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In-surface confinement of topological insulator nanowire surface states

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The bandstructures of [110] and [001] Bi2Te3 nanowires are solved with the atomistic 20 band tight binding functionality of NEMO5. The theoretical results reveal: The popular assumption that all topological insulator (TI) wire surfaces are equivalent is inappropriate. The Fermi velocity of chemically distinct wire surfaces differs significantly which creates an effective in-surface confinement potential. As a result, topological insulator surface states prefer specific surfaces. Therefore, experiments have to be designed carefully not to probe surfaces unfavorable to the surface states (low density of states) and thereby be insensitive to the TI-effects. © 2015 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4931975]

Topological insulator (TI) materials such as Bi2Te3 have extraordinary surface properties.1–3 These make them a unique class of materials for applications such as low power electronic devices,4 spintronics,3 and quantum computation.5,6 TIs host surface states with the spin perpendicular to the surface normal, spin-locked relative to the electronic in-plane momentum. Backscattering of such surface electrons requires spin-flip processes. In TI devices that are free of magnetic impurities, surface electron backscattering is therefore unlikely. Then, the surface conductance is expected to be limited by the Fermi velocity.7 Experimental values of the Fermi velocity of Bi2Te3 surfaces show more than 25% variation.8–10 Experiments that determine Fermi velocities and other TI surface properties are often implicitly assuming different TI surfaces host the same physics.8,11,12 Even many theoretical studies of TI wires assume all wire surfaces are equivalent due to rotational wire symmetry.13–17 This assumption is only true for wires grown along [001] direction. In contrast, fabricated Bi2Te3 nanowires are grown in [110] direction and often have rectangular cross sections.18,19 The crystal structure of [110] Bi2Te3 nanowires shows different chemical surface composition: Some surfaces are composed of Te atoms only and other surfaces contain both Te and Bi atoms. To capture this important feature of the surface chemistry requires atomistic simulations. Only then, the important effect of in-surface confinement of surface states can be simulated. It is shown in this work for Bi2Te3 nanowires that this effect confines surface states to wire surfaces with specific chemistry. It is expected that similar situations hold for other TI-materials and geometries. If experiments are set to surfaces that are unfavorable to the surface states, where the topological insulator surface states have a low density of states, the experimental setup can be effectively insensitive to the TI physics.

In this work, atomistic sp3d5s* (20 band, spin-orbit coupling included) tight binding bandstructure calculations of Bi2Te3 nanowires are presented. In agreement with literature, the band gap of the Bi2Te3 nanowires is observed to close when the magnetic flux through the wire cross section is a half-integer flux quanta.13–17 Deviations from literature are found in the details of the surface state energies and surface Fermi velocities: Fermi velocities of chemically different surfaces differ and create an effective surface-state confining potential around the wire surface. Guided by the atomistic results, the analytical Fermi velocity model of Ref. 13 is augmented to cover these differences of the wire surface chemistry.

In Section II, the two methods used in this work are presented. This covers the atomistic tight binding features of NEMO5 and the analytical model of Ref. 13 augmented to cover variations in the wire surface chemistry. In Section III, the atomistic tight binding bandstructure results of NEMO5 for Bi2Te3 nanowires in the presence of magnetic fields are verified against literature.13 Bandstructures of rectangular Bi2Te3 nanowires with different ratios of pure Te and mixed atom type surfaces are presented then. These bandstructures serve as fitting targets for the surface Fermi velocities of the analytical model. Confinement effects of the surface states are shown after that. The analytical model is then used to explain this confinement of the wire surface states. Finally, the paper concludes with a summary of the finding.

In this work, atomistic sp3d5s* (20 band, spin-orbit coupling included) tight binding bandstructure calculations of Bi2Te3 nanowires are calculated with the multipurpose NanoElectronics Modelling Tool (NEMO5).20,21 A quintuple layer of Bi2Te3 consists of a sequence of five atomic layers: Te1, Bi, Te2, Bi, and Te1. “Te1” and “Te2” both denote Tellurium, but they differ in the chemical surrounding: The neighbor layers of Te1 consist of Te1 and Bi, whereas the Te2 atom layer lies between two Bi atom layers. Tight binding parameters for Bi2Te3 are taken from Ref. 22. The framework of NEMO5 and tight binding parameters give a surface Fermi velocity of 4.04 × 107 m/s for a 15 nm thick Bi2Te3 layer which agrees well with experimental data of Ref. 23.

