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Quasiparticle dynamics and spin-orbital texture of the SrTiO₃ two-dimensional electron gas

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Two-dimensional electron gases (2DEGs) in SrTiO₃ have become model systems for engineering emergent behaviour in complex transition metal oxides. Understanding the collective interactions that enable this, however, has thus far proved elusive. Here we demonstrate that angle-resolved photoemission can directly image the quasiparticle dynamics of the *d*-electron subband ladder of this complex-oxide 2DEG. Combined with realistic tight-binding supercell calculations, we uncover how quantum confinement and inversion symmetry breaking collectively tune the delicate interplay of charge, spin, orbital and lattice degrees of freedom in this system. We reveal how they lead to pronounced orbital ordering, mediate an orbitally enhanced Rashba splitting with complex subband-dependent spin-orbital textures and markedly change the character of electron-phonon coupling, co-operatively shaping the low-energy electronic structure of the 2DEG. Our results allow for a unified understanding of spectroscopic and transport measurements across different classes of SrTiO₃-based 2DEGs, and yield new microscopic insights on their functional properties.

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The ubiquitous perovskite oxide SrTiO₃, a wide-gap band insulator, hosts varied bulk properties including quantum paraelectricity, dilute doping-induced superconductivity and high thermoelectric coefficients. These reflect a subtle competition between interactions of the underlying quantum many-body system. Intriguingly, thermodynamic and transport measurements^{1–6} indicate that the balance of these interactions can be tuned to engineer striking emergent properties when quantum confinement and doping are combined to create a two-dimensional electron gas (2DEG)^{1,2,7}. A diverse and attractive array of properties have been uncovered to date in this system, including gate-tuned superconductivity^{3,4,8}, its coexistence with ferromagnetism^{5,6} and enhanced Seebeck coefficients⁹, establishing SrTiO₃-based 2DEGs as a model platform for use in future multifunctional electronic devices¹.

They are most commonly realised at a polar interface to another band insulator LaAlO₃, creating a narrow conducting channel that resides solely within the SrTiO₃ (ref. 10). Similar 2DEGs can also be created by interfacing SrTiO₃ to a wide array of other band or Mott insulators including NdAlO₃ (ref. 11), LaTiO₃ (ref. 12) and GdTlO₃ (ref. 13), by chemical doping of electrons into narrow SrTiO₃ channels^{14,15}, analogous to δ -doping of semiconductors such as Si, and by field-effect doping in a transistor-style configuration⁸. Moreover, the recent discovery of a 2DEG formed at the free surface of a bulk SrTiO₃ crystal opens new avenues for its advanced spectroscopic investigation^{7,16}.

Exploiting this, here we present unified angle-resolved photoemission (ARPES) measurements and tight-binding supercell calculations revealing new richness of the electronic structure of this model oxide 2DEG. We show how a pronounced orbital ordering mediates an unconventional spin splitting, giving rise to strongly anisotropic and subband-dependent canted spin-orbital textures. The orbitally enhanced Rashba effect explains the pronounced spin splittings previously inferred from magneto-transport in this system, while simultaneously revealing a breakdown of the conventional picture used to describe these. We uncover how this complex ladder of subband states are further renormalized by many-body interactions. This reconciles

previous discrepancies between effective masses estimated from ARPES and quantum oscillations, unifying the properties of surface and interface SrTiO₃ 2DEGs, and reveals a strikingly different nature of electron–phonon coupling compared with bulk SrTiO₃.

Results

Orbital ordering. Figure 1 summarizes the generic electronic structure of SrTiO₃ 2DEGs, as revealed by ARPES from a SrTiO₃(100) surface with saturated band bending⁷. Consistent with previous reports from both surface and interface 2DEGs^{7,16–19}, we find a broad bandwidth that extends up to ≈ 250 meV below the Fermi level. Here we can resolve a ladder of at least three light subband states that contribute concentric circular Fermi surface sheets, co-existing with just a single heavy electron band ($m^* = 14 \pm 3m_e$) that has a much shallower binding energy of < 50 meV and gives rise to elliptical Fermi surfaces oriented along $\langle 10 \rangle$. From this Fermi surface topology together with the polarization dependence of our measured intensities (Supplementary Fig. 2), we assign not only the lowest¹⁶ but rather the whole ladder of observed light states as having dominantly d_{xy} orbital character, while the heavy states derive from $d_{xz/yz}$ orbitals. This immediately indicates a strong breaking of the t_{2g} orbital degeneracy that is present in the bulk electronic structure of SrTiO₃ (ref. 20), driving a pronounced orbital ordering with a polarization $P = \frac{n(d_{xy}) - n(d_{xz/yz})}{n(d_{xy}) + n(d_{xz/yz})}$, which exceeds 30%, a lower limit derived from our experimentally resolved Fermi surface areas.

