Transport properties of a 3D topological insulator based on a strained high mobility HgTe film

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We investigated the magnetotransport properties of strained, 80 nm thick HgTe layers featuring a high mobility of \( \mu \sim 4 \times 10^5 \text{cm}^2/\text{V}\cdot\text{s} \). By means of a top gate the Fermi-energy is tuned from the valence band through the Dirac type surface states into the conduction band. Magnetotransport measurements allow to disentangle the different contributions of conduction band electrons, holes and Dirac electrons to the conductivity. The results are in line with previous claims that strained HgTe is a topological insulator with a bulk gap of \( \approx 15 \text{meV} \) and gapless surface states.

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The discovery of two (2D)- and three-dimensional (3D) topological insulators (TI), a new material class with insulating bulk and topologically protected, conducting surface states, has opened a new and exciting research field in condensed matter physics \([1]–[10]\). Although quite a number of different, especially Bi based materials \([11]–[14]\), belong to this category, materials which combine high charge carrier mobility and insulating bulk are still scarce. This is mostly due to the fact that Bi-based 3D TIs are heavily-doped alloy films with a very low mobility \( \mu \sim 10^3 \text{cm}^2/\text{V}\cdot\text{s} \) and a high bulk carrier density of \( 10^{17} \) - \( 10^{19} \text{cm}^{-3} \). HgTe based 2D TI, on the other hand, are characterized by very high mobilities enabling the discovery of the quantum spin Hall effect \([15]\). A recent analysis of the sequence of quantum Hall plateaus suggests that also strained HgTe layers constitute a 3D TI. The strain opens a gap in the gapless semimetal HgTe, so that the TI properties can be explored by tuning the Fermi energy into the bulk gap and probing the transport properties of the gapless surface states. Although the strained HgTe film has a much higher mobility \( \mu = (3–4) \times 10^4 \text{cm}^2/\text{V}\cdot\text{s} \), the high bulk carrier density and the absence of a top gate has complicated the detection of the 3D TI so far \([16]–[17]\).

The strain in HgTe layers, grown by molecular beam epitaxy (MBE), stems from a 0.3% lattice mismatch between HgTe and CdTe. The corresponding critical film thickness for pseudomorphic growth is larger than 100 nm meaning that thinner films adopt the substrate lattice constant. Due to this strain a small gap of \( \sim 15 \text{meV} \) opens (see below) in the bulk energy spectrum of the film. Within the bulk gap the gapless surface states reside. The charge neutrality point of the corresponding Dirac cone is located in the valence band \([16]\). Thus Dirac electrons with higher energy dominate electrical transport if the Fermi energy is located in the bulk gap. This allows us to map the bandstructure in the vicinity of the bulk gap by magnetotransport experiments as is outlined below.

In this Letter we report on the transport properties of high mobility, 80 nm wide, strained HgTe films equipped with a gate. The low disorder, manifested in high charge carrier mobilities, together with the possibility to tune the Fermi energy from the valence via the gap into the conduction band, enables us to probe the 2D Dirac surface states when the Fermi energy is in the energy gap of the bulk. Since HgTe films grown on CdTe suffer from dislocations due to the lattice mismatch, our 80 nm thick HgTe films were separated from the CdTe substrate by a 20 nm thin Cd_{0.7}Hg_{0.3}Te buffer layer. This buffer layer increases the electron mobility by an order of magnitude (up to \( 4 \times 10^5 \text{cm}^2/\text{V}\cdot\text{s} \)) and reduces the bulk impurity concentration to values of order \( 10^{16} \text{cm}^{-3} \) (see supplemental material). We fabricated and investigated two types of devices: one with the upper HgTe surface uncapped and the other one covered with a 20 nm Cd_{0.7}Hg_{0.3}Te cap layer (see Fig. 1(a) and supplement). Since all measured major properties (magnetotransport traces, bulk energy gap etc.) of capped and uncapped films were found to be similar, we will only focus on the uncapped HgTe film below. For transport measurements the films were patterned into Hall bars supplied with top gates. Cross-sections of the devices are sketched in Fig. 1(a). Each Hall bar consists of three 50 \( \mu \text{m} \) wide segments of different lengths (100, 250, and 100 \( \mu \text{m} \)) with eight voltage probes. Ohmic contacts to the active layer were formed by alloying indium. For gating two types of dielectric layers were used, giving similar results: 100 nm SiO\(_2\) and 200 nm of Si\(_3\)N\(_4\) grown by plasma chemical vapor deposition of SiH\(_4\) + N\(_2\)O at 100°C or 80 nm Al\(_2\)O\(_3\) grown by atomic layer deposition. In both cases TiAu was deposited as metallic gate. Magnetotransport measurements were performed at temperatures \( T \) between 1.5 K and 15 K and in magnetic fields \( B \) up to 10 T. Several devices from the same wafer have been studied.

