variation of the photoemission matrix elements with \( k_z \) and cannot be used to determine \( k_{\parallel} \). For another PA-LAO/STO sample, where the post-growth annealing time was increased to 24 h, we found a somewhat smaller \( k^0 \sim 0.40 \) Å\(^{-1}\) that anyway reduced to \( k^1 \sim 0.29 \) Å\(^{-1}\) in the second BZ. Furthermore, we have compared our results on PA-LAO/STO with previous resonant SX-ARPES measurements on bare STO (supplementary in reference [22]) and have found in this reference system remarkably similar \( k^0 \sim 0.40 \) Å\(^{-1}\) and \( k^1 \sim 0.29 \) Å\(^{-1}\).

This difference of the apparent \( k_{\parallel} \) between the two BZs is actually a clear manifestation of the 3D character of the \( d_{xy} \) band in both PA-LAO/STO and STO. This situation is sketched in figure 3(c), showing the \( d_{xy} \)-derived FS sheet measured in our experiment. In this case the \( k_{\parallel}=\)coordinate of \( k \) indeed depends on \( k_z \), which decreases as \( k_{\parallel} \) increases into the next BZ. The effect of the relatively small change of \( k_{\parallel} \) between the two BZs on the apparent \( k_z \) is amplified by the strong elongation of the FS sheets along the \( k_z \)-axis, with light- and heavy-electron axes being \sim 0.12 and 0.8 Å\(^{-1}\), respectively, as measured in the previous polarization-dependent studies on for PA-LAO/STO [14, 22] and STO [17, 18]. We note that our \( hv = 466.4 \) eV at the Ti \( L_2\)-edge resonance corresponds to \( k_z = 6.99 (2\pi/a) \), i.e. brings \( k_z \) almost ideally to the \( \Gamma \)-point of bulk STO where the
Figure 2. The \(d_{xz}\)-derived FS in PA-LAO/STO as a function of \(k_z\) in an extended \(hv\)-range. The corresponding \(hv\)-values are marked. The intensity is normalized to the angle-integrated intensity for each \(hv\). The 3D-ity of the out-of-plane MES states, suggested by rounded FS contours, is confirmed by the \(k_y\) variations across the BZs in figure 3.

Figure 3. 3D-MES in PA-LAO/STO: (a) resonant ARPES image through two BZs measured at \(hv = 466.4\) eV; (b) corresponding gradient \(dI_W/dk\) of the bandwidth-integrated intensity, where the dashed lines mark \(k_y\) of the \(d_{yz}\) bands. As sketched in (c), the difference in the apparent \(k_y\) of the \(d_{yz}\) bands between the two BZs identifies the out-of-plane dispersion and thus 3D-MES; (d) schematics of the sharp interfacial band bending, screened by the additional electrons supplied by the VO s, and of the corresponding wavefunctions of the \(d_{xy}\) and \(d_{xz}\) states. The \(t_{2g}\) electrons, propagating into the STO bulk, form the 3D-MES.

MES is located. Had the \(L\)-edge intra-atomic resonance energy not coincided with the \(\Gamma\)-point, the spectral intensity from the MES would have hugely dropped, and the ARPES data acquisition would have required much longer acquisition time similarly to the dataset presented in figure 2.

Our SX-ARPES results on PA-LAO/STO and STO can be compared to the VUV-ARPES results of Plumb et al [49] on STO surfaces prepared by annealing in vacuum. In this case, the emergent MES can be related to the VO s only [14–22], without any intrinsic polar-discontinuity contribution. In line with our findings, the experimental \(k_z\) dispersions measured in an \(hv\) range below 100 eV have also demonstrated the 2D-ity of the Ti \(d_{xy}\) states and 3D-ity of the \(d_{xz}/yz\) ones, with the latter forming closed contours in the out-of-plane FS map [49]. An indirect confirmation of the 3D character of the x-ray generated MES is an observation that such MES has a higher mobility compared to the intrinsic one [38] that can be attributed to its extension from the interface into the defect-free STO bulk. Furthermore, spectroscopic signatures of delocalized charge carriers,
that can be associated with the MES, have also been observed with resonant x-ray scattering (RIXS) \cite{51, 52}; given the large probing depth of RIXS, the sizable magnitude of this spectral feature is consistent with the large spatial extension of the MES.

