Bulk Fermi surface coexistence with Dirac surface state in Bi$_2$Se$_3$: a comparison of photoemission and Shubnikov-de Haas measurements

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Shubnikov de Haas (SdH) oscillations and Angle Resolved PhotoEmission Spectroscopy (ARPES) are used to probe the Fermi surface of single crystals of Bi$_2$Se$_3$. We find that SdH and ARPES probes quantitatively agree on measurements of the effective mass and bulk band dispersion. In high carrier density samples, the two probes also agree in the exact position of the Fermi level $E_F$, but for lower carrier density samples discrepancies emerge in the position of $E_F$. In particular, SdH reveals a bulk three-dimensional Fermi surface for samples with carrier densities as low as $10^{17}$ cm$^{-3}$. We suggest a simple mechanism to explain these differences and discuss consequences for existing and future transport studies of topological insulators.

Recently, a new state of matter, known as a topological insulator, has been predicted to exist in a number of materials: Bi$_{1-x}$Sbx$_2$, Bi$_2$Se$_3$, Bi$_2$Te$_3$ and Sb$_2$Te$_3$[1, 2]. This state of matter is characterized by a full band gap in the bulk of the material, but with a gapless, dissipationless surface state. The surface state is comprised of counter-propagating spin states, which create a dispersion of a single, massless Dirac cone that is protected by time-reversal symmetry. The experimental realization of this state could mean significant advances in spintronic devices, quantum computation and much more besides.

As a result there has been great excitement in the last year after the discoveries of various ARPES experiments[3], and more recently from scanning-tunneling measurements[4] that such a state appears to exist in nature. Amidst this flurry of recent results, it is easy to forget that these same materials have been the subject of careful and thorough research for much of the 20th century. However, common to all the unambiguous measurements of the Dirac cone is the use of surface-sensitive probes. Only recently have transport measurements emerged specifically investigating the surface state (Refs [5]), all of which note the dominance of the bulk conductivity. It is thus of great interest to perform a coordinated study of these materials using both bulk transport experiments and surface sensitive ARPES experiments. Here we report results of these investigations. The transport experiments reveal quantum oscillations that indicate a bulk band structure and Fermi surface volume that monotonically change with doping. For carrier densities in the range $\sim$10$^{19}$ cm$^{-3}$, the transport extracted band structure is in quantitative agreement with the bulk band structure determined by ARPES which also observes the Dirac dispersion of the surface state. The quantitative agreement between ARPES and SdH provides additional support for the existence of novel band structure in these materials. For lower carrier density samples down to $10^{17}$ cm$^{-3}$ we observe SdH oscillations which unambiguously pin the Fermi level in the bulk conduction band, with a high level of consistency across all samples measured from the same batch. While ARPES places $E_F$ near the SdH level for some samples, there are others from the same batch whose $E_F$ is found to reside into the bulk gap. We discuss possible explanations for these discrepancies and the implications for transport studies of surface Dirac Fermions in samples near a metal-insulator transition.

The material Bi$_2$Se$_3$ can be grown without the introduction of foreign dopants as either n or p type[12, 13] though is more commonly found as the former because the dominant defects tend to be Se vacancies. Quantum oscillatory phenomena, which provides evidence of bulk metallic behavior has been reported by Kohler et al. [12] on low carrier density samples and more recently by Kulkbachinskii [14] on high carrier density samples. Below a

![FIG. 1:](image-url) (a) Temperature dependence of two typical samples of Bi$_2$Se$_3$ with carrier densities differing by two orders of magnitude. (b) Shows the carrier density $N_c$, (c) the resistivity $\rho_0$ at $T=2K$ and (d) the mobility for samples of different thicknesses. Each sample was a cleave from a parent sample, so that the surface area of each sample was kept constant.
carrier density of $7 \times 10^{18} \text{cm}^{-3}$, the band structure is well approximated by a single parabolic band, making the interpretation of transport measurements transparent [14]. Two $n$-type samples with carrier densities differing by two orders of magnitude are shown in Figure 1. For the low carrier density samples an upturn in the resistivity is seen, which levels off at sufficiently low temperature. This behavior has been attributed to the presence of an impurity band whose thermally activated conductivity is comparable to the band conductivity until carriers freeze out at around 30K [13, 14]. This behavior is not apparent in the higher carrier density materials, where the band conductivity always dominates. Even though we have reduced the carrier density by 2 orders of magnitude, the resistivity increases by one, suggesting that the mobility has increased in the low carrier density samples, consistent with previous measurements [14]. The low carrier density samples are around an order of magnitude smaller than those of reported topological insulators including Sn doped Bi$_2$Te$_3$ ($n_e \sim 1.7 \times 10^{13} \text{cm}^{-3}$) [14] or Ca doped Bi$_2$Se$_3$ ($n_e \sim 5 \times 10^{18} \text{cm}^{-3}$) [3] and as a result may be better candidates in which to observe the transport properties dominated by the topological surface state.