Fig. 1(b) shows the typical resistivity \( \rho_{xx} \) at \( B = 0 \) and Hall resistance \( \rho_{xy} \) at \( B = 1 \text{T} \) as a function of gate...
at the same value of $V_g$ approximately: For $V_g > 2 V$ the $\rho_{xx}$ does hardly change with $T$ but changes dramatically for $V_g < 2 V$ where $\rho_{xx}$ varies by a factor 2 between $T = 1.9 K$ and $15 K$. This behavior is ascribed to strong Landau scattering which coexisting electron and hole states, similar to the one observed in HgTe quantum wells [16].

Another feature in $\rho_{xx}(V_g)$ emerges at $V_g = 4 V$ (Fig. 1(b)), accompanied by a change of the slope of $N_{xx}^\text{s-Drude}(V_g)$ (Fig. 1(c), extracted from low-field Shubnikov-de Haas (SdH) oscillations (see below)). We suggest that this feature marks the gate voltage at which the Fermi level starts to enter the conduction band. Therefore, the data presented in Fig. 1 and Fig. 2a imply that the gap opens between 2 and 2.5 V (top of valence band) and closes around 4 V (bottom of conduction band). Then only Dirac states localized at the two surfaces of the strained 80 nm HgTe film contribute to transport. Outside this $V_g$ region Dirac electrons and bulk electrons (holes) conduct in parallel. A sketch of the correspond-

FIG. 1. (a) Cross-section of our heterostructures. (b) Left vertical axis: $V_g$ dependence of $\rho_{xx}$ at $T = 1.9 K$ and for $B = 0$. Right vertical axis: Hall resistance $\rho_{xy}(V_g)$ for the second sample. (c) Electron $N_x(V_g)$ and hole $P_x(V_g)$ densities at different $V_g$ extracted from the Drude model and high-field SdH oscillations. The electron density of the top layer, $N_{xx}^\text{top}$ is extracted from low-field SdH oscillations (see text). (d) Electron and hole mobilities, averaged over all participating charge carriers, at different gate voltages. Between $V_g = 0.5 V$ and 2 V the fits to the two-carrier Drude model were not reliable enough to extract electron and hole mobilities (see supplement).
ing density of states versus energy is shown in the inset of Fig. 2(a). Using the electron densities extracted at \( E_c \) and \( E_v \) and the calculated k-linear dispersion of Dirac electrons \([16]\) we estimate a gap size of \( \approx 15 \text{ meV} \). This value is very close to the one calculated in \([16]\) for a strained HgTe film. The same value we found for the \( \text{Cd}_{0.7}\text{Hg}_{0.3}\text{Te} \) capped HgTe film, indicating that the surface states are not affected by the precise nature of the interface.