This is a direct consequence of the real-space anisotropy of the orbital wavefunctions combined with inversion symmetry breaking by the electrostatic potential that defines the 2DEG by creating a steep asymmetric quantum well along the z direction (Fig. 1c). As shown by our self-consistent tight-binding supercell calculations (Fig. 1b, see Methods), the resulting quantized subbands that derive from planar d_{xy} orbitals have wavefunctions reminiscent of the envelope functions of a semiconductor quantum well, except that in SrTiO₃ they are much more localized in real space, almost to within a single unit cell for the lowest subband state. In contrast, the potential variation acts as a

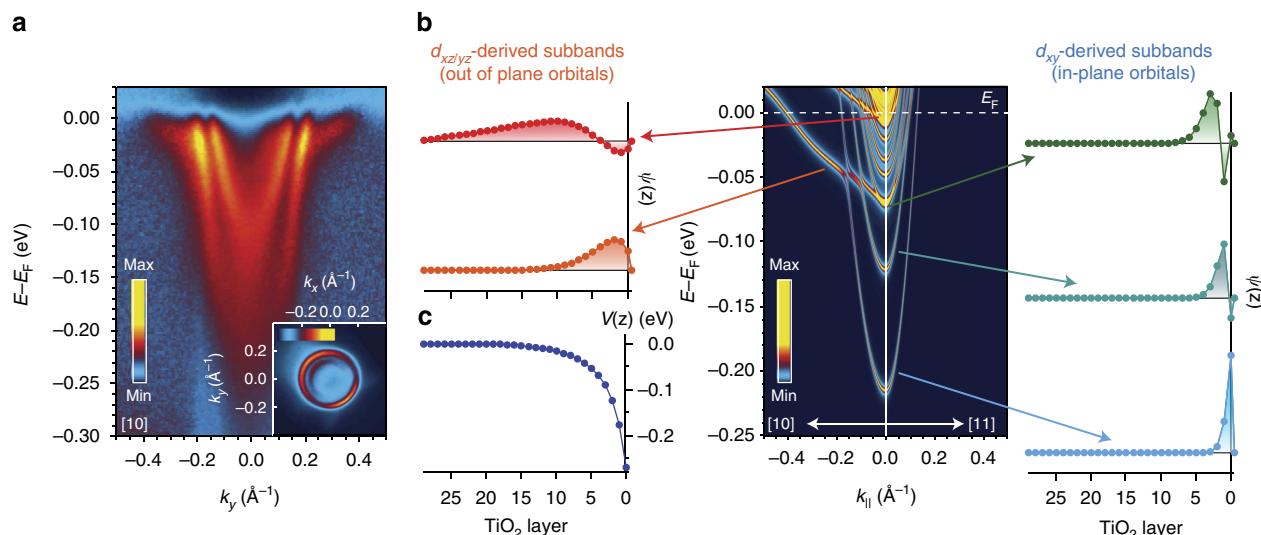


Figure 1 | Orbital ordering of a 2D electron gas in SrTiO₃. (a) E versus k dispersion from ARPES ($h\nu = 50$ eV, measured along the $[10]$ direction), revealing a multi-orbital subband structure comprising co-existing ladders of light and massive d -electron subband states. The respective circular and faint elliptical Fermi surface pockets (measured with $h\nu = 51$ eV and the polarization along $[11]$) are shown in the inset. For the dispersion plot, a normalization (division by the average MDC) has been applied to better reveal the massive band, as shown in Supplementary Fig. 1. (b) This electronic structure is well described by a self-consistent tight-binding supercell calculation. The spatial dependence of the subband wavefunctions along the confinement direction, $\Psi(z)$, reveal a pronounced real-space orbital ordering, a direct consequence of near-surface band bending (c).

much weaker perturbation on the out-of-plane $d_{xz/yz}$ orbitals, which have much larger hopping amplitudes along the z direction. The resulting subbands sit close to the top of the potential well, leading to wavefunctions that penetrate much deeper into the bulk. This disparate spatial extent of the subband states is consistent with their relative spectral weight in our surface-sensitive ARPES measurements.

