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Phys. Rev. B 97, 035147 — Published 22 January 2018

DOI: 10.1103/PhysRevB.97.035147
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(Dated: December 21, 2017)

We report magnetotransport properties of BaZnBi$_2$ single crystals. Whereas electronic structure features Dirac states, such states are removed from the Fermi level by spin-orbit coupling (SOC) and consequently electronic transport is dominated by the small hole and electron pockets. Our results are consistent with three dimensional (3D) but also with quasi two dimensional (2D) portions of the Fermi surface. The spin-orbit coupling-induced gap in Dirac states is much larger when compared to isostructural SrMnBi$_2$. This suggests that not only long range magnetic order but also mass of the alkaline earth atoms A in ABX$_2$ (A = alkaline earth, B = transition metal and X=Bi/Sb) are important for the presence of low-energy states obeying the relativistic Dirac equation at the Fermi surface.

I. INTRODUCTION

Rapid development of graphene and topological matter could induce transformational changes in both information and energy science.1–4 Main physical observables are Dirac states with Berry phases and high mobility due to suppressed backscattering.5 This generates large linear magnetoresistance (MR), quantum Hall effect and is of high interest for electronic and optical device fabrication.6–14

Dirac states could be found not only at the surface of topological insulators or graphene but also in multiband bulk crystals, giving rise to magnetoresistivity mobility up to $\mu_\text{MR}\sim 3400 \text{ cm}^2/\text{Vs}$, comparable to that in graphene and topological insulators.15–19 In contrast to Na$_3$Bi and C$_3$As$_2$, ABX$_2$ materials feature quasi-two-dimensional (quasi-2D) electronic transport of Dirac states.15,16,20 Moreover, ABX$_2$ incorporate strongly correlated alternating Mn-Bi magnetic layers and sometimes rare earth atoms on A site.21–23 Localized magnetic moments with sufficient hybridization could provide a possible route to the Kondo effect and to transforming Dirac into a Weyl semimetal, separating the degenerate Dirac cones.24 This could result in spin-dependent band splitting and quasi-2D spin-polarized transport.

Coupling of magnons to Dirac states is indicated by Raman measurements in CaMnBi$_2$, where the enhanced interlayer exchange coupling drives a charge gap opening.25 In contrast, magnetic dynamics can be described by the Heisenberg model and is not influenced by Dirac fermions.26 It is of interest to establish conditions for Dirac states at the Fermi surface in ABX$_2$ materials since the momentum-anisotropy of Dirac states might tailor Dirac current for applications.17,27–30 In this study we have synthesized BaZnBi$_2$ single crystals that feature crystal structure identical to SrMnBi$_2$ and BaMnBi$_2$. Our first-principle calculations show presence of Dirac states near Fermi level and portions of Fermi surface with quasi-2D character, similar to other AMnBi$_2$ materials. However, SOC opens a larger gap in Dirac states when compared to SrMnBi$_2$, leaving small hole and electron pockets that dominate the electronic transport.

II. EXPERIMENTAL DETAILS

Single crystals of BaZnBi$_2$ were grown from an excess Bi flux.31 Ba, Zn and Bi granules were mixed in the ratio of Ba:Zn:Bi = 1:1:10, put into an alumina crucible and then sealed in a quartz tube. The quartz tube was heated to 1073 K, held there for 6 h, then cooled to 593 K at a rate of 2.4 K/h where the excess Bi flux was decanted using a centrifuge. Shiny needle-like single crystals with typical size $5 \text{ mm} \times 1 \text{ mm} \times 1 \text{ mm}$ were obtained. The element analysis was performed using an energy-dispersive x-ray spectroscopy (EDX) in a JEOL LSM-6500 scanning electron microscope, confirming 1:1:2 stoichiometry. X-ray diffraction (XRD) data were obtained by using Cu $K_\alpha$ ($\lambda = 0.15418 \text{ nm}$) radiation of a Rigaku Miniflex powder diffractometer on crushed crystals. Heat capacity and magnetotransport measurement up to 9 T were conducted in a Quantum Design PPMS-9 on cleaved and polished single crystals in order to remove residual Bi droplets on the crystal surface [Fig. 1(a)]. Thin Pt wires were attached to electrical contacts made with Epotek H20E silver epoxy, producing contact resistance of about 10 $\Omega$. Sample dimensions were measured with an optical microscope Nikon SMZ-800 with 10 $\mu\text{m}$ resolution. Magnetotransport in high magnetic field up to 18 T was conducted at the National High Magnetic Field Laboratory (NHMFL) in Tallahassee. Resistivity was measured using a standard four-probe configuration for both $\rho_\alpha$ and $\rho_\perp$. Both measurements were conducted with 0.3 mA 16 Hz ac current excitation. All magnetotransport measurements were performed in transverse configuration with current always perpendicular to magnetic field.
The de Haas van Alphen (dHvA) oscillation experiments were performed at NHMFL Tallahassee. The crystals were mounted onto miniature Seiko piezoresistive cantilevers which were installed on a rotating platform. The field direction can be changed continuously between parallel (θ = 0°) and perpendicular (θ = 90°) to the c-axis of the crystal. For first-principles band structure calculations, we applied the WIEN2K\textsuperscript{32} implementation of the full potential linearized augmented plane wave method in the generalized gradient approximation\textsuperscript{33} of density-functional theory with SOC treated in a second variational method. The basis size was determined by $R_{\text{mt}}K_{\text{max}} = 7$ and the Brillouin zone was sampled with a regular $13 \times 13 \times 13$ mesh containing 196 irreducible $k$ points to achieve energy convergence of 1 meV. The Fermi surface was plotted in a 10 000 k-point mesh.