Remarkably, the ST-samples show a qualitatively different pattern of the ARPES variations with $k_{ll}$. Figure 4(a) represents an ARPES image generated from the dataset of reference \cite{11} which was acquired for an ST-LAO/STO sample with a LAO thickness of 4 u.c. under the same conditions of the ARPES experiment as above. Although the spectral intensity of the $dyz$ bands again strongly varies through the BZs, the corresponding $dW/dk$ extremes (b) show constant $k_F \sim 0.33 \text{ Å}^{-1}$, identifying the 2D-ity of the MES. In passing, we note a significantly larger polaronic renormalization of the $dyz$ dispersion at the ST-LAO/STO interfaces compared to PA-LAO/STO, where the electron–phonon interaction is screened by the larger MES density. Also the two hot spots of the ARPES intensity, where the $dxy$ band intersects the $dyz$ one, are more pronounced for ST-LAO/STO; the larger smearing of the $dxy$ states in PA-LAO/STO can manifest a larger disorder in its top STO layer due to the random x-ray generated VO$_x$ in this region.

We note that recent results by Soltani et al.\cite{53} suggest that a 2D-MES may also be realized at bare STO surfaces. This has been identified from a splitting of the $dz^2/ryz$-derived states, suggesting their quantum confinement and thus 2D-ity. Obviously, the dimensionality of the MES is determined by the depth and extension of the surface/interfacial QW which, in turn, is sensitive to the stoichiometry and preparation of the samples, which was \textit{in situ} cleavage in the experiments by Soltani et al.\cite{53} vs \textit{in situ} annealing in the experiments by Plumb et al.\cite{49}.

6. Discussion

6.1. Bulk and interfacial oxygen deficiency in STO

The VO$_x$s, developing under x-ray irradiation, should be one of the key factors of the observed 3D-ity of the MES in PA-LAO/STO, in contrast to the conventional 2D-MES in ST-LAO/STO. First of all, we note that LAO/STO interfaces grown on top of OD-STO have previously been studied by a number of techniques other than ARPES such as magnetotransport \cite{8} and conducting-tip atomic force microscopy of the interfacial cross-section \cite{9}. It has been noted that their $n_s$ can be two-three orders of magnitude larger than $n_s = 0.5e$ per u.c. area predicted by the electrostatic arguments. This fact alone necessitates the existence of an extended and thus 3D component of the MES in OD-samples, and it has indeed been observed that sufficiently high concentrations of VO$_x$s result in a dimensionality transformation of the MES from 2D to 3D \cite{8–10}. Electron mobility in these systems could significantly exceed that of the paradigm 2DES in ST-LAO/STO \cite{8, 32, 54}; this phenomenon could trace back to stronger electron screening of the polaronic interactions in LAO/STO by larger electron density \cite{11}. However, the VO distribution profile in these OD-samples would extend into STO by a few thousands of
Å and more [55], also affected by oxygen trapping and diffusion at domain walls (see, for example, references [56–58]).

