A Systematic Investigation of the Coupling between One-Dimensional Edge States of a Topological Crystalline Insulator

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The interaction of spin-polarized one-dimensional (1D) topological edge modes localized along single-atomic steps of the topological crystalline insulator Pb0.7Sn0.3Se(001) has been studied systematically by scanning tunneling spectroscopy. Our results reveal that the coupling of adjacent edge modes sets in at a step-to-step distance \(d_{ss} \leq 25\) nm, resulting in a characteristic splitting of a single peak at the Dirac point in tunneling spectra. Whereas the energy splitting exponentially increases with decreasing \(d_{ss}\) for single-atomic steps running almost parallel, we find no splitting for single-atomic step edges under an angle of 90°. The results are discussed in terms of overlapping wave functions with \(p_x, p_y\) orbital character.

Introduction — The discovery of two-dimensional (2D) topological insulators (TIs), such as HgTe/CdTe [1] or InAs/GaSb quantum wells [2], generated significant attention, especially due to the existence of edge states protected by time-reversal symmetry. The spin–momentum locking pertinent to these states mandates that charge carriers with a given spin can only propagate in a predetermined direction, thereby inhibiting disorder scattering and greatly reducing dissipation in electrical transport [3–7]. Further material classes with topologically protected band structures have been predicted and realized, such as topological crystalline insulators (TCI) [8,11] and Weyl semimetals [12,17]. However, although the ruggedness of the respective topological states is often praised as one of their main advantages, the reliable realization of one-dimensional (1D) topological conductance channels highly relevant for practical applications remains a challenge. Some systems with topological 1D states, like higher-order topological hinge states [18,19], or surface step edges of weak TIs [20] or TCIs [21], could be identified, but systematic studies which address the interaction of adjacent channels remain elusive. As a result the question up to which density topological 1D channels may be packed without compromising the signal quality has not yet been answered.

In this context hybridization effects of topological states play a particularly important role. For example, the close proximity to the underlying substrate may lead to a significant interaction of topological edge states with substrate bulk states [22,24]. Calculations predict that—if tuned appropriately—these effects may even be utilized, such as in topological-insulator–ferromagnetic-metal (TI–FM) heterostructures, where hybridization-induced interface states lead to a large spin-transfer torque [24]. Strong hybridization between the top and the bottom surfaces, however, can also result in band crossings resulting in a trivial TCI phase [24] or—for very thin films of three-dimensional TIs—to the complete quenching of the topological surface state [24].

In this contribution we systematically study the interactions of one-dimensional topological states which arise at odd-atomic step edges of (001)-terminated (Pb,Sn)Se single crystals. Whereas the rock salt compound PbSe is a trivial insulator, the order of \(p\)-derived cation and anion surface states bands is inverted for Pb1−xSnxSe at \(x \geq 0.2\), i.e., when substituting a sufficient fraction of Pb atoms with Sn [19,25]. Band inversion results in four Dirac cones per surface Brillouin zone (SBZ) centered in close proximity to the \(\Gamma\) and \(\Delta\) points which are protected by (110) mirror symmetry [29,32].

Clean (001) surfaces of Pb1−xSnxSe are typically created by cleaving bulk crystals. As first shown in Ref. [33] tunneling spectra of flat (Pb,Sn)Se (001) surfaces exhibit a V-shaped dip with two symmetrically arranged side peaks which were assigned to the Dirac point and two Van Hove singularities caused by saddle points in the dispersion, respectively. However, the cleaving process unavoidably also results in a variety of steps of different height and shape. Whereas the electronic structure of double- or other even-atomic step edges is virtually identical to a flat terrace, see Ref. [31] (Note 1), the observation of a 1D topological edge mode at odd-atomic step edges, e.g., steps with a height of one or three atomic layers, which are characterized by a strong peak in the LDOS at the Dirac energy, was reported in Ref. [24] and confirmed in Ref. [35]. As detailed in Ref. [36], adjacent extended (001) terraces of (Pb,Sn)Se must—due to the displacement by one atomic layer in a two-atomic unit cell—be described by Hamiltonians with chiral symmetry and an inverted order of states. The resulting surface states are divided into two classes with an even or odd number of layers. Therefore, odd-atomic step edges host a doubly-degenerate flat band at the Dirac energy which is protected by the mirror plane symmetry of the film [36].
In this paper we systematically investigate the interaction between closely adjacent 1D edge states of the (001) surface of the TCI (Pb,Sn)Se. Hybridization is observed for step-to-step distances \( d_{ss} \lesssim 25 \text{ nm} \), resulting in the splitting of the single peak in tunneling spectra at the Dirac point into a double-peak. Our results reveal that the energetic splitting exponentially increases with decreasing \( d_{ss} \). It is largely insensitive to the details of a particular configuration of step edges as long as the steps run approximately in parallel. In contrast, for single-atomic steps crossing at an almost perpendicular angle, we do not observe any splitting.