Unconventional Rashba spin splitting. The same breaking of inversion symmetry that drives this orbital ordering can additionally lift the spin degeneracy through a Bychkov–Rashba-type spin–orbit interaction^{21–23}. Focussing near the band bottom of the lowest d_{xy} band (Fig. 2a,b), we indeed find a small characteristic splitting between the calculated energy of spin-up and spin-down states, $\delta E = 2\alpha k_{\parallel}$, with a Rashba parameter, $\alpha = 0.003 \text{ eV\AA}$. The non-negligible splitting found here, despite the modest spin–orbit interaction in $3d$ transition metals, is indicative of the very strong electric field gradient of the confining potential. From our self-consistent band-bending potential calculation (Fig. 1c), we estimate that this exceeds $2 \times 10^8 \text{ V m}^{-1}$ within the first two unit cells where the lowest subband state is confined. For the more delocalized second d_{xy} subband, whose wavefunction extends into regions of shallower band bending, we find a slightly smaller Rashba parameter $\alpha = 0.0014 \text{ eV\AA}$, confirming that the strength of this spin splitting is controlled by the confining electric field. This should therefore

be directly tuneable by electrical gating, suggesting a potential route towards spintronic control in oxides.

Unlike typical Rashba systems such as the Au(111) surface, however, here the interplay between orbital ordering and spin–orbit coupling leads to a significantly richer spin structure of the 2DEG states. Close to the crossings of the light d_{xy} and heavy $d_{xz/yz}$ subbands, the spin splitting increases by approximately an order of magnitude, concomitant with a strong mixing of their orbital character (Fig. 2a–c). This rationalizes an increased spin splitting reported from transport when the $d_{xz/yz}$ subbands become populated in electrically gated $\text{SrTiO}_3/\text{LaAlO}_3$ interface 2DEGs²⁴. Moreover, the crossover from k -linear to strongly enhanced spin splitting that we find here readily explains the approximately k^3 dependence of the splitting that has been reported²⁵. Our layer-projected calculations indicate that the subband wavefunctions become more delocalized in the z direction close to these band crossings, a natural consequence of the stronger overlap of neighbouring $d_{xz/yz}$ orbitals along z (Supplementary Fig. 3). This delocalization would naively be expected to reduce the strength of the Rashba effect. In contrast, its significant enhancement here points to a dominant role of inter-orbital hopping in driving such surprisingly large spin splittings. Similar enhancements have recently also been observed in other calculations, mainly based on model 3-band Hamiltonians^{26–28}, which are qualitatively entirely consistent with our results. Our calculations demonstrate how this is a direct consequence of orbital ordering in the