In contrast to these initially-OD samples, the post-annealing of our samples in oxygen, albeit ex situ, ensures that before the x-ray irradiation they were at most stoichiometric. As discussed above, this fact is evidenced by the initial absence of the IGS peak and Ti3+ core-level component in the ARPES spectra. The x-ray generated VOs should then locate mostly in the top layer of STO because, while easy out-diffusion of oxygen atoms from STO through the LAO overlayer can be explained by typically relaxed crystallinity of the latter [59], the out-diffusion from deeper layers of STO should be practically prohibited due to vanishing diffusion coefficient of oxygen at our low sample temperature. Our case should therefore be different from creation of VOs by deposition of thin metal films on STO at higher temperature where the VOs could be distributed over a depth of ~1 nm [60, 61]. This top-layer location of the VOs in our case is consistent with the above observation that the Ti3+ spectral weight in the TEY-XAS spectra in figure 1(a), which probe a significant depth into STO, is much smaller than in the core-level photoemission spectra in (b), which probe a much narrower vicinity of the interface. Furthermore, the top-layer location of the VOs is consistent with the angular dependence of the Ti3+ core-level component in hard-x-ray photoemission experiments by Sing et al [62].

### 6.2. Extended-QW scenario

In view of the 2D-ity of the x-ray generated VOs, the observed 3D-MES in our PA-LAO/STO samples seems puzzling. Which particular kind of the electron states and V(z) shape is needed to realize such 3D-MES? We will start from the well-known picture of the band bending at the ST-LAO/STO interface [4, 6, 55] sketched in figure 3(d). The in-plane dxy orbitals from each TiO2 plane form a sequence of the dxy electron states localized in the out-of-plane direction, whose energies climb V(z) to cross Ek at a finite distance from the interface. The out-of-plane dxy orbitals from each TiO2 plane hybridize into electron states delocalized in the out-of-plane direction, which quantize in V(z) to form a ladder of the the dxy states (where normally only the lowest one is occupied). With no occupied t2g states propagating into the STO bulk, the MES confined in the interfacial V(z) is purely 2D.

One possibility to realize the 3D behaviour (or rather quasi-3D in this particular context) is to increase the depth extension of V(z), which we will call the extended-QW scenario. In this case the 3D states will form as the convergence limit of a ladder of 2D states, quantized in the long-range confining potential, as the extension of this potential increases. Such a 2D- to 3D-dimensionality transformation has previously been demonstrated, for example, for multilayer graphene [63] and anatase TiO2 [64]. A similar analysis for the bare STO surface [65] has suggested that whereas a confining potential with an extension of a few monolayers can form a 3D-like ladder of QW states out of the dxy states having large perpendicular m∗, it forms only one QW state out of the dxy states with their large interlayer hopping and thus smaller perpendicular m∗. This result for the STO surface, however, is at odds with the experimental kF dispersions demonstrating that, other way around, the dxy states are 2D [18, 49] and the dxy ones are 3D [49]. These experimental results go in line with our observations of the 3D-ity of the dxy states in PA-LAO/STO.

Another way to reconcile the extended-QW scenario with our experimental results might be if the MES contribution injected by the top-layer VOs is significantly more delocalized than the intrinsic polar-discontinuity contribution. To check this scenario, we have performed supercell DFT + SIC electronic structure calculations where a 400-atom supercell incorporating 16 TiO2 layers, 6 LaO layers and a (2 × 2) interlayer resolution in a superlattice geometry was employed. As shown in figure 5(a), the oxygen deficiency was simulated with a VOs placed in the top TiO2 layer. Figure 5(b) presents the results of this calculation (red line) in comparison with the stoichiometric interface (blue). We observe that although the VOs somewhat enhances the asymptotics of the MES towards the STO bulk, it essentially does not change its overall depth extension of ~2 u.c. compared to ST-LAO/STO, where our ARPES experiment has found a purely 2D intrinsic MES.