Single crystals of Bi$_2$Se$_3$ have been grown by slow cooling a binary melt. Elemental Bi and Se were mixed in alumina crucibles in a molar ratio of 35:65 for batch S1 ($n_e = 5 \times 10^{17}$), 34:66 for batch S2 ($n_e = 3 \times 10^{17}$), 34:66 for batch S3 ($n_e = 2.3 \times 10^{17}$), and 40:60 for batch S4 ($n_e = 2.3 \times 10^{19}$). The mixtures were sealed in quartz ampules and raised to 750°C and cooled slowly to 550°C, then annealed for an extended period. Crystals can be cleaved very easily perpendicular to the (0 0 1) axis. Measurements of the resistivity and Hall effect were measured in a 14T PPMS using a standard 4-probe contact configuration and Hall measurements were performed using a 6-probe configuration. For the latter, only data which was linear in the low field limit was used to avoid mixing with longitudinal components. In addition to this precaution, signal from positive and negative field sweeps was subtracted to extract the odd (Hall) components of the signal, after which the carrier density is extracted in the usual way. ARPES measurements were performed at beam line 10.0.1 of the Advanced Light Source (ALS) at Lawrence Berkeley National Laboratory. Measurement pressure was kept $<3 \times 10^{11}$ Torr, and data were recorded by Scienta R4000 analyzers at 15K sample temperature. The total convolved energy and angle resolutions were 16meV and 0.2° (i.e. $<0.007(A^{-1})$ or $<0.012(\Lambda^{-1})$ for photoelectrons generated by 48eV photons), at which energy the cross-section for both surface state and bulk bands is strong.

In Figure 2 we show complimentary ARPES and SdH data on samples from the same batch, with carrier density determined by the Hall effect of $n_e = 2.3 \times 10^{19} \text{cm}^{-3}$. The SdH reveals an anisotropic pocket of frequency 155 T, corresponding to a filling of around 160meV (the band structure is not parabolic at this filling and so we assume similar band structure parameters as Kohler et al. [12] characterizing similar carrier density samples of Bi$_2$Se$_3$). ARPES results on samples from the same batch, show the Fermi level 150meV above the bottom of the conduction band in good quantitative agreement. Similarly, the effective mass (see below) extracted from SdH is in good quantitative agreement with that measured by ARPES.

In Figure 3 we illustrate angle dependent SdH data (a) taken at 1.8K, on a sample from batch S1 with a lower carrier density of $10^{17} \text{cm}^{-3}$. The SdH signal reveals a pocket that is approximately an ellipsoid elongated along the $c$ axis, consistent with measurements by Kohler et al. from the 1970’s [12] on samples with similar carrier densities. For a two-dimensional pocket expected from the surface state, quantum oscillations should vary as $1/\cos \theta$, where $\theta$ is the angle between the $c$ axis and the field direction, so the present observations must originate from a 3D Fermi surface existing in the bulk. It has been shown by Kohler et al. and more recently by Kulbachinskii et al. that the conduction band structure for these low carrier densities is approximately parabolic [12, 14], and so the band filling can be estimated by $E_F = \frac{4\pi^2 k_F^2}{m_e^*}$, where $A_K$ is the area of the Fermi surface in Fourier space. We estimate the Fermi energy to be 18meV above the bottom of the conduction band.