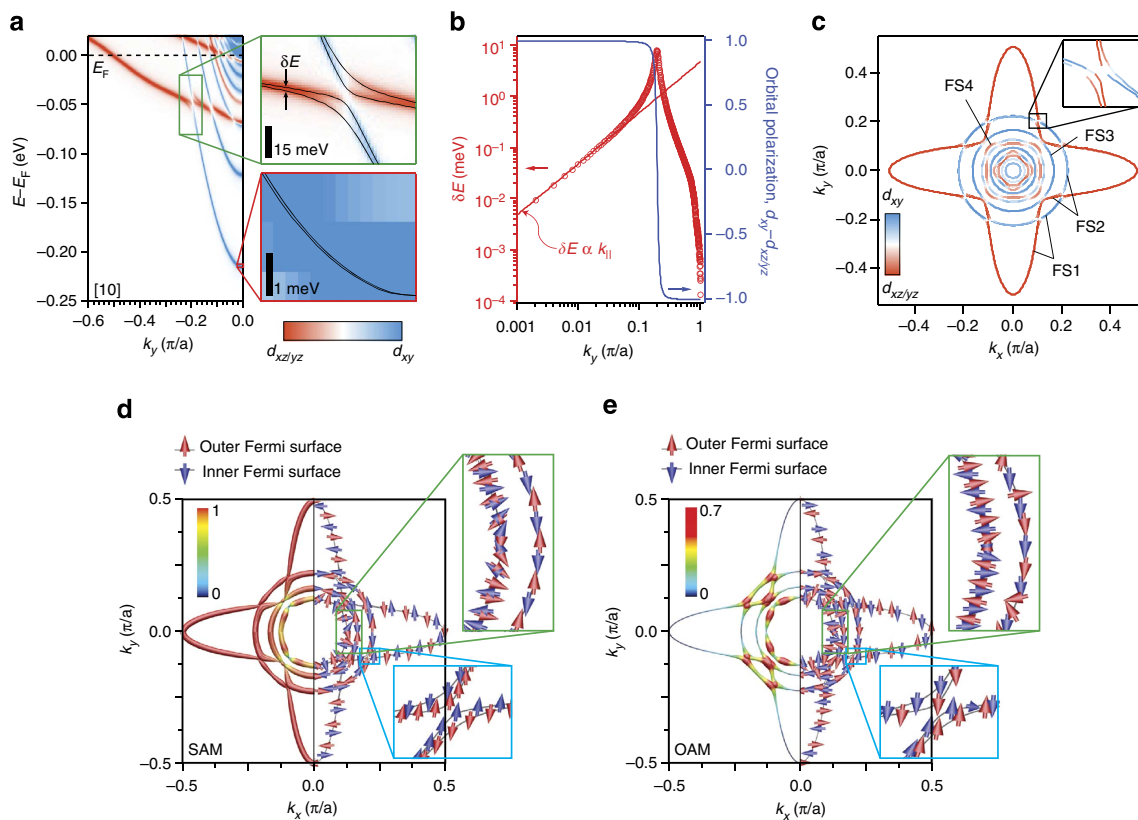


Figure 2 | Orbitaly enhanced spin splitting. (a) Orbitaly resolved electronic structure of the 2DEG along the [10] direction. Magnified views reveal a weak Rashba-type spin splitting around the band bottom, which becomes enhanced by approximately an order of magnitude at the crossings of the light and heavy subband states where the orbital character becomes strongly mixed, as quantified in (b) for the lowest subbands. (c) The calculated Fermi surface shows how similar orbital mixing and pronounced spin splittings occur at the crossings of circular d_{xy} - and elliptical $d_{xz/yz}$ -derived Fermi surface sheets. This gives rise to an exotic anisotropic and subband-specific coupled spin–orbital texture of the 2DEG, as evident from the magnitude (left) and direction (right) of the (d) spin, SAM and (e) orbital, OAM, angular momenta of the four largest Fermi surface sheets. The magnitude of the SAM (OAM) is represented on a false colour scale in units of $\hbar/2$ (\hbar), respectively, and is shown for additional Fermi surfaces in Supplementary Fig. 4.

real experimentally confirmed multi-subband structure of the SrTiO₃ 2DEG.

Moreover, as shown in Fig. 2d,e, they reveal an exotic coupled spin–orbital texture of the resulting 2DEG Fermi surfaces. While an approximately perpendicular spin-momentum locking ensures tangential spin winding around the circular d_{xy} sections of Fermi surface, it leads to spins aligned almost perpendicular to the Fermi surface for large sections of the extremely anisotropic d_{xz} (d_{yz}) sheets. Around the crossings of d_{xy} and $d_{xz/yz}$ states, the spins of neighbouring Fermi surfaces align (anti-)parallel with a $|\downarrow\rangle|\uparrow\rangle|\uparrow\rangle|\downarrow\rangle$ ordering, ensuring maximal hybridization gaps are opened. At the same time, rather than being quenched to zero as might have naively been expected, we find a finite orbital angular momentum (OAM, L) emerges. This is relatively small ($\lesssim 0.05\hbar$) for the isolated d_{xy} and $d_{xz/yz}$ sections of Fermi surface (Fig. 2e and Supplementary Fig. 3), but grows as large as $0.7\hbar$ around the band crossings. Moreover, we find that the OAM is oriented either parallel or antiparallel to the corresponding spin angular momentum (SAM, S), and so this increase in OAM maximally enhances $L \cdot S$, by a factor of ~ 14 , at and in the vicinity of the hybridization gaps, comparable to the corresponding increase in spin splitting (Fig. 2b).

This therefore provides a natural basis to understand the large spin splittings inferred from magnetotransport^{29–31}, despite the small atomic spin–orbit interaction of SrTiO₃, in terms of an orbital Rashba effect^{32,33}. For isolated d_{xy} sections of Fermi surface, we find that the OAM of the inner and outer spin-split branches have the same helicity, consistent with a weak spin–orbit coupling limit³³. For the $d_{xz/yz}$ sections of Fermi surface, however, the inner and outer branches have opposite OAM, reflecting additional richness as compared with model systems such as noble metal surface states³³. This causes mixed helicities of the OAM around the inner branch of each Fermi surface sheet, as compared with a complete 2π winding for the outer branches (see arrows in Supplementary Fig. 4). Importantly, we find that the dominant inter-band interactions cause the winding direction of both the OAM and SAM of the outer d_{xy} -derived Fermi

surfaces to abruptly switch sign across the hybridization gaps. For several inner Fermi surfaces whose orbital character is strongly mixed, continuously evolving between d_{xy} - and $d_{xz/yz}$ -like around the Fermi surface (Fig. 2c), this leads to strongly frustrated spin and orbital textures, rapidly canting from tangential to radial alignment as a function of Fermi surface angle (Fig. 2d,e). This is quite distinct from the functional form of conventional Rashba splitting²¹ and provides strong constraints for the influence of spin splitting on magnetism³⁴ and superconductivity³⁵.

