Large spin-orbit coupling and helical spin textures in 2D heterostructure [Pb$_2$BiS$_3$][AuTe$_2$]

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Two-dimensional heterostructures with strong spin-orbit coupling have direct relevance to topological quantum materials and potential applications in spin-orbitronics. In this work, we report on novel quantum phenomena in [Pb$_2$BiS$_3$][AuTe$_2$], a new 2D strong spin-orbit coupling heterostructure system. Transport measurements reveal the spin-related carrier scattering is at odds with the Abrikosov-Gorkov model due to strong spin-orbit coupling. This is consistent with our band structure calculations which reveal a large spin-orbit coupling gap of $\varepsilon_{so} = 0.21$ eV. The band structure is also characterized by helical-like spin textures which are mainly induced by strong spin-orbit coupling and the inversion symmetry breaking in the heterostructure system.

The motion of an electron through a lattice potential gives rise to a coupling between the electron’s momentum and its spin. This relativistic effect is known as spin-orbit coupling (SOC). SOC plays an important role in topologically nontrivial electronic structures$^{1-3}$ as well as those which can realize Majorana fermions$^{4-6}$. SOC also offers a unique route to tune the spin degree of freedom that has potential impact on spintronics devices$^{7-9}$. Understanding SOC in confined systems such as interfacial boundaries and nanowires is particularly important in order to exert quantum control on electronic degrees of freedom in miniaturized devices. Extensive studies have been conducted in the 2D electron gas such as GaAs/Ga$_{1-x}$Al$_x$As$^{10}$ and HgTe/CdTe/HgTe quantum wells$^{11}$. More emergent properties can be expected in similar systems with sufficiently strong SOC such as naturally formed crystalline heterostructures that adopt a structure similar to those of epitaxially grown heterostructure films. A strong advantage of naturally formed heterostructures is that a large number of organic and inorganic materials adopt this type of structure, for example the intergrowth chalcogenides$^{12}$ and the inorganic-organic hybrid systems such as halides$^{13}$. These crystal heterostructures provide a parallel platform to epitaxial films to uncover novel quantum phenomena derived from strong SOC and inversion symmetry breaking$^{10,11}$.

Recent work on the naturally formed 2D mineral [Pb$_2$BiS$_3$][AuTe$_2$] shows heterostructure electronic states featuring a conductive [AuTe$_2$] layer sandwiched between two insulating [Pb$_2$BiS$_3$] sheets$^{14}$. The building blocks [Pb$_2$BiS$_3$]$^{-1}$ and [AuTe$_2$]$^{-1}$ are alternately stacked along the crystal’s c-axis to form an intergrowth structure, shown in Fig. 1a$^{14-16}$. A number of intriguing properties such as an extremely large electrical anisotropy, high mobility 2D carriers, and linear energy band dispersions were discovered$^{14}$. The SOC strength in this material is expected to be strong, because many of the component materials are heavy elements. However, as we will discuss below, the SOC in [Pb$_2$BiS$_3$][AuTe$_2$] is anomalous in that it lies beyond the expected scaling properties predicted by the Abrikosov-Gorkov model$^9$. Furthermore, our calculations on atomically thin films unveil helical-like spin textures when the spin vectors traverse around the closed 2D Fermi surfaces. The helical-like spin textures could be an effect arising from the strong SOC and the unique heterostructure in [Pb$_2$BiS$_3$][AuTe$_2$].

Results and Discussion

A direct measurement of a material’s SOC strength is difficult because SOC is a relativistic effect. However, SOC manifests itself in quantum interference phenomena such as weak antilocalization (WAL)$^{17,18}$. WAL provides an effective approach to quantitatively investigate the SOC strength of a material using standard electrical

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characterization. In the diffusion regime and in the case of self-intersecting electron scattering paths, the interference between the electron's time reversed paths is pronounced and tends to localize electrons. This phenomenon is known as weak localization (WL). WL occurs when strong SOC suppresses WL effects.

Figure 1b shows the temperature dependent resistance of a cleaved single-crystal thin layer. The resistance upturn below 150 K demonstrates the poor metallic property of the samples. Below 40 K, the resistance increases linearly with decreasing temperature. This behavior could be related to electron-electron interaction that has been extensively discussed in disordered semiconductors and topological insulators. Our magnetoconductivity measurements focus on the low field regime, for example B < 0.6 T. In this regime, the Hall resistivity (ρ_{xy}) is far smaller than the longitudinal resistivity (ρ_{xx}) and the square conductance tensor takes the form

\[ \rho_{xy} \approx \frac{-\alpha}{\pi} \left[ \Psi \left( \frac{1}{2} \right) + \frac{B_e}{B} \right] - \ln \left( \frac{B_e}{B} \right) \]

where \( \alpha = 1/2 \), e is the electron charge and \( h \) is the Planck constant. The parameters \( \tau_{\omega}, \tau_{\omega}, \) and \( \tau_{\varphi} \) are the electronic elastic scattering time, spin-orbit scattering time and phase coherence time, respectively. The quantity \( \Psi(z) \) is the digamma function. We define a field strength \( B_e = h/(4e^2) \) which is the characteristic field associated with the coherence length \( l_{\varphi} \), where \( l_{\varphi} = \sqrt{D(T)} \) and \( D \) is the diffusion constant. Figure 1c shows a fitting curve which virtually superposes on the experimental data. This fitting confirms that Pb2BiS3AuTe2 has strong SOC.

In Fig. 1d, we compare conductance