### III. RESULTS AND DISCUSSION

The unit cell of BaZnBi\textsubscript{2} crystals can be indexed in the $I4/mmm$ space group by RIETICA software [Fig. 1(a)].\textsuperscript{34} The lattice parameters $a = b = 0.4855(2)$ nm and $c = 2.1983(3)$ nm agree with the previously reported values.\textsuperscript{35} Hence, the BaZnBi\textsubscript{2} [Fig. 1(a) inset] features an identical structure, but a somewhat compressed c-axis when compared to SrMnBi\textsubscript{2} and BaMnBi\textsubscript{2}. The heat capacity of BaZnBi\textsubscript{2} [Fig. 1(b)] is approaching the Einstein value of $4 \cdot 3 \cdot R = 99$ J mol$^{-1}$ K$^{-1}$, where $R$ is the universal gas constant 8.31 J mol$^{-1}$ K$^{-1}$. From $C/T$ as a function of $T^2$ at low temperature [Fig. 1(b) inset] we obtain an electronic heat capacity $\gamma = 0.48(1)$ mJ mol$^{-1}$ K$^{-2}$ and a slope of 2.61 mJ mol$^{-1}$ K$^{-4}$. From the latter, the Debye temperature $\Theta_D = 155(1)$ K is
SOC reveals Dirac-like bands derived from the 5p orbitals of Bi square lattices near the X points in the Brillouin zone, which are fairly close to the Fermi level [Fig. 3(a)]. The density of states near the Fermi level [Fig. 3(b)] is dominated by partial contribution of Bi square lattices and displays a pseudogap feature, again indicating that the Dirac points are close to the Fermi level.

FIG. 3. (Color online). (a) Band structure calculations for hypothetical BaZnBi$_2$ where Ba is 5% closer to Bi2 plane, all other parameters remaining unchanged. This leaves mass of A the same, while mimicking steric effects (lattice parameter change) due to different A size. (b) Band structure calculation for BaZnBi$_2$ in its true unit cell but where spin-orbit coupling is selectively switched off on the Ba atoms only.

obtained from $\Theta_D = (12\pi^4 NR/5\beta)^{1/3}$ where $N$ is the atomic number in the chemical formula. The value of $\gamma$ is much smaller than that of the isostructural antiferromagnetic compound BaMnBi$_2$. Fig. 1(c) shows the magnetic field dependence of in-plane magnetoresistance MR = $[\rho(B) - \rho(0)]/\rho(0) \times 100\%$ at different temperatures up to 50 K. The linear unsaturated magnetoresistance is evident at low temperatures and in high magnetic fields. Linear unsaturated magnetoresistance could be a consequence of magnetotransport of the lowest Landau level where all components of the resistivity tensor should be linear in temperature. Other possible mechanisms of the large linear magnetoresistance include the mobility fluctuations in a strongly inhomogeneous system, $\omega_c^2 \tau \sim 1$ limit in thin films (where $\omega_c = eB/m^*\alpha$ is cyclotron frequency and $\tau^{-1}$ is scattering rate) and charge neutrality in a compensated semimetal. The former should not apply in BaZnBi$_2$ since our sample is stoichiometric crystal without doping/disorder. To get more insight into magnetotransport properties below we present results of first principle calculations.