Many-body interactions. In Fig. 3, we further uncover a pronounced role of electron–phonon interactions on this complex hierarchy of electronic states. Unlike in bulk-doped SrTiO₃, where the Fermi energy is typically only a few meV and the electron–phonon interaction is thus non-retarded, the occupied widths of different subbands of the 2DEG range from almost zero up to values greater than the highest phonon frequency of ≈ 100 meV. This is an unusual situation, neither described by the adiabatic ($\hbar\omega_D \ll E_F$) nor the anti-adiabatic ($\hbar\omega_D \gg E_F$) approximation, and points to a complex influence of electron–phonon coupling in this system. We extract the corresponding self-energy, $\Sigma_{e-ph}(\omega) = \Sigma'(\omega) + i\Sigma''(\omega)$, from our ARPES measurements of the lowest subband along the [11] direction (see methods), where we resolve an isolated band all the way up to E_F . The slope of Σ' at the Fermi level yields an electron–phonon coupling strength of $\lambda = 0.7(1)$, while its broad maximum between ≈ 20 – 60 meV is indicative of coupling to multiple modes. Indeed, the experimentally determined self-energy is in excellent agreement with a calculation within Eliashberg theory that assumes a coupling function $\alpha^2F(\omega)$ proportional to the entire phonon density of states associated with the motion of oxygen and Ti ions³⁶ and includes the realistic 2DEG electron density of states from our tight-binding calculation.

Together with a moderate correlation-induced mass enhancement of ≈ 1.4 that we estimate from a Kramers–Kronig transform of a Fermi-liquid contribution to the imaginary part

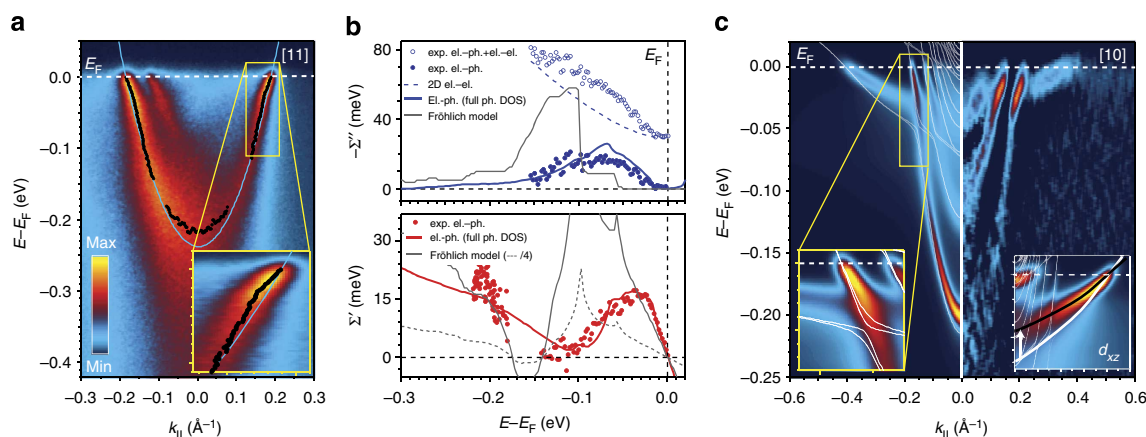


Figure 3 | Quasiparticle dynamics of the subband states. (a) ARPES measurements along the [11] direction ($h\nu = 51$ eV), together with the peak positions of fits to MDCs and EDCs (black dots) and a cosine ‘bare band’ dispersion (blue line). The data reveal pronounced signatures of electron–phonon coupling. Particularly apparent is a low-energy kink in the dispersion at $\omega \approx 30$ meV, shown magnified inset. (b) Real, Σ' , and imaginary, Σ'' , parts of the extracted self-energy. The open blue symbols show Σ'' before subtraction of an electron–electron scattering contribution to the total measured imaginary part of the self-energy, approximated here by the expression $\Sigma''_{e-e}(\omega) = \beta\omega^2(1 + 0.53\ln(\omega/E_F))$ for a 2D Fermi liquid (dashed line, $\beta = 1.5 \text{ eV}^{-1}$, $E_F = 0.25 \text{ eV}$). The solid lines are calculated electron–phonon self energies with the Eliashberg function proportional to the full phonon density of states (red/blue) or for coupling to three LO phonons (black), which dominate the interaction in lightly doped bulk SrTiO₃. (c, left) Calculated electron–phonon spectral function, $A(k, \omega)$, along the [10] direction (see Methods). This reveals complete bandwidth renormalization of the heavy states (particularly apparent when $A(k, \omega)$ is projected only onto d_{xz} orbitals as shown in the right inset), and kinks in the light d_{xy} subband dispersions at their crossings with this renormalized heavy state (left inset), both in good agreement with our experimental measurements, shown in the right half using curvature analysis⁴⁶ of the raw data.