To check the validity of our picture further we resort to magnetoresistance measurements in a perpendicular magnetic field, displayed in Fig. 2(b). The magnetoresistance (MR) within classical Drude theory for two groups of carriers (labeled by index 1 and 2) is, for small fields, proportional to \( B^2 \). Its magnitude, normalized to \( \rho_{xx}(B=0) \) is proportional to 

\[
\frac{\sigma_i}{\sigma_1 + \sigma_2} (\mu_1 \mp \mu_2)^2, \quad \text{where } \sigma_i \text{ and } \mu_i \text{ are conductivity and mobility of the respective carrier species } i \text{ at } B = 0.
\]

The sign between \( \mu_1 \) depends on the carriers’ polarity: in case the carriers have the same polarity (e.g. Dirac electrons in the conduction band) the "+" sign applies while in the case of different polarity (electrons and holes coexist) the mobilities add, thus resulting in a strong MR. Hence the maximum of the MR magnitude is expected to occur in the semimetal state with the Fermi level residing in the valence band near the charge neutrality point, where \( N_s \approx P_s \) and \( \sigma_1 \) and \( \sigma_2 \) become comparable. A significant change of the MR magnitude is expected when the Fermi energy moves from the valence band into the gap where only Dirac electrons reside. For this single carrier type the MR is small. Corresponding normalized \( \Delta \rho_{xx}(V_g) \) and \( \Delta \rho_{xx}(B) \) data, displayed in Fig. 2(b), are in accord with this expectation. For \( V_g < 2 \text{ V} \) a large parabolic MR is observed with a pronounced MR maximum at \( V_g = 1 \text{ V} \). In contrast, for \( V_g > 2 \text{ V} \), i.e. where Dirac electrons prevail, the MR drops by a factor of up to 10. The MR is expected to rise again when the Fermi level moves from the gap into the conduction band, as two (or more) groups of carriers with different mobility are involved, i.e. Dirac and bulk electrons. Bulk electrons near the bottom of the conduction band are expected to have a much lower mobility than Dirac electrons. A corresponding increase is indeed visible in Fig. 2(b) (marked by "\( E_c \)."

Important extra information can be obtained from experiments in quantizing magnetic fields. SdH oscillations and quantized Hall steps can be seen in Fig. 3(a) as a function of \( V_g \) for \( B = 4 \text{ T} \). The maximum in \( \rho_{xx} \) at \( V_g \approx 1 \text{ V} \) corresponds to the charge neutrality point at which the Hall resistance (voltage) changes sign. On the left hand side, where low mobility holes dominate, no quantum Hall steps appear. In contrast, on the right hand side where Dirac fermions and conduction band electrons prevail, quantized Hall steps develop.

Surprisingly, the Hall steps extend into a \( V_g \) region where conduction band and Dirac electrons coexist. The electron density, extracted from high-field SdH oscillations, plotted in Fig. 1(c), is identical to the one obtained from classical Drude fits. This means that the filling factors at high \( B \)-fields are determined by the total electron concentration \( N_{\text{tot}} \), i.e. bulk and surface electrons. Similarly, at large negative \( V_g \) and high \( B \), the hole density extracted from SdH oscillations and from Drude theory are nearly equal, while at smaller bias the SdH data deliver smaller hole densities. This suggests that the filling factor in the valence band is given by the difference of bulk hole and surface electron density. This is similar to the situation observed in GaSb/InAs heterojunctions where electrons and holes coexist \([23]\). A central observation is thus that bulk (which in our case is a 80 nm wide HgTe quantum well) and surface charge carriers determine jointly the high-\( B \) LL filling factors. The \( \rho_{xx}(B) \) minima do not vanish indicating parallel conduction probably due to the sides of the HgTe layer, which are oriented parallel to the applied magnetic field \([16]\). The quantized Hall steps on the electron side display, as in Brüne’s work \([16]\), even and odd integer plateau values, thus indicating different carrier densities for top and bottom surface.

This is not surprising as it is a consequence of screening, i.e., a part of the electric field gets screened by the top layer of Dirac electrons. Applying an electric field to the top gate hence results in different filling rates.