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degenerate states are combined into symmetric and anti-symmetric states. Magnetic fields are included with the Peierl’s phase in symmetric gauge. All presented atomistic calculations are numerically very intense and require typically about $1 \times 10^6$ CPUs on the Blue Waters supercomputer.

To ease understanding of the tight binding results, the analytical model of Refs. 13–17 is augmented to support nanowires that are not rotationally symmetric along the transport axis. Typical examples for such wires are grown along [110] direction and are rectangular in the cross section. Such wires have two different types of facets: one type contains atoms of all types (“mixed surface”), while the second consists of Te$_1$ atoms (referred as “Te$_1$ surface”). For these surface types, different Fermi velocities are assumed: $v_{f1}$ for the mixed surface and $v_{f2}$ for the pure Te$_1$ one. The energy difference of the wire surface states $\Delta E$ with vanishing momentum ($k = 0$) is assumed to be

$$\Delta E = \begin{cases} \frac{v_{f1} \hbar}{P} & \text{(Type I, } W > T) \\ \frac{v_{f2} \hbar}{P} & \text{(Type II, } T > W) \end{cases}$$

Here, $P$ is the perimeter of the wire, equals to $(2W + 2T)$; $W$ is the dimension of one mixed surface (“width of the wire”) and $T$ is the dimension of one pure Te$_1$ surface (“thickness of the wire”). The two Fermi velocities are fit to match the surface state quantization of the tight binding results.

If not stated otherwise, all figures show tight binding results. All considered Bi$_2$Te$_3$ wires are grown along [110] direction.

It is an accepted rule in literature that the band gap of TI nanowires closes when the magnetic flux through the wire cross section agrees with half integer multiples of the magnetic flux quantum ($\Phi = \frac{h}{e}$). The largest band gap of the TI wires is expected with magnetic fluxes equal to integer multiples of the flux quantum. However, this knowledge is based on non-atomistic models (i.e., envelope function approximations). Figure 1 show the atomistic tight binding bandstructures of $12 \times 48$ nm$^2$ Bi$_2$Te$_3$ nanowires with varying magnetic fields along the wire growth direction. Here, the smaller facets are pure Te$_1$ type. The atomistic calculations indeed follow the rule of vanishing and maximal band gaps as a function of the magnetic flux. Equivalent behavior was observed for atomistic tight binding calculations of Bi$_2$Te$_3$ nanowires for a great variety of cross sections (ranging from $6 \times 24$ nm$^2$ to $60 \times 150$ nm$^2$). Atomistic tight binding calculations showed that different geometries and facet configurations do not alter the rule for band gap maxima and minima.

Although different wire configurations follow the same rule for the band gap with magnetic fields, the band structure details depend significantly on the ratio of pure Te$_1$ and mixed facet dimensions. This is exemplified in Fig. 2(a): compare the atomistic structure of two $12 \times 48$ nm$^2$ Bi$_2$Te$_3$ nanowires that differ in the size of the pure Te$_1$ and mixed facets. For later reference, wires with larger mixed than pure Te$_1$ facets are termed “Type I”, the other cases as “Type II”. The bandstructures of the two cases in Fig. 2(b) show an energy difference of the wire surface states with vanishing magnetic flux quantum $\Delta E$ of 6.2 meV for the type I and 10.5 meV for the type II nanowire of Fig. 2(a). This difference in $\Delta E$ is in contrast to non-atomistic models that cannot distinguish wires of type I and type II.

Figure 3 shows the $\Delta E$ as a function of the perimeter of Bi$_2$Te$_3$ nanowires grown in [110] (a) and [001] (b) direction. In all cases, the longer edge of the wire cross section is kept constant. All presented atomistic calculations showed that different geometries and facet configurations do not alter the rule for band gap maxima and minima.

FIG. 1. Bandstructures of $12 \times 48$ nm$^2$ Bi$_2$Te$_3$ nanowires (type I) with varying magnetic field along the wire axis. The wire bandstructure without a magnetic field (a) or with a magnetic field corresponding to the magnetic flux quantum (c) has the largest band gap and every state is double degenerate. Bandstructures with magnetic fields corresponding to 0.5 (b) and 1.5 (d) magnetic flux quanta have disappearing band gap and are only degenerate at $k = 0$.