of the self-energy, our analysis suggests an overall mass enhancement arising from many-body interactions of $m^*/m_{\text{band}} \approx 2.1$, close to the values deduced for lightly doped bulk SrTiO₃ from measurements of the electronic specific heat³⁷ and optical spectroscopy³⁸. The nature of the electron–phonon coupling, however, is very different. In lightly doped bulk SrTiO₃, it is dominated by long-range coupling to longitudinal optical (LO) phonons as described by the Fröhlich model^{38,39}. This model predicts much weaker coupling to low-energy modes than observed here, but a significantly stronger coupling to the highest LO phonon at 100 meV, as evident from a calculation employing coupling strengths from bulk SrTiO₃ (refs 39,40), which yield an electron–phonon self-energy in clear contrast to our experimental findings (Fig. 3b).

The electric field that confines the 2DEG is known to dramatically reduce its dielectric constant^{7,41}. This, together with higher carrier densities as compared with bulk SrTiO₃, will lead to shorter electronic screening lengths for the 2DEG, explaining the observed suppression of long-range coupling to LO modes. The enhanced coupling to low-energy phonons for the 2DEG instead leads to a pronounced kink in the dispersion of the d_{xy} subbands at an energy around 30 meV. We resolve these along both the [10] and [11] directions for the first two d_{xy} subbands. Crucially, the resulting enhanced quasiparticle mass, which we estimate as 1.1(2) m_e from our measured Fermi velocities, rectifies the discrepancy between the light masses around 0.6 m_e reported in earlier ARPES studies of SrTiO₃ surface 2DEGs^{7,16} and recent quantum oscillation experiments that revealed effective masses typically around 1 m_e (refs 42,43).

Intriguingly, along [10] the kink energy coincides almost exactly with the crossing of the light d_{xy} and heavy d_{xz} subband states. This behaviour is well captured by our spectral function simulations calculated from our tight-binding bare dispersions and electron–phonon self-energy. These illustrate a very different effect of electron–phonon interactions on the heavy compared with the light subbands of the 2DEG, the former coupling to phonons with frequencies ranging from below to above the bare bandwidth. Our calculations reveal that electron–phonon coupling essentially results in an overall bandwidth renormalization of these states, in agreement with our experimental data where we find the band bottom of the heavy state substantially above the value predicted by our model tight-binding calculations.

Discussion

The combination of electron–phonon coupling with orbital ordering therefore effectively pins the crossings of the d_{xy} and $d_{xz/yz}$ subbands to the low-energy peak in the phonon density of states. As demonstrated above, however, we additionally find orbitally enhanced spin–orbit splittings that become maximal around these band crossings. Our direct spectroscopic measurements, together with our theoretical calculations, therefore demonstrate how a co-operative effect of orbital ordering and electron–phonon coupling sets the relevant energy scale for dominant spin splitting in this system.

Together, this reveals an intricate hierarchy of interactions and orderings governing the low-energy electronic structure of the SrTiO₃ 2DEG (Fig. 4). Electrostatic screening in response to a surface or interface charge generates an electron accumulation layer confined to a narrow potential well. The resulting quantum size effects drive pronounced orbital ordering, creating multiple intersections of light d_{xy} - and heavy $d_{xz/yz}$ -derived subband states. This leads to orbital mixing and induces a significant local orbital angular momentum, which in turn permits a pronounced spin splitting to emerge, despite modest atomic spin–orbit coupling. Many-body interactions, of strikingly different form to the bulk,