![FIG. 2. (a) \( \rho_{xx}(V_g) \) for different temperatures. The inset sketches the density of states in our system. (b) \( \rho_{xx}(B)/\rho_{xx}(B = 0) \) vs. \( V_g \) for different magnetic field values. "\( E_c \)" and "\( E_v \)" mark the onset of valence and conduction band.](image-url)
$dN_s^{\text{top(bot)}}/dV_g$. These rates can be easily estimated if $E_F$ is in the gap, i.e. between $\sim 2$ V and 4 V. Then the change of top and bottom electron density is given by 
\[ \frac{\Delta N_s^{\text{top}}}{\Delta N_s^{\text{bot}}} = 1 + \frac{q^2D_{\text{HgTe}}}{\varepsilon_{\text{HgTe}}\varepsilon_0}, \]
where $D$ is the density of states of Dirac electrons on the top surface, and $d_{\text{HgTe}}$ and $\varepsilon_{\text{HgTe}}$ are thickness and dielectric constant of the HgTe layer, respectively. Inserting typical values we obtain $\Delta N_s^{\text{top}}/\Delta N_s^{\text{bot}} = 3 - 5$. Below we show that the experimentally observed difference in top and bottom layer filling rate is close to this expected value. Besides $N_s$, we plot in Fig. 1(c) also the electron density of the top surface as a function of $V_g$. The corresponding data are obtained from the following consideration: Assuming that the carrier densities are equal at the flat band condition, i.e. at $V_g = 0$ V, the electron density on the top surface becomes significantly higher for $V_g > 2$ V. The higher carrier density of the top surface is expected to be connected with a higher electron mobility $25$. This offers an opportunity to separate the electron density of top and bottom layer experimentally. As the higher mobility of the top layer is connected to a smaller Landau level broadening, Shubnikov de Haas commences at lower magnetic fields and dominate the low field magnetoresistance oscillations in Fig. 3(c). This is reflected in different periods of SdH oscillations in low and high $B$-fields, displayed in Fig. 3(d). The $1/B$ positions of the SdH minima vs. filling factors can be fitted by two straight lines corresponding to the carrier density $N_s^{\text{top}} = 1 \times 10^{11}$ cm$^{-2}$ of the top layer and the total carrier density $N_s^{\text{tot}} = 1.4 \times 10^{11}$ cm$^{-2}$. $N_s^{\text{top}}$, extracted from low-field SdH-oscillations for different $V_g$ is shown in Fig. 1(c). In both cases we assumed spin-resolved LLs. The reduced slope of $N_s^{\text{top}}(V_g)$ for $V_g > 4$ V is a clear signature that $E_F$ moves in the conduction band. Since $dN_s^{\text{tot}}/dV_g$ is constant, $dN_s^{\text{top}}/dV_g$ decreases when the bulk electron density $N_s^{\text{bulk}}$ starts to appear, i.e., when $dN_s^{\text{bulk}}/dV_g > 0$. For $B > 1$ T Landau quantization gets resolved in the lower mobility bottom layer, too and, due to electron redistribution, the two surfaces (and for $V_g > 4$ V also the bulk electrons) act like a single 2DEG with density $N_s^{\text{tot}}$. The redistribution of electrons in the TI state is possible via contacts, ungated conducting regions and via side facets of the HgTe film.

With $N_s^{\text{tot}}$, acquired from high-field SdH oscillations, and $N_s^{\text{top}}$ we can, for $E_F$ between 2 V $< V_g < 4$ V, i.e. in the TI regime, calculate the carrier density of the bottom layer, $N_s^{\text{bot}}$. Corresponding data are also shown in Fig. 1(c). The slope of $N_s^{\text{top}}(V_g)$ is by a factor of $\sim 3$ higher than the one of $N_s^{\text{bot}}(V_g)$. This is in line with the effect of screening, discussed above.

In summary we have shown that an analysis of magnetotransport data in strained high-mobility HgTe layers brings out the different carrier types contributing to transport at different Fermi level positions. A strong magnetoresistance, observed in the valence band where electrons and holes coexist, is absent in the gap where only surface electron contribute, and rises again when the Fermi energy enters the conduction band. An analysis of high- and low-field quantum oscillations highlights the interplay of the different carrier types and allows to probe the carrier density of top and bottom layer in the TI regime separately.