**Methods** — Experiments have been performed in an ultra-high vacuum system equipped with a cryogenic STM (operation temperature \( T = 4.9 \text{ K} \)). Pb\(_{0.7}\)Sn\(_{0.3}\)Se p-doped single crystals grown by the self-selecting vapor growth method [9, 21] (see Ref. 34, Note 2, for further information) have been cleaved in UHV at a base pressure \( p < 10^{-10} \text{ mbar} \) and immediately transferred into the STM. Tunneling conductance \( \text{d}I/\text{d}U \) maps and spectra were measured by lock-in technique with modulation voltage \( U_{\text{mod}} = 2.5 \text{ mV} \) at a frequency \( f = 790 \text{ Hz} \).

**Results** — Figure 1(a) shows a typical STM topography image of the cleaved (001) surface of the TCI Pb\(_{0.7}\)Sn\(_{0.3}\)Se. Numerous wedge-shaped atomically flat plateaus can be recognized. The line profile measured along the green line in Fig.1(a), presented in Fig.1(b), reveals that most step edges are about 3 Å high, corresponding to half of the two-atomic unit cell. Fig.1(c) shows typical \( \text{d}I/\text{d}U \) spectra measured on a flat terrace far away from any step edge (1), on a double-atomic step edge (2), and at two single-atomic step edges [points (3),(4)]. Spectrum (1) exhibits a minimum at the Dirac point \( E_D = (125 \pm 5) \text{ meV} \) and two shoulders (\( L^- \) and \( L^+ \)), which are associated with the two saddle points in the dispersion [30, 33]. Spectrum (2) taken on a double-step edge is virtually indistinguishable from (1), indicating the absence of any edge state at step edges equivalent to an even number of atomic layers. In agreement with our earlier publication [21], a pronounced peak at the Dirac energy indicative for the 1D edge state can only be found at odd-atomic step edges, such as spectrum (3) measured on a single-atomic step edge. However, we find that the spectra of single-atomic step edges at some locations deviate from this simple single-peak structure. For example, the spectrum measured at (4) close to the termination point of an acute-angled wedge-shaped plateau reveals a pronounced double-peak feature in the vicinity of the Dirac point with a peak splitting of about 25 mV.

To understand the physical origin of this peak splitting we performed detailed spectroscopic investigations at numerous single-atomic step edge structures on a variety of topological Pb\(_{0.7}\)Sn\(_{0.3}\)Se surfaces. Our results indicate that the peak splitting observed in Fig.1(c) on the single-atomic step edge is not caused by the presence of kinks, defects, or random variations of the local doping level [32]. Instead, the data reveal that the splitting occurs when two single-atomic step edges are in close proximity. A particularly illustrative case is displayed in Fig.2. The topographic STM image of Fig.2(a) shows a wedge-shaped plateau bound by two single-atomic step edges which converge under an acute angle of 3.4° ± 0.3°. As a result, the step-to-step distance \( d_{ss} \) slowly decreases over a total length of about 500 nm from \( d_{ss} > 30 \text{ nm} \) in the bottom part to almost zero at the apex of the wedge. Both steps have rough edges which clearly deviate from high symmetry directions. Nevertheless, the \( \text{d}I/\text{d}U \) map measured at a bias voltage corresponding to the Dirac point presented in Fig.2(b) shows several unique features. First, in the bottom part of Fig.2(b) where the two single-atomic step edges are far apart an intense \( \text{d}I/\text{d}U \) signal can be recognized along the step edges, indicating the presence of the topological edge state. Second, as \( d_{ss} \) falls below about 15 nm, indicated by two arrows in Fig.2(b), the intensity of the \( \text{d}I/\text{d}U \) signal noticeably decreases. Third, close to the top of the wedge where \( d_{ss} \lesssim 7 \text{ nm} \) (marked by a hatched ellipse) it becomes essentially indistinguishable from the signal measured on the surrounding terraces.