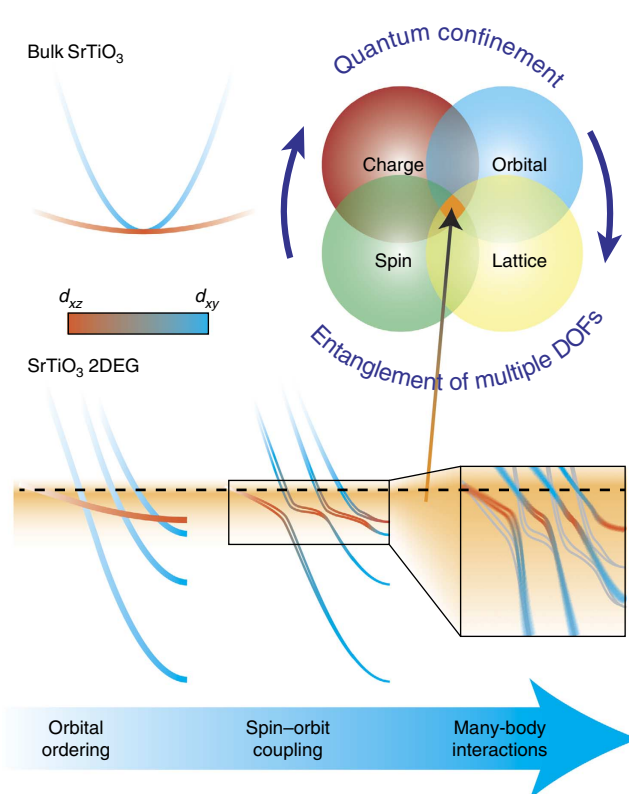


Figure 4 | Hierarchy and interplay of underlying degrees of freedom.

In the formation of the 2DEG, quantum confinement reconstructs the bulk electronic structure into a rich array of intersecting orbitally ordered subband states. The interplay of this with spin–orbit coupling lifts the spin degeneracy of these bands, particularly strong around their crossings, through an orbitally enhanced Rashba-like interaction. Electron–phonon and electron–electron interactions further renormalise these spin-split dispersions, increasing the quasiparticle masses close to the Fermi level and causing a complete bandwidth narrowing of the shallow $d_{xz/yz}$ states that pins the orbitally ordered band crossings close to the energy of the lowest phonon mode. This, in turn, sets the energy scale for enhanced spin–orbit splittings around these crossings, strongly entangling the charge, spin, orbital and lattice degrees of freedom of the 2DEG on a low-energy scale (orange shading).

enhance the quasiparticle masses of these spin-split subbands, reduce their bandwidths and renormalize the energetic locations of their intersections, thus modulating their unconventional spin splitting.

Together, this interplay strongly entangles the charge, spin, orbital and lattice degrees of freedom of the SrTiO₃ 2DEG on an energy scale ≤ 30 meV. This will dominate both its transport and thermodynamic properties. Already we have shown how this explains enhanced quasiparticle masses observed from quantum oscillations as well as signatures of spin splitting in magnetotransport, unifying electrical and spectroscopic measurements from surface- and interface-based SrTiO₃ 2DEGs. More generally, it establishes how quantum size effects can dramatically manipulate the underlying electronic landscape of interacting electron liquids, setting the stage for engineering new emergent properties by dimensional confinement in transition metal oxides.

Methods

Angle-resolved photoemission. ARPES measurements were performed at the CASIOPEE beamline of SOLEIL synchrotron, the SIS beamline of the Swiss Light Source, and beamline 10.0.1 of the Advanced light source using Scienta R4000

hemispherical electron analysers, and with base pressures $<5 \times 10^{-11}$ mbar. Single crystal SrTiO₃ commercial wafers were cleaved *in situ* at the measurement temperature of $T = 20\text{--}30$ K along notches defining a (100) plane. Measurements were performed on stoichiometric transparent insulating samples as well as very lightly La-doped samples (Sr_{1-x}La_xTiO₃ with $x = 0.001$) to help avoid charging. 2DEGs were induced at the bare surface by exposure to intense UV synchrotron light⁷. The samples were exposed to irradiation doses $\gtrsim 1,000 \text{ J cm}^{-2}$ to saturate the formation of the 2DEG, and we experimentally confirmed that saturation was reached before starting any of the measurements presented here. The data shown here was measured using *s*-polarized photons of 51 or 55 eV, except for the Fermi surface maps shown in Supplementary Fig. 2 that used 43-eV *s*- and *p*-polarized light. All measurements were performed in the second Brillouin zone.

Self-energy determination. To determine the electron–phonon self-energy experimentally, we fit momentum distribution curves (MDCs) and energy distribution curves (EDCs) of the lowest d_{xy} subband measured along the [11] direction. We chose this band as its dispersion does not intersect that of the heavy $d_{xz/yz}$ subbands up to the chemical potential along this direction (Fig. 1b), allowing us to perform a quantitative analysis over an extended energy range, free from complications associated with the hybridization of different subbands. The real part of the self-energy, $\Sigma'(\omega)$, is given by the difference between our extracted dispersion and that of a ‘bare’ band. In order to derive a Kramers–Kronig consistent self-energy, we take the cosine bare band shown in Fig. 3a, which includes a moderate bandwidth renormalization due to electron correlations. We extract the imaginary part of the self-energy, $\Sigma''(\omega) = \Delta k(\omega)/2 \cdot \partial \epsilon / \partial k$, where Δk is the full width at half maximum of Lorentzian fits to MDCs, and $\partial \epsilon / \partial k$ is the bare band dispersion. This results in an imaginary part that includes a contribution from electron–electron interactions, which we approximate by the expression for a two-dimensional Fermi liquid $\Sigma''_{e-e}(\omega) = \beta \omega^2 (1 + 0.53 \ln(\omega/E_F))$. Subtracting this contribution with realistic parameter values of $\beta = 1.5 \text{ eV}^{-1}$ and $E_F = 0.25 \text{ eV}$ from our total extracted Σ'' (see Fig. 3b) yields the imaginary part of the electron–phonon self-energy.