In Figure 4 (b) we show the derivative of the longitudinal magnetoresistance of a sample from batch S3 and a fit of the entire data set using the usual Lifshitz-Kosevich formalism, to extract the effective mass and Dingle temperature $T_D$, with fit shown in (c). Fitting the entire data set, which is often more accurate than tracing the amplitude of the Fourier transform, our fit yields $m^* = 0.15m_e$ and $T_D = 3.5K$, for this frequency (F=14T). Similar data for samples from batch S4 give $m^* = 0.125m_e$, $T_D = 4K$ and F=155T. The mean free

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FIG. 2: (a) ARPES band dispersion on samples of Bi$_2$Se$_3$ with carrier density $2.3 \times 10^{19} \text{cm}^{-3}$ (batch S4). (b) Due to the quantization of the energy spectrum into Landau levels (LLs), oscillations appear in the magnetoresistance known as SdH oscillations. The SdH oscillations here are for a sample taken from the same batch as in (a) at $\theta = 0$, corresponding to a oscillatory frequency of $F = 155T$, consistent with $E_F \sim 160\text{meV}$. ARPES and SdH are in good agreement for these high carrier density samples.
path is calculated using the orbitally averaged velocity and scattering time extracted from $T_D$ yields $l_{S3} \sim 60\text{nm}$ and $l_{S4} \sim 220\text{nm}$. This data is wholly consistent with the very complete SdH studies of Kohler et al.\cite{12,13} and more recently by Kulbachinskii et al.\cite{14}. In addition, the data was reproduced with high consistency on a number of samples from the same batch, and even on samples from different batches with similar growth parameters.

ARPES data on samples from the same batches as those shown in Figure 3 determining the effective mass as $m^\ast - 0.13$ in very good quantitative agreement with SdH. However, the exact placement of the Fermi level in the band structure reveals some disagreement. In Figure 4 we illustrate photoemission data for two separate samples from batch S1. In (a) the Fermi level is near the bottom of the conduction band in agreement with SdH, while in (b) it is in the gap (about 60meV below the conduction band), crossing the Dirac cone with apparently no bulk contribution. A number of samples from similar batches, such as batch S2 and S3, also have shown similar variation in $E_F$ crossing the gap on some samples. While $E_F$ determined by photoemission appears to show some variation, it is important to note that the measured $E_F$ from ARPES is either near or below the SdH $E_F$.

Despite the good agreement of SdH and ARPES on high carrier density samples, and the agreement of the effective mass and other band parameters on the low carrier density samples, the discrepancy in the position of the Fermi level requires explanation. Such differences can occur for a number of reasons, for example due to sample variation within a batch, or perhaps due to variation in the exposure of cleaved surfaces before a photoemission measurement. However, it should be noted that the SdH frequency does not appear to vary significantly within a batch for up to 20 samples measured in the present study and so the former seems an unlikely scenario. Another reason for the discrepancy may be that atmospheric exposure of transport samples has contaminated them with an $n$-type dopant causing them to appear bulk $n$-type. Figure 4 (c) illustrates photoemission data for a sample cleaved in air. The Dirac cone of the surface state remains robust and the bulk conduction band appears partially occupied. Such doping may lead to a 3D Fermi surface pocket appearing in SdH oscillations if the contamination is deep enough and allows for sufficiently long mean free paths.

To investigate this possibility further we measure the thickness dependence of the transport by systematically thinning a single sample. Cleaving was achieved with tape, keeping the surface area of the resulting samples relatively constant and allowing direct comparison of data sets of each cleave. Though the samples are vulnerable to deformations between cleaves, only data from mirror-like flat samples is presented. In most cases these samples still exhibited quantum oscillatory phenomena, confirming the high quality of the cleaved samples. Figure 4: ARPES data on samples of Bi$_2$Se$_3$ from batch S1. The horizontal lines show the crossing of the Fermi level ($E - E_F = 0$) and the Dirac crossing. (a) Band structure measured by ARPES results on samples from batch S1 showed the Fermi level near the SdH level of $\sim 15\text{meV}$ from the bottom of the conduction band. (b) Measurement on another sample from batch S1 showed the Fermi level in gap. Some other samples from S2 and S3 also showed the Fermi level in the bulk gap. The variation might be due to the lower carrier density of these samples, and the surface band structure is more susceptible to small amounts of surface contamination. c) Band structure of a sample also from batch S1 which was cleaved in atmosphere and exposed for 10s, showing significant $n$-type doping with large bulk conduction band pocket.