The first-principles band structure in the absence of SOC reveals Dirac-like bands derived from the 5p$_x$ and 5p$_y$ orbitals of Bi square lattices near the X points in the Brillouin zone, which are fairly close to the Fermi level [Fig. 2(a)]. The density of states near the Fermi level [Fig. 2(b)] is dominated by partial contribution of Bi square lattices and displays a pseudogap feature, again indicating that the Dirac points are close to the Fermi level. However, spin-orbit coupling introduces sizeable gaps into the Dirac states possibly [Fig. 2(c,d)] creating more small hole and electron pockets, while the electron states near the X points remain quasi-two-dimensional. Corresponding calculated Fermi surfaces [Fig. 2(e,f)] are quite different. This result suggests that linear MR [Fig. 1(c)] is unlikely to arise from the lowest Landau level magnetotransport.

It appears that the SOC effect on the A atoms predominates the steric effect due to different sizes of A atoms, since the same structural data were used in the calculations with and without SOC. To further distinguish the two A-site effects, we performed non-SOC calculations for BaZnBi$_2$ with Ba atoms moved 5% closer to the Bi2 planes and all the other parameters remaining unchanged. In this case, the steric effect was enhanced considerably but it has little impact on the Bi2 bands showing Dirac points [Fig. 3(a)]. In addition, we carried out calculations for the experimental structure with SOC on the Bi atoms only. We found [Fig. 3(b)] that the gapping out of the Dirac state along the $\Gamma$ - X line is nearly vanishing. This shows that the large SOC on the A-site atoms do facilitate the gapping out of the Dirac state in BaZnBi$_2$.

Figure 4(a) shows the temperature dependences of in-plane ($\rho_a$) and out-of-plane ($\rho_c$) resistivity at 0 and 9 T for BaZnBi$_2$ single crystal. Temperature dependence of resistivity is very anisotropic ($\rho_c/\rho_a \sim 55$ at 2 K and $\sim 30$ at 300 K). The broad hump at $\sim 150$ K at $\rho_c$ indicates the crossover from high-T incoherent to low-T coherent conduction in a quasi-2D system. The temperature dependence of magnetoresistance (MR) for electric current along a- and c-axes in magnetic fields up to 9 T are shown in the Fig. 4(b). The linear transverse MR is established for both current directions, however at much lower field for $\rho_c$. The temperature-dependent cantilever signal [Fig. 4(c)] shows clear oscillations below 40 K. The oscillatory component obtained by subtracting a smooth background is periodic in 1/B [Fig. 4(d)]. The fast Fourier transform (FFT) analysis yields four frequencies at $F_a = 175$ T, $F_b = 210$ T, $F_\gamma = 237$ T, and $F_\eta = 280$ T, as shown in Fig. 4(e). From the Onsager relation $F = (\Phi_0/2\pi^2)A_F$, where $\Phi_0$ is the flux quantum and $A_F$ is the orthogonal cross-sectional area of the Fermi surface, we estimate $A_\alpha = 1.7$ nm$^2$, $A_\beta = 2.0$ nm$^2$, $A_\gamma = 2.3$ nm$^2$, and $A_\eta = 2.7$ nm$^2$, corresponding to about 1.0%, 1.2%, 1.4%, and 1.6% of its total Brillouin zone (BZ) in the (001) plane, respectively. These are likely to arise from the small pockets in spin-orbit induced compensated Fermi surface [Fig. 4(f)]. The cyclotron mass can be obtained from fitting the temperature dependence of the oscillation amplitude at different frequencies using Lifshitz-Kosevich formula: $A \sim \{m^*\alpha(T/B)/\sinh(\alpha m^*T/B)\}$ where $\alpha = 2\pi^2 k_B/e\hbar \approx 14.69$ T/K, $m^* = m/m_e$ is the cyclotron mass ratio ($m_e$ is the mass of free electron). The fitting results are presented in Fig. 4(f), the rather small cyclotron masses corresponding to different frequencies are estimated to be $m^*_a = 0.14 m_e$, $m^*_\alpha = 0.12 m_e$, $m^*_\beta = 0.11 m_e$, and $m^*_\gamma = 0.11 m_e$, in agreement with heat capacity result [Fig. 1(b)].