Electronic structure and self-energy calculations. To calculate the subband structure, we start from a relativistic density functional theory calculation of bulk SrTiO₃ using the modified Becke–Johnson exchange potential and Perdew–Burke–Ernzerhof correlation functional as implemented in the WIEN2K programme⁴⁴. The muffin-tin radius of each atom R_{MT} was chosen such that its product with the maximum modulus of reciprocal vectors K_{max} become $R_{MT}K_{max} = 7.0$. The Brillouin zone was sampled by a $15 \times 15 \times 15$ *k*-mesh. We downfold this using maximally localized Wannier functions to generate a set of bulk tight-binding transfer integrals, and then incorporate these into a 30-unit cell supercell with additional on-site potential terms to account for band bending via an electrostatic potential variation. We solve this self-consistently with Poisson’s equation, incorporating an electric field-dependent dielectric constant⁴¹, to yield the bare band dispersions including Rashba-type spin splitting⁴⁵ of the 2DEG. We stress that the total magnitude of the band bending is the only adjustable parameter, and yields a realistic electronic structure in good agreement with our spectroscopic measurements apart from our observed signatures of electron–phonon interactions that are not included at the level of density functional theory. To incorporate these, we calculate the self-energy $\Sigma_{e-ph}(\omega)$ in an Eliashberg model,

$$\Sigma_{e-ph}(\omega) = \int_{-E_F}^{\infty} N(\epsilon) d\epsilon \int_0^{\tilde{\omega}_{max}} d\tilde{\omega} \alpha^2 F(\tilde{\omega}) \times \left\{ \frac{f(-\epsilon, T) + n(\tilde{\omega}, T)}{\omega - \epsilon - \tilde{\omega} + i\delta^{\pm}} + \frac{f(\epsilon, T) + n(\tilde{\omega}, T)}{\omega - \epsilon + \tilde{\omega} + i\delta^{\pm}} \right\}$$

where $N(\epsilon)$ is the bare electronic density of states determined from our tight-binding calculations and $f(\epsilon, T)$ and $n(\tilde{\omega}, T)$ are the Fermi and Bose occupation factors. For the coupling function $\alpha^2 F(\tilde{\omega})$ we use two different models. The blue line in Fig. 3b assumes $\alpha^2 F(\tilde{\omega})$ proportional to the entire O- and Ti-derived phonon density of states from ref. 36, while the black line is a calculation for the coupling strengths given in refs 39,40 for the three LO phonons that were found to dominate the electron–phonon interaction in lightly doped bulk samples. We then calculated the spectral function

$$A(k, \omega) = -\frac{1}{\pi} \frac{\Sigma''(\omega)}{[\omega - \epsilon(k) - \Sigma'(\omega)]^2 + [\Sigma''(\omega)]^2}$$

where the bare band dispersion $\epsilon(k)$ is taken from our tight-binding calculation. To better compare with our experimental data in Fig. 3c, we project this onto different atomic orbitals, and include contributions from d_{xy} and d_{xz} but not d_{yz} orbitals to account for transition matrix elements in our experimental geometry. We additionally project the calculation onto different layers of our supercell, and incorporate an exponential attenuation of signal with depth below the surface in photoemission, assuming an inelastic mean free path of 5 Å, into our simulation. Finally, we convolve the simulated spectral function with a 2D Gaussian to account for an experimental energy and momentum resolution of 0.01 eV and 0.015 Å⁻¹, respectively.

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Author contributions

The experimental data was measured by P.D.C.K., S.M.W., A.T., A.d.I.T., T.E., P.B., W.M. and F.B., and analysed by P.D.C.K., S.M.W., A.T. and F.B. P.D.C.K. and M.S.B. performed the electronic structure calculations and A.T. performed the electron-phonon self-energy calculations. S.-K.M. maintained the ARPES endstation at the Advanced Light Source and provided experimental support. P.D.C.K. and F.B. were responsible for overall project planning and direction, and wrote the manuscript with input and discussion from all co-authors.

Additional information

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