### 6.3. Bulk-metallicity scenario

Another possibility to realize 3D-MES would be that under x-ray irradiation the whole STO depth within the light absorption depth becomes metallic, and the depth extension of V(z), other way around, shrinks to enable the ARPES experiment to directly access the bulk electronic structure. This bulk-metallicity scenario is sketched in figure 3(d). In our case such short-range V(z) seems more relevant than the long-range one in the previous scenario, because the top-layer VOs release additional MES electrons, as manifested by the observed increase of both kF and lateral conducting fraction [14] under x-ray irradiation. The increased electron density will then more effectively screen the interfacial-potential discontinuity, reducing the depth over which V(z) saturates to below the probing depth in our ARPES experiment. If upon further progression into the STO bulk the CBM stays below Ek, the flat V(z) potential in this region forms the 3D-MES. This scenario explains the 3D-ity of the dxy states observed by the ARPES experiment in PA-LAO/STO. Due to the flat potential, in the STO bulk the dxy states degenerate with the dxy. Importantly, the bulk metallicity in STO required for such a scenario can
Figure 5. DFT+GGA calculations: (a) the supercell incorporating VO in the top and bulk TiO₂ layers; (b) calculated layer-resolved MES density for the stoichiometric and OD-LAO/STO interfaces. Note that only the Ti t₂g contribution enters here, since this is the major contribution to the MES.

only be realized if the x-ray irradiation causes some effective doping throughout the STO bulk. As we discuss below, such metalliclicity is essentially the well-known (albeit poorly understood) photoconductivity observed in STO-based systems. The scenario of short-range V(z) combined with the bulk metalliclicity naturally extends from PA-LAO/STO to bare STO surfaces; the 3D-MES in that case cannot be explained only by the VO [19, 20] or any atomic rearrangements [49] in the STO layer.

One possibility for the bulk metalliclicity of PA-LAO/STO might be that already before the x-ray irradiation the deeper STO layers contained a minute amount of VO below the sensitivity limit of our ARPES and XAS experiments. Figure 5(b) (green line) shows our calculations which included, in addition to the VO in the top STO layer, another VO in the STO-slab centre between the eighth and ninth TiO₂ layer, representing the STO bulk. These results indicate that the extension of the MES induced by the bulk VO is similar to the interfacial ones, restricted within a sphere with a radius of ∼2 u.c. The corresponding wavefunctions will start to overlap, forming the 3D character of the MES, at a concentration of only 1.5% which is below the detection limit of our experiment. The previous DFT calculations by Li et al. [10] showed a roughly twice larger spatial extension of the VO-induced MES, with the difference to our calculations plausibly attributed to the neglect of electron correlations. Transport measurements [24] indicate a yet smaller VO-concentration of 0.03% which is necessary to transform STO into 3D metal. Such a minute amount of VO could likely stay in our samples after their post-annealing. With STO substrates being typically slightly OD, the same mechanism could be at play to form the 3D-MES at bare STO surfaces. We note that the above picture of the photoinduced DX-centers as electron donors forming an extended MES is actually similar to the conventional semiconductors, where tiny dopant concentrations on the promille level can form delocalized electron gas. The exhaustive picture of the OD-LAO/STO interfaces, including the entangled phenomena of the interfacial electrostatic field, VO, intrinsic and VO-induced MES and photoconductivity, still awaits accurate theoretical description.

The bulk metalliclicity in STO suggested by this scenario may be mediated by the well-known giant photoconductivity of STO-based systems (see, for example, references [34, 37, 66, 67]) that may exceed that in semiconductors by few orders of magnitude. This phenomenon strongly depends on the sample preparation, with the defects including the VO as well as the cubic-tetragonal phase transition [37] and lattice relaxation [39] playing the main role, and for some LAO/STO [66, 67] and OD-STO [68] samples can even become persistent on a time scale of up to days and more. In our experimental setup, we observed the photoconductivity as a drain current to the sample holder through the whole 0.25 mm thickness of our nominally insulating STO substrates which switched on and off under x-ray irradiation. Although the exact physics of photoconductivity in STO remains elusive yet, it is often associated with DX-centers (donor defect) located ∼200 meV below the conduction-band minimum. This energy position would allow the photoinduced (in our case x-ray induced) electron states to effectively hybridize with the MES wavefunctions, providing the coupling between the VO-induced MES puddles in STO necessary for the 3D-ity of the whole system. Therefore, although the photoconductivity states, disordered and having relatively small electron density, are less visible in ARPES, they may be a crucial element of the observed 3D-ity.