FIG. 3: (a) Magnetotransport for samples from S1. As the angle is swept the frequency of the oscillation varies according to the topology of the Fermi surface. For a two-dimensional pocket the expected dependence is $1/\cos\theta$ (shown in green in the inset). The observed angle dependence is clear evidence for a closed ellipsoidal Fermi surface pocket, similar to that observed by Kohler et al.\cite{12}. Similar SdH data was gathered on batch S2 and S3 on a number of samples. Samples from batch S3 showing the temperature dependence of the derivative of the SdH signal in (b) and a fit to the data shown in (c) from which the effective mass, Dingle temperature and oscillatory frequency can be extracted.
A function of thickness at the density of carriers and mobility. Within our error bars, each quantity seems to vary weakly down to 3μm in thickness. Although the residual resistivity and carrier density varies slightly (possibly from disorder related to slight sample deformation, despite the precautions mentioned above), the mobility remains almost constant as a function of thickness at μ ~ 1m²/Vs. In summary, the transport is insensitive to the thickness, suggesting that the SdH oscillations are not a consequence of atmospheric contamination, but originate from the intrinsic band conductivity of the bulk.

A final scenario for the discrepancy is that the band structure is distorted near the surface due to space-charge accumulation. This is known to occur in many semiconductors, such as InSb or CdTe[12, 16], whereby the bulk band structure bends as the surface is approached. Typically, such bending occurs over a surface depletion layer zd, which can be calculated by solving the Poisson equation to yield zd = κe0ΔV/εns[17], where κ is the DC dielectric permittivity (estimated from these samples as ~ 113[18]) and ΔV is the difference in energy between the surface and bulk state. We estimate zd ~ 40nm for the low carrier density samples and zd ~ 2nm for the high carrier density samples. A schematic representation of the band bending is shown in Figure 5. The present argument suggests that discrepancies between ARPES and SdH can be explained, even expected for low carrier density samples. In addition, due to the small value of ns, these samples are likely more susceptible to a small amount of surface contamination, especially if the un-contaminated surface Ef is in the gap (as illustrated in by ARPES on atmosphere exposed samples). This may help explain why there is some variability in the Fermi level of ARPES data but not in the SdH data.

Much theoretical work has emerged on the dramatic consequences of the surface state on transport properties[19]. Yet over several decades of experimental study, such properties have not been observed. Recently, Aharonov-Bohm and universal conductance fluctuations have been observed which may be due to the surface state [10, 11], but even in these cases the conductance appears bulk at the temperatures considered. Conventionally, such intrinsically doped materials can become ‘insulators’ by either a Mott-like or an Anderson transition. The first can occur when the Bohr radius ab = κℏ²/m∗e² falls below the Thomas-Fermi screening length λTF, so that wavefunctions cannot overlap. This can be estimated using λTF = κε0/(2πε²g(EF)), where g(EF) is the number of states per unit volume per unit energy, estimated by Middendorff et al. [20]. In the present case, the large κ and small m∗ tend to make ab very large. For the lowest carrier density samples investigated here ab ~ 3nm and λTF ~ 4nm, which places is this material on the metal-insulator boundary. The carrier density can also be reduced by introducing foreign dopants which ‘drain’ the excess carriers and pin the chemical potential μ in the gap. For hydrogenic like impurities this can be very effective, but in the present materials impurity bands often form instead. At high enough impurity densities the carriers may become Anderson localized. Such samples are characterized by a high carrier density with very low mobility, leading to a negative gradient in the temperature dependence of the resistivity. This may be the case for example in Bi2SbPbSe3 which has μ ~ 30m²cm⁻¹ carrier density ns ~ 5×10¹⁸cm⁻³[21]. An Anderson insulator is generally bad news for topological insulators, because even though at zero temperature the bulk conductivity σ = 0, at finite temperature the transport may remain dominated by bulk hopping mechanisms.