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[20] Note that the experiment delivers averages over both surfaces and, in certain voltage regions, also over part of the bulk. This can be seen from the band scheme (Fig. S5, supplement) which shows the position of $E_F$ at different $V_g$.
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[25] Comparison of CdHgTe capped and uncapped heterostructures (see supplement for details) proves that they have the same average mobility for $E_F$ in the gap. We thus conclude that for the same carrier density mobilities on top and bottom surface are approximately the same.
Supplemental material for “Transport properties of a 3D topological insulator based on a strained high mobility HgTe film”

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SAMPLES DETAILS

Heterostructure and Hall bar

We fabricated and investigated several devices based on two types of heterostructures. Both types were grown by molecular beam epitaxy on a (013)-oriented CdTe substrate and have very similar structure: the essential part is a strained 80 nm thick HgTe film (Fig. S1(a)). The two heterostructures types differ in the top layer sequence: in the first structure (named “uncapped”) the HgTe is uncapped on top while the second one (“capped”) is covered with a 20 nm Cd_{0.7}Hg_{0.3}Te cap layer. Since HgTe films grown on CdTe suffer from dislocations due to lattice mismatch, thus causing low mobility (< 4 × 10^4 cm^2/V·s) and high impurity concentration (> 10^{17} cm^{-3}) in the bulk [16, 17], our 80 nm thick HgTe films were separated from the CdTe substrate by a 20 nm thin Cd_{0.7}Hg_{0.3}Te buffer layer (Fig. S1(a)). The buffer layer increases the electron mobility by an order of magnitude (up to 4 × 10^5 cm^2/V·s) and reduces the bulk impurity concentration to a value about 10^{16} cm^{-3}, estimated from the carrier density in ungated samples (see below).

For transport measurements the films were patterned into Hall bars. The central part of several Hall bars is covered with a top gate. A top view of the devices is sketched in Fig. S1(b). Each Hall bar consists of three 50 µm wide segments of different lengths (100, 250, and 100 µm) with eight voltage probes. The ohmic contacts to the active layer were formed by alloying indium. In our devices we used two types of dielectric layers for gating: 100 nm SiO_2 and 200 nm of Si_3N_4 grown by plasma chemical vapor deposition of SiH_4 + N_2O at 100°C or 80 nm Al_2O_3 grown by atomic layer deposition. In both cases TiAu was deposited as metallic gate.

Fig. S1. (a) Cross-section of devices with top gate. We fabricated and investigated also ungated devices made from both kinds of structures. (b) Schematic top view of the Hall bars studied.

Ungated sample properties

Fig. S2(a) displays \( \rho_{xx}(B) \) and \( \rho_{xy}(B) \) traces of one of the ungated samples. These traces are typical for a system with coexisting electrons and holes. Fitting these traces by using the classical two-component Drude model (see next paragraph) allows to determine the values of 2D electron and hole densities and mobilities. Electron and hole densities of the 80 nm thick HgTe layer are expressed as 2D carrier densities. Values of carrier densities and mobilities extracted from such fits are presented in the boxes underneath the figures in Fig. S2. In comparison with previous results [16, 17] our samples exhibit much smaller carrier densities of typically \((1−2) \times 10^{14} \text{cm}^{-2}\) and higher mobilities (> 10^5 cm^2/V·s). Both, density decrease and mobility increase indicate that the amount of charged impurities has been significantly reduced by introducing of the CdHgTe buffer layer. The amount of charged impurities in the
structure can be estimated from charge neutrality, i.e. $N_{\text{imp}} \approx |N_s - P_s| = 1.6 \times 10^{11}$ cm$^{-2}$. Taking the thickness of the HgTe layer into account this results in a volume impurity density of $N_{\text{imp}}^{\text{3D}} \approx 2 \times 10^{16}$ cm$^{-3}$.