This modification of the LDOS at the Dirac energy correlates with a systematic alteration of the local STS data. Some exemplary spectra measured at various step-to-step distances ranging from \( d_{ss} = 30 \text{ nm} \) down to \( d_{ss} = 7 \text{ nm} \) show...
FIG. 2. (a) Topographic STM image of a wedge-shaped plateau surrounded by single-atomic step edges. Note, that the x- and y-direction are scaled differently, i.e., the wedge shape is even more acute-angled than it appears. (b) $dI/dU$ map measured at the Dirac energy of the same region shown in (a). An enhanced intensity at the position of the step edges is clearly visible in the bottom part, indicating the presence of the topologically protected edge state. As the wedge-shaped plateau becomes more narrow, the intensity of the $dI/dU$ signal measured at both step edges decreases (see region above arrows) and eventually vanishes close to the termination point (ellipse). (c) Tunneling spectra measured at the left (blue) and right (red) single-atomic step edge at various step-to-step distances indicated in (a). (d) Color-coded raw $dI/dU$ spectra measured along the green hatched line in (a). (e) Numerically calculated second derivative of the $dI/dU$ map measured along the green spectral (d set) or subtle double humps (red curve) can be recognized. With $d_{ss}$ reduced to 14 nm (curve 3) a clear splitting in two peaks becomes apparent, which amounts to $\Delta E \approx 18$ mV. The splitting further increases with decreasing step-to-step distance reaching $\Delta E \approx 26$ meV at $d_{ss} = 11$ nm (curve 4). At $d_{ss} = 7$ nm the peak intensities have drastically dropped (curve 5), even though the splitting still appears to increase ($\Delta E \approx 43$ meV). Eventually, at even smaller $d_{ss}$ the peaks become almost undistinguishable from the spectra of the terrace (curve 6 and dashed line, respectively).

Figure 2(d) represents the complete data set of color-coded raw $dI/dU$ spectra measured along the green hatched line in Fig. 2(a). This line section follows the contour of the left single-atomic step edge for about 500 nm from the bottom of Fig. 2(a) ($L = 0$) all the way to its termination point at the upper part of the the wedge-shaped plateau. For better contrast the numerically calculated second derivative of the $dI/dU$ signal, i.e., $d^2I/dU^2$, is presented in Fig. 2(e). At $0 \leq L \leq 200$ nm the data confirm the existence of a single peak at the Dirac energy $E_D = eU \approx 125$ meV. Local fluctuations by about $\pm 5$ mV are probably caused by statistical variations of the substitutional dopant concentration [37, 38]. As indicated by arrows in Fig. 2(d) and (e), for $L \geq 250$ nm the single peak which is known to represent the topological edge state of (Pb,Sn)Se [21] splits into two diverging branches. The data presented in Fig. 2(c) reveal the peak splitting increases with decreasing step-to-step distance up to $L \leq 400$ nm where $\Delta E \approx 28$ meV at $d_{ss} \approx 10$ nm. Above this line both peaks experience significant peak-broadening and strongly decrease in intensity, possibly due to the onset of attenuation. As presented in detail in Ref. [34] (Note 3), the results obtained on the right step edge are fully consistent with the results presented for the left step edge in Fig. 2. Furthermore, these main findings are independent of the specific type and shape of the step edges. For example,
Discussion — Similar observations of a coupling-dependent energy splitting have been made for zero-dimensional quantum dots [39] and two-dimensional layered van der Waals heterostructures [40]. In the latter case the opening of band-gaps was particularly prominent for out-of-plane–oriented MoS₂ orbitals which strongly overlap with graphene π-bands [10]. In fact, periodic boundary calculations presented in Fig. 6 of Ref. [39] suggest that the topological 1D edge state of (Pb,Sn)Se remains largely unaffected for terrace width \( d_{ss} \geq 86 \text{ nm} \), in reasonable agreement with our experimental data, cf. Fig. 2. Similar to the tight binding calculations presented in Ref. [21] a weak splitting may be visible for \( d_{ss} \geq 43 \text{ nm} \). For narrower terraces, however, a collapse of the double Dirac cone into a single cone with a crossing at the \( X \) point was predicted [36], in clear disagreement with our results which show the persistence of a split peak well below \( d_{ss} \leq 10 \text{ nm} \), cf. Fig. 2.

To simulate the observed behavior, we assume that the formation of the edge state results in an increased local density of states, crammed in a narrow region around the step edge. This regions of accumulated charge are— in a first-order approach—modeled by narrow identical quantum wells with a widths \( \sim 1 \text{ nm} \) perpendicular to the step edge and much larger along the step edge, determined as the intra-row coherence length. For a perfect step edge the topological edge mode would be completely delocalized along the step edge. The fact that the energy splitting observed at different points along the step edge depends on the step–to–step distance suggests that we don’t deal with a single, infinitely extended quantum state, but with electron wave functions which are localized to a certain extent due to their finite coherence length, potentially induced by disorder. Two quantum wells are separated by a distance \( d \) and affect the edge state, see Fig. 2(f). The well depth is assumed to \( \sim 100 \text{ meV} \), corresponding to the bulk band gap of the Pb₀.₇Sn₀.₃Se [31]. As shown in detail in Ref. [24] (Note 6), both quantum wells host wave functions which extend beyond their respective boundaries and decay exponentially with \( \exp(-\frac{d}{\xi}) \) outside the well, with the decay length \( \xi \).