In conclusion, the present study reveals substantial agreement between transport and ARPES measurements of the Fermiology of Bi2Se3, in particular for samples with large carrier densities. However, for samples with carrier densities approaching 10¹⁷cm⁻³, discrepancies emerge as to the exact position of the Fermi level. We have confirmed the bulk nature of the transport by the thickness dependence of the Hall effect, resistivity and mobility. Furthermore SdH data is highly consistent between different samples from the same batch. Interestingly, the carrier densities measured here are an order of magnitude smaller than those of the topological insulators recently reported in the literature[3, 5, 11, 12]. ARPES and STM have been invaluable tools in revealing the physics of topological insulators, providing compelling evidence for the presence of the topologically protected Dirac surface state. The present results should stimulate further theoretical work as to the consequences of the coexistence of bulk and surface states in a single sample as well as innovation in novel ways to fabricate these materials so the bulk state can be cleanly eliminated.

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[1] J. C. Y. Teo, L. Fu, and C. L. Kane, Physical Review B (Condensed Matter and Materials Physics) 78, 045426 (2008).

[2] H. Zhang, C. Liu, X. Qi, X. Dai, Z. Fang, and S. Zhang, Nat Phys 5, 438 (2009), ISSN 1745-2473.

[3] D. Hsieh, Y. Xia, L. Wray, D. Qian, A. Pal, J. H. Dil, J. Osterwalder, F. Meier, G. Bihlmayer, C. L. Kane, et al., Science 323, 919 (2009).

[4] Y. L. Chen, J. G. Analytis, J. Chu, Z. K. Liu, S. Mo, X. L. Qi, H. J. Zhang, D. H. Lu, X. Dai, Z. Fang, et al., Science 325, 178 (2009).

[5] D. Hsieh, Y. Xia, D. Qian, L. Wray, J. H. Dil, F. Meier, J. Osterwalder, L. Patthey, J. G. Checkelsky, N. P. Ong, et al., Nature 460, 1101 (2009), ISSN 0028-0836.

[6] Z. Alpichshev, J. G. Analytis, J. H. Chu, I. R. Fisher, Y. L. Chen, Z. X. Shen, A. Fang, and A. Kapitulnik, 0908.0371 (2009).

[7] P. Roushan, J. Seo, C. V. Parker, Y. S. Hor, D. Hsieh, D. Qian, A. Richardella, M. Z. Hasan, R. J. Cava, and A. Yazdani, Nature 460, 1106 (2009), ISSN 0028-0836.

[8] K. K. Gomes, W. Ko, W. Mar, Y. Chen, Z. Shen, and H. C. Manoharan, 0909.0921 (2009).

[9] A. Taskin and Y. Ando, Physical Review B 80 (2009).

[10] H. Peng, K. Lai, D. Kong, S. Meister, Y. Chen, X. Qi, S. Zhang, Z. Shen, and Y. Cui, Nat Mater 10.1038/nmat2609 advance online publication (2009).

[11] J. G. Checkelsky, Y. S. Hor, M. H. Liu, D. X. Qu, R. J. Cava, and N. P. Ong, Physical Review Letters 103, 246601 (2009).

[12] H. Kohler, Physica Status Solidi (b) 58, 91 (1973).

[13] H. Kohler and A. Fabbicius, physica status solidi (b) 71, 487 (1975).

[14] V. A. Kulbachinskii, N. Miura, H. Nakagawa, H. Arimoto, T. Ikaida, P. Lostak, and C. Drasar, Physical Review B 59, 15733 (1999).

[15] P. D. C. King, T. D. Veal, M. J. Lowe, and C. F. McConville, Journal of Applied Physics 104, 083709 (2008).

[16] R. K. Swank, Physical Review 153, 844 (1967).

[17] W. Monch, Semiconductor surfaces and interfaces (Springer, 2001), ISBN 3540679022, 9783540679028.

[18] U. R. O. Madelung and M. Schulz, in Non-Tetrahedrally Bonded Elements and Binary Compounds I (1998), pp. 1–12.

[19] D. Lee, Physical Review Letters 103, 196804 (2009).

[20] A. Middendorff, H. Kohler, and G. Landwehr, Physica Status Solidi (b) 57, 203 (1973).

[21] J. Kasparova, C. Drasar, A. Krejcova, L. Benes, P. Lost’ak, W. Chen, Z. Zhou, and C. Uher, Journal of Applied Physics 97, 103720 (2005).