![FIG. S2. $\rho_{xx}(B)$ (black dots) and Hall resistance $\rho_{xy}(B)$ (red dots) of ungated (a) and gated (b) samples. Solid lines are fits to the Drude model for a two-component 2D electron-hole gas. Extracted electron and hole densities and mobilities are shown in the boxes underneath.](image)

**Gated and ungated samples**

Whether the Dirac surface states are affected by gate fabrication we studied with some care. In Fig. S2 we compare $\rho_{xx}(B)$ and $\rho_{xy}(B)$ of the ungated sample (Fig. S2(a)) and of the gated one (Fig. S2(b)) for an applied gate voltage $V_g = -1.4$ V. The most important feature here is the overall similarity of the corresponding traces in Fig. S2(a) and (b): by applying an appropriate gate voltage we can bring the gated system into the same state as the ungated one. Direct comparison of electron and hole densities and mobilities, extracted from Drude fits, shows that they are within 20% the same. This is the same level of difference found between samples fabricated the same way. This indicates that gate fabrication does not change the band diagram of the structure qualitatively but introduces charged impurities and/or defects in the system with a density of order of $(1 - 2) \times 10^{11}$ cm$^{-2}$, varying from sample to sample.

**CdHgTe-capped and open HgTe films**

Besides investigating the influence of the gate fabrication on the sample’s charge state we also checked how capping of the HgTe film with a CdHgTe film affects the transport properties (see Fig. S1(a)). In addition we compared different kinds of dielectric layers under the gate. The following results show that the properties of the HgTe film are only weakly dependent on the presence of a cap or different dielectric layers. A comparison between two samples is presented in the Fig. S3. The first one (labeled “uncapped” in Fig. S3) has an uncapped HgTe film with an insulating SiO$_2$ layer on top and the second one (“capped” in the Fig. 3) is a HgTe film with CdHgTe cap layer and a Al$_2$O$_3$ insulator under the gate. Since dielectric constant and insulator thickness are different for these samples the change of carrier density $N_s - P_s$ with gate voltage $V_g$ differs correspondingly: $d(N_s - P_s)/dV_g^{\text{SiO$_2$}} \approx 0.75 \times 10^{11}$ cm$^{-2}$/V for the Si$_3$N$_4$-covered sample and $d(N_s - P_s)/dV_g^{\text{Al$_2$O$_3$}} \approx 2.5 \times 10^{11}$ cm$^{-2}$/V for the Al$_2$O$_3$-covered one. In order to compare the samples one has to rescale the gate voltage axis with the ratio of the respective specific capacitances. In doing so, the experimental traces in Fig S3(a), (c) and (d) of the two samples follow each other closely. Only some features in $\rho_{xx}$ and $\rho_{xy}$ in the presence of a magnetic field and near the charge neutrality point in Fig. S3(b) and (c) differ slightly. In spite of these small differences the general properties of the samples are still the same: all the observed features in transport of the uncapped sample are more or less visible in the measurements of the capped sample and even the estimated gap value between conduction and valence bands is the same – $\Delta \approx 15$ meV. The
reproducibility between the two samples made from different wafers is amazing and points to some kind of rigidity of the band structure and Dirac surface states with respect to external disturbances.

FITTING OF MAGNETORESISTANCE AND HALL RESISTANCE WITHIN THE TWO-COMPONENT CLASSICAL DRUDE MODEL

We used the classical Drude model to extract electron and hole densities and mobilities by fitting $\rho_{xx}(B)$ and $\rho_{xy}(B)$ at small magnetic field and for a fixed gate voltage (see Fig. S4). A detailed description of the model can be found in [18] and many other textbooks. Here we present only its application to our system. The model is based on the assumption that, in the system studied, $n$ groups of noninteracting carriers exist. Each group $i = 1...n$ has charge $q_i = \pm q$ (electrons or holes), density $n_{2D}^i$ and mobility $\mu_i$. The conductivity of the system is given as the sum over the conductivities of each group: $\sigma_{xx} = \sum_{i=1}^{n} \sigma_{xx}^i$ and $\sigma_{xy} = \sum_{i=1}^{n} \sigma_{xy}^i$. After tensor inversion one obtains the corresponding expressions for $\rho_{xx}(B)$ and $\rho_{xy}(B)$.