When the wave functions overlap at low \( d \), see green area in Fig. 2(f), the exponentially decaying tails of the wave functions couple, resulting in an energy splitting \( \Delta E \). Indeed, a plot of \( \Delta E \) values extracted from Fig. 2(e) versus \( d_{ss} \) confirms the expected exponential dependence, see Fig. 2(g). The decay length of the 1D edge mode perpendicular to the step direction is determined to \( \xi = (10 \pm 1) \text{ nm} \). By taking \( \hbar \delta k \propto h/2\xi \) and \( \delta E \approx 10 \text{ meV} \) as the width of the single peak in spectrum 1 of Fig. 2(c), we estimate the group velocity to \( v_{gr} \approx \frac{\delta E}{\hbar \delta k} \approx 4.8 \cdot 10^5 \text{ m/s} \), in good agreement with earlier band dispersion measurements [31, 42].

Orthogonal step edges — Our investigations of wedge-shaped islands like those presented in Figs. 1 and 2 suggest that wedge angles are always \( \lesssim 30^\circ \). In contrast, much larger adjacent angles we regularly observed at points where two single-atomic steps intersect. For example, the STM data of Fig. 3(a) show two steps oriented along the planar high-symmetry (100) directions intersecting under an angle of \( \approx 90^\circ \). Again, we find a strongly enhanced \( dI/dU \) signal at the Dirac energy at the location of odd-atomic step edges and a strong reduction at the step crossing point, Fig. 3(b). In contrast to the situation observed for wedge-shaped step edges in Fig. 2 however, the tunneling spectra measured along the white arrow in panel Fig. 3(a) show no hints for any peak splitting, see Fig. 3(c) and (d) for raw spectra and \( d^3I/dU^3 \), respectively. The full data set is presented in Ref. [34] (Note 7).

The results presented in Figs. 2, 3 and Ref. [34] show that the topological edge state characteristic for odd-atomic step edges of the (Pb,Sn)Se (001) surface splits for adjacent steps (\( d_{ss} \geq 25 \text{ nm} \)) running in parallel or under an acute angle. In contrast, for step edges crossing perpendicularly no such peak splitting is observed. This latter behavior hints at wave functions which are orthogonal such that the solution of the Schrödinger equation becomes \( \Phi_n^* \Phi_m dV = \delta_{n,m} \), where the indices \( n,m \) stand for quantum numbers and \( \delta_{n,m} \) is the Kronecker symbol. Obvious choices for this quantum number might be the spin or the orbital moment of the edge state. We
are convinced that we can exclude the spin since results obtained within a tight binding [21] or envelope function model [36] indicate an out-of-plane or vanishing spin polarization of the edge state, respectively, both insufficient to explain the absence of a peak splitting for orthogonal edge states. In contrast, the contribution of a Se $p_z$ and $p_y$ orbital momentum Dirac points at $X$ and $Y$ has been described theoretically [36] [42] [43] and nicely corresponds with our experimental findings. In order to test this interpretation we investigated intersections of two single-atomic steps at angles below but close to 90°. Indeed, for single-atomic steps that cross under a 60° angle we observe a very weak but detectable splitting, see Ref. [34].

(Note 8). It is not clear, however, if the spin-orbit coupling in (Pb,Sn)Se plays any role. This question remains to be clarified by future theoretical investigations.

**Conclusion**—In conclusion, we have studied the interaction of adjacent 1D spin-polarized edge states which are localized at single-atomic step-edges of the TCI Pb$_{0.7}$Sn$_{0.3}$Se(001). Coupling at a step-to-step distance $d_{ss} \lesssim 25 \text{ nm}$ leads to the energy splitting of the single peak in LDOS at the Dirac point into two. The energy splitting increases exponentially with the reducing $d_{ss}$ and reveals the localized character of the topological edge mode, instead of single quantum state. At a very small separation ($d_{ss} \lesssim 10 \text{ nm}$) the two peaks strongly decrease in intensity. No peak splitting is found for perpendicular single-atomic step edges, suggesting an important role of the Se $p_z$ and $p_y$ orbital momentum of the edge state.

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