FIG. 2. (a) Atomic structures of $12 \times 48$ nm$^2$ Bi$_2$Te$_3$ nanowires grown along the [110] direction in the two possible surface configurations. In the type I wire, the pure Te$_1$ facet is much smaller than in the type II configuration. (b) Bandstructures of the nanowires of (a) show different quantization energies $\Delta E$ for the surface states at $k = 0$. 

FIG. 3. $\Delta E$ as a function of the perimeter of Bi$_2$Te$_3$ nanowires grown in [110] (a) and [001] (b) direction.
constant to 48 nm, while the smaller edge varies. For [110] grown wires of type I, the mixed surface is kept constant, while for type II wires, the pure Te₁ surface is constant. Note that all surfaces of [001] wires are of mixed atom type and equivalent. Therefore, a type I and type II distinction is meaningless for [001] wires, i.e., the surfaces A and B are equivalent. Therefore, nanowires grown in [110] direction show a strong dependence of AE on the facets ratio configuration, while AE of [001] wires is virtually independent of that. Due to the linear nature of all results shown in Fig. 3, the analytical model discussed in the method section gives an almost perfect fit to the numerical atomistic tight binding data. It turns out, the Fermi velocities for the [110] wires are \( v_{f1} = 1.28 \times 10^5 \text{ m/s} \) and \( v_{f2} = 4.36 \times 10^5 \text{ m/s} \) and for the [001] wires \( v_{f1} = v_{f2} = 1.28 \times 10^5 \text{ m/s} \). Note that mixed surfaces that are chemically equivalent (i.e., having the same portion of Te₁, Te₂, and Bi atoms) have always the same Fermi velocity, irrespective of the wire growth direction.

The values of the Fermi velocities \( v_{f1} \) and \( v_{f2} \) vary with the size of the bigger facet (kept constant in Fig. 3). For instance, if the bigger facet has the dimension of 120 nm the following Fermi velocities are found: \( v_{f1} = 1.39 \times 10^5 \text{ m/s} \) and \( v_{f2} = 4.50 \times 10^5 \text{ m/s} \).

This change in the Fermi velocity indicates confinement effects within the wire surface. This in-surface confinement is illustrated in Fig. 4, which show the absolute squared wavefunctions for the first 3 surface states of the type I nanowire in Fig. 2 with energies above about 0.12 eV and momentum \( k = 0.025 \text{ nm}^{-1} \). The surface states are delocalized over the total wire surface, but they are mainly located at the mixed type facets. The number of minima of the surface states envelope (shown in Fig. 4(b)) increases with the state’s energy—similar to confined electronic states in quantum wells. The calculations also show stronger in-surface confinement effect with increasing momentum. The in-surface confinement vanishes at the \( \Gamma \)-point. This finding can be understood with the analytical model of Eq. (1): The dispersion difference of the two different nanowire facets (pure Te₁ and mixed type) yields an effective, momentum dependent potential between the facet types (see schematic of Fig. 5(a)). This potential vanishes at the \( \Gamma \)-point and increases with finite momenta. This potential effectively creates a system of 2 quantum wells within the wire surface (see schematic of Fig. 5(b)). The surface states envelopes’ confinement is typical for such quantum wells.

In this work, NEMO5’s atomistic tight binding models are applied on Bi₂Te₃ nanowire bandstructures for wires grown along [110] and [001] direction. The atomistic representation unveils for experimentally common, rectangular [110] nanowires two chemically different types of surfaces. It is imaginable that TI wires with other than rectangular geometries still host in-surface confinement if the facets are chemically distinct enough. Chemically distinct surfaces host topological insulator surface states with different Fermi velocities. The surface states spread over all facets, but they are subject to momentum-dependent in-surface confinement. This finding is critical for experiments on TI-surfaces: TI-properties should be measured only on TI-favorable surfaces. It is imaginable that variations of measured Fermi velocities in Bi₂Te₃ might trace back to the different Fermi velocities of the different surface kinds. This situation is different in [001] wires due to their chemically equivalent facets. Both situations can be well reproduced with an analytical model presented in this work as well.

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