In our system we can distinguish four main groups of carriers: bulk holes in the valence band, bulk electrons in the conduction band and two groups of Dirac fermions (DF) which exist at the two surfaces of the HgTe film. This picture is based on band structure calculations of Brüne et al. [10], supplementary). A simplified band scheme which describes the position of the Fermi energy at different gate voltages $V_g$ is presented in Fig. S5. Though the Drude model can, in principle, be used for any number $n$ of carrier groups, a reasonable fit is only possible for $n = 1$ (trivial case) and $n = 2$. Already for three groups of carriers the expressions become complicated and contain 6 fitting parameters. Then fitting become ambiguous. Due to the gap in the bulk spectrum we have in our case no more than three groups of carriers at $E_F$ (see Fig. S5). To reduce the number of groups further we treat the Dirac electrons of top and bottom surface as one group. This is justified as they have the same charge and since the difference in mobility and carrier density causes only weak corrections to $\rho_{xx}(B)$ and $\rho_{xy}(B)$. When extracting carrier density and

FIG. S3. Comparison of the gated sample with uncapped HgTe film and with SiO$_2$+Si$_3$N$_4$ insulator ("uncapped", black lines) and the sample with the HgTe film capped by a CdHgTe layer and with Al$_2$O$_3$ insulator ("capped", red lines). (a) Resistivity $\rho_{xx}$ vs gate voltage at zero magnetic field and $T = 1.9$ K. (b) $\rho_{xx}(V_g)$ at $B = 1$ T. (c) Hall resistance $\rho_{xy}(V_g)$ at $B = 1$ T. (d) Charge $N_s(V_g) - P_s(V_g)$ in the system determined by fitting of $\rho_{xx}(B)$ and $\rho_{xy}(B)$ at fixed gate voltages by the two-component Drude model.
mobility of surface electrons we always obtain an average of top and bottom surface. Within this approximation the number \(n\) of Drude groups is reduced to 2 (bulk holes and Dirac electrons or bulk electrons and Dirac electrons when \(E_F\) is located in the valence or conduction band, respectively). With \(E_F\) in the bulk gap \(n\) is even reduced to one as then only Dirac electrons exist.

In case of two of groups of carriers (with index 1 and 2) the system displays a classical parabolic magnetoresistance (MR) with magnitude \(d^2\rho_{xx}(B)/dB^2\big|_{B=0} = \frac{\sigma_1 \sigma_2}{\sigma_1 + \sigma_2}(\mu_1 \pm \mu_2)^2\), where \(\sigma_i\) and \(\mu_i\) are the corresponding conductivities of each group \(i\) at zero magnetic field. The sign between the \(\mu_i\)'s depends on the carriers’ polarity: in case the carriers have the same polarity (e.g., two groups of electrons) the """" sign applies while in the case of different polarity (electrons and holes coexist when Fermi level resides in the valence band) the mobilities add, thus resulting in a strong MR. Hence two groups of Dirac electrons with similar values of their mobility will cause only a small MR effect. This is actually the reason why we can treat top and bottom surface electrons as one species. In contrast, we expect a maximum magnetoresistance when the Fermi level is in the valence band near the the charge neutrality point where \(N_e \approx N_h\) and \(\sigma_1 \approx \sigma_2\) holds. Note that the charge neutrality point is not equal to the Dirac point as the latter is located in the valence band. A significant change of the MR magnitude is expected when the Fermi level moves from the valence band into the gap where only Dirac electrons reside (topological insulator state) and the magnitude of MR drops by up to a factor of 10. The data in Fig. S4(b) show this drastic change in MR depending on \(V_g\), i.e. \(E_F\). The drastic change of \(\rho_{xy}(B)\) between 1 V and 2 V in Fig. S4(b) also heralds this change in the contributing charge carriers. At 2 V \(\rho_{xy}(B)\) becomes nearly linear indicating that for \(V_g > 2\) V only the electrons of the surface states contribute.