Although the above mechanisms explain the emergence of the 3D electron states, the concentration of VO and the density-of-states of the corresponding MES in the deeper STO layers stays negligible compared to the interfacial region. Therefore, these mechanisms cannot directly explain the whole large density of the
x-ray generated 3D-MES. One possible explanation might be that x-rays generate VO$_{1s}$ in STO bulk through creation of Frenkel pairs, where the O atom would become interstitial without the necessity to diffuse over a large distance. This behavior may be facilitated by the repulsion of the x-ray generated mobile electrons from the ones localized at the VO$_{1s}$ [39]. Furthermore, the Frenkel-pair mechanism would be consistent with the fact pointed out above that an increase of temperature above $\sim$100 K quenches the spectroscopic signatures of the VO$_{1s}$. However, we could not computationally confirm or rule out this scenario within any reasonable computational effort. Full understanding of the x-ray induced formation of the 3D-MES in LAO/STO requires therefore further experimental and theoretical investigations. Remnant cation non-stoichiometry on both LAO and STO sides, however small, can in principle also be involved in the observed dimensionality effects.

7. Summary

We used soft-x-ray ARPES to investigate electronic structure of LAO/STO samples grown at low oxygen pressure and post-annealed ex situ till their (nearly) complete stoichiometric recovery. Under x-ray irradiation at low temperatures below $\sim$100 K, the ARPES spectra of these samples show a rapid development of the Ti$^{3+}$ component of the Ti 2$p$ core levels and IGS peak, characteristic of the VO$_{1s}$, in parallel with scaling up of the MES spectral weight. Given that at this temperature the oxygen out-diffusion is suppressed, these VO$_{1s}$ should be located mostly in the top STO layer. However, the out-of-plane electron dispersions evaluated from the off-resonant ARPES data as well as the variation of the $k_F$ values over the BZs, evaluated by the gradient method from the Ti 2$p$ resonant data, evidence that the x-ray generated MES in our PA-LAO/STO samples is 3D, extending deep into the STO bulk. This behavior, essentially identical to the x-ray generated MES in STO, is clearly distinct from the intrinsic MES in ST-LAO/STO samples, fairly immune to x-ray irradiation, where our ARPES experiment finds pure 2D-ity of the MES. Our DFT+SIC calculations indicate that the spatial extension of the MES induced by VO$_{1s}$ in the STO bulk as well as in the top STO layer is about 2 u.c. which can hardly explain the 3D-ity of the MES in PA-LAO/STO. One scenario of this phenomenon, the extended-QW one, involves formation of a dense ladder of QW states in a long-range interfacial potential. More likely in our case is however the bulk-metallicity scenario of the observed 3D-ity, where the x-ray-induced bulk metallicity in STO is accompanied by a short-range interfacial $V(z)$ allowing the ARPES experiment to directly access the bulk states. In this case the formation of the 3D-MES may involve the minute concentrations of VO$_{1s}$ remnant in the STO bulk and the well-known photoconductivity of STO, and even more exotic mechanisms such as x-ray induced formation of Frenkel pairs. The short-range character of $V(z)$ can be caused by a large concentration of the x-ray generated VO$_{1s}$ and thus MES density in the top STO layer, where it effectively screens the interfacial $V(z)$ discontinuity.

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Data availability statement

The data that support the findings of this study are available upon reasonable request from the authors.