The response of the carriers to the applied magnetic field and the magnitude of MR is determined by the mobility in the plane orthogonal to the applied magnetic
FIG. 4. (Color online). (a) Temperature dependence of the in-plane resistivity $\rho_a(T)$ and c-axis resistivity $\rho_c(T)$ of the BaZnBi$_2$ single crystal in B = 0 and 9 T magnetic fields, respectively. Large resistivity anisotropy indicates that the c-axis interlayer coupling of conductive Bi crystallographic square layers is small. (b) Field dependence of transverse magnetoresistance at 2 K for $\rho_a$ (filled circles) and $\rho_c$ (filled squares), respectively. (c) Cantilever oscillation as a function of magnetic field at different temperatures and its oscillatory component as a function of $1/B$ (d). (e) FFT spectra at various temperatures, data in (c), (d), and (e) use same legend. (f) Temperature dependence of the FFT amplitudes at different frequencies, the solid line is the fitting curve using Lifshitz-Kosevich formula.

There should be no significant angle-dependent MR in the case of isotropic three-dimensional Fermi surface. For quasi-2D electronic systems, the 2D states respond only to the perpendicular component of the magnetic field $B|\cos(\theta)|$; hence MR oscillations should be observed in quasi-2D conducting states. Figures 5(a,b) show the angular dependence of the $\rho_a$. The rotating magnetic field [Fig. 5(a) inset] is parallel (perpendicular) to the crystallographic c-axis when $\theta = 0^\circ$ ($90^\circ$), respectively. The $MR(\theta)$ at 18 T is consistent with the $|\cos(\theta)|$ angular dependence [solid red line in Fig. 5(b)], commonly observed in AMnBi$_2$ (A = Ca, Sr, Ba). This reflects the contribution of quasi-2D cylinder-like states away from the $\Gamma$ point in the Brillouin zone [Fig. 2(f)].

Angular-dependent quantum oscillation offer further insight into the geometry of the Fermi surface. Cantilever oscillatory components are shown in Fig. 5(c), corresponding FFT spectra are shown in Fig. 5(d). The small oscillation amplitude at $8^\circ$ reflects the angle error bar in measurement in which amplitude of oscillation should be zero at strict $0^\circ$ and $90^\circ$. As shown in Fig. 5(e), $F_\beta$ and $F_\gamma$ can be fitted very well by $1/\cos(\theta)$, indicating the quasi-2D feature of $\beta$ and $\gamma$ Fermi surface.
$F_3$ and $F_4$ with similar angular dependence might come from the cylindrical states near A point, and $F_5$ can be attributed to small pockets near the center of BZ. On the other hand, it is difficult to trace the weak angular dependence of $F_n$ which could come from the large hole pockets around $\Gamma$ point.

It is instructive to compare mechanism of Dirac states formation with isotropical materials $\text{AMnBi}_2$ ($A = \text{Sr}$, $\text{Ba}$). We note that nearly-linear Dirac-like energy dispersion is evident in first-principle calculation in all materials. However, a small SOC-induced gap of about 0.05 eV appears near the Dirac point when SOC is included in SrMnBi$_2$.\textsuperscript{17,27} SOC-induced gap in Bi$_2$-derived states of about (0.2 - 0.3) eV is much larger in BaZnBi$_2$ [Fig. 2(c)], probably due to hybridization with heavier Ba near square Bi layers. This creates small pockets near $\Gamma$ point in the Brillouin zone. Together with the presence of more 3D-like Zn states not separated away from the Fermi level due to magnetic ordering, this results in the conventional magnetotransport and the absence of Dirac states in BaZnBi$_2$.

In summary, we show that spin-orbit coupling induces gap in Dirac states, causing their removal from the Fermi surface. Resulting electronic transport is governed by cylindrical states away from the $\Gamma$ point and small electron and hole-like pockets near Brillouin zone center. Our results suggest that the mass of the alkaline earth atoms $A$ near Bi$_2$ square network, in addition to magnetic order is essential for formation of Dirac cones in ABX$_2$ materials.

**ACKNOWLEDGEMENTS**

We thank John Warren for help with SEM measurements. This work was supported by the U.S. DOE-BES, Division of Materials Science and Engineering, under Contract No. DE-SC0012704 (BNL) and the National Natural Science Foundation of China under Grant Nos. 51671192 and 51531008 (Shenyang). Work at the National High Magnetic Field Laboratory is supported by the NSF Cooperative Agreement No. DMR-1157490, and by the state of Florida.