FIG. S4. Resistivity \(\rho_{xx}(B)\) (top) and Hall resistance \(\rho_{xy}(B)\) (bottom) for different gate voltages. Fits to the two-component Drude model are drawn as solid lines. In the semimetal state (a) \((V_g = -3...0\) V\) it is possible to fit \(\rho_{xy}\) over the \(B\)-range and \(\rho_{xx}\) around \(B = 0\). (b) Gate voltage region corresponding to the transition from the valence band (semimetal state) to the topological insulator state. Between \(V_g = 0\) V and \(V_g = 2\) V the fits work less well, most probably to to the proximity to the charge neutrality point, at which small inhomogeneities are more detrimental. (c) Magnetoresistance and Hall resistance in the topological insulator region \((V_g = 2...4\) V\) and at the transition to the bulk electron metal \((V_g > 4\) V\). For all gate voltages \(\rho_{xy}(B)\) shows essentially a linear dependence on \(B\). This indicates that the current is carried by only one sort of charge carriers. A more detailed analysis of \(\rho_{xy}(B)\) shows a slight nonlinearity (see derivative \(d\rho_{xy}(B)/dB\) in the insert) near \(B = 0\) indicating the presence of two (two surfaces with Dirac electrons in the TI state) or more (Dirac electrons on top and bottom surface and bulk electrons in the conduction band) groups of carriers with the same charge but with different mobilities. However the magnitude of the nonlinearity is so small making it impossible to distinguish these different groups reliably within the Drude model. For \(V_g \geq 2\) V we determined only the overall electron density and their average mobility without differentiation between several groups.
FIG. S5. Schematic band diagrams of our system for different gate voltages. The Dirac point is located in the valence band indicated by the onset of the yellow bars which illustrate the Dirac electrons on top and bottom surface. In the band structure shown here, flat band occurs at $V_g = 0\,\text{V}$. Applying a bias voltage results in band bending. When $E_F$ is in the gap in (e) and (f) the screening on the electron density of the back surface can be estimated (see main text). The band diagram under different bias condition show that the electron (hole) densities and mobilities are always averages over top and bottom surfaces and, depending on bias, also involving bulk electrons (see (g)) and bulk holes (see (b)).

The next change of the MR behavior is expected when the Fermi level enters the conduction band. (Fig. S4(c)). There, the presence of low-mobility bulk electrons results in a rise of the MR magnitude (see, e.g., $\rho_{xx}(0.1\,\text{T}) - \rho_{xx}(0)$).

The monotonous decrease of $d^2\rho_{xx}(B)/dB^2$ (curvature around $B = 0$) with increasing $V_g$ is reversed in the small voltage interval between $V_g = 4\,\text{V}$ and $V_g = 4.5\,\text{V}$. Because of the relatively small bulk electron density when $E_F$ just enters the conduction band and their small contribution to the overall conductivity, the resulting MR is weaker compared to the situation in the valence band where carriers of different polarity are involved. For further increasing $V_g$ the bulk electrons’ contribution to the conductivity increases but, on the other hand, the mobility increases also resulting in a reduced MR magnitude for $V_g > 5\,\text{V}$, as displayed in Fig. S4(c).
Thereby the presence of the bulk electrons, starting to appear at \( \sim 4 \) V is observed in classical transport. However, the Drude fits are not sensitive enough to disentangle density and mobility of surface and bulk electrons and one could determine the gate voltage corresponding to crossing of Fermi level with the bottom of conductive band but the sensitivity of the Drude fitting technique is not enough to determine exact values of bulk electrons density and mobility.