Appendix. Gradient vs intensity method for the $k_F$ determination

A crucial element of our further analysis of the ARPES data is accurate determination of the MES’ $k_F$. In fact, its signatures in the ARPES intensity can be strongly distorted not only by sharp variations of the photoemission matrix element but also by broadening of the experimental spectra due incoherent electron scattering (including defects) and limited experimental resolution in $k$ and energy. Conventionally, one determines from the maxima of the momentum-distribution curve of the spectral intensity at $E_F$ (Fermi intensity $I_F$). However, Straub et al [69] have shown that the applicability of this so-called intensity method is restricted to wide-band systems, where the occupied quasiparticle bandwidth $W$ much exceeds $\Delta E$ the experimental resolution, and that in the general case, including narrow-band systems, extremes of the gradient $dI_W/dk$ of the spectral intensity integrated through the whole $W$ resolve $k_F$ with much higher accuracy. In fact, this so-called gradient
Figure A1. Determination of $k_F$ at different experimental $\Delta E$ vs the occupied bandwidth $W$: (a)–(c) band dispersion superimposed with the ARPES intensity (bottom) and the corresponding characteristics of $k_F - I_F(k)$, bandwidth-integrated $dI_W/dk$, and $dI_F/dk$—for different $\Delta E/W$ values; (d) deviation of the peaks in these curves from the true $k_F$ as a function of $\Delta E/W$. The $dI_W/dk$ peaks deliver the most accurate $k_F$ value through all $\Delta E/W$ values.

method is based on the theoretical many-body definition of the Fermi surface (FS) as restricted by the extremal gradients of the momentum distribution function $n(k)$ [69].

To identify the optimal $k_F$-determination method in our case, we performed ARPES simulations for a 2D (or 2D cross-section of 3D) parabolic band whose parameters $W = 50$ meV and $k_F = 0.4$ Å$^{-1}$ closely resembled the $dyz$ band along $k_x$. With the electron lifetime broadening being negligible close to $E_F$, the corresponding spectral function $A(E, k)$ was modelled as the $\delta$-function centered at the band's dispersion $E(k)$ and terminated at $E_F$. Its ARPES response was simulated by Gaussian convolution with FWHM = 50 meV in the $E$-direction describing the experimental $\Delta E$, and 0.1 Å$^{-1}$ in the $k$-direction describing the combined effect of disorder and experimental $k$-resolution.

The results of these simulations, corresponding to the case $W = \Delta E$, are presented in figure A1(b) as the ARPES intensity with the corresponding $I_F(k)$ and $dI_W/dk$ curves on top. Whereas the intensity method to determine $k_F$ from the peaks of $I_F$ underestimates its value as much as 14%, the extremes of the gradient $dI_W/dk$ yield its value with a remarkable precision of $\sim 0.25%$. Important for the gradient-method accuracy is that the spectral intensity is integrated over the whole bandwidth, because the gradient $dI_F/dk$ of the Fermi intensity only, also shown in figure A1(b), overestimates the actual $k_F$ by 2.5%. Therefore, our ARPES simulations clearly demonstrate that the gradient method to determine $k_F$ is optimal in our case.

Our ARPES simulations extended to smaller and larger $\Delta E/W$ values are shown in figures A1(a) and (c), respectively. Obviously, the intensity method improves its accuracy towards small $\Delta E$, whereas the $I_F$ gradient does so towards large $\Delta E$ compared to $W$. This trend calculated over a wide range of $\Delta E/W$ is presented in the panel (d) which shows the relative deviation of the $I_F(k)$, $dI_W/dk$ and $dI_F/dk$ extremes from the true $k_F$. A reduction of the spectral $k$-broadening will increase the accuracy of the intensity and $I_F$-gradient methods, in particular of the latter. However, the $dI_W/dk$ gradient method stays remarkably accurate through the whole range of $\Delta E$ compared to $W$, with a negligible rms deviation of only 0.3% even with the relatively large $k$-broadening used in our simulations. The gradient method to determine $k_F$ can therefore be deemed the most universal. An added advantage of this method is that the narrower width of the $dI_W/dk$ peaks compared to $I_F(k)$ makes it less susceptible to the photoemission matrix-element variations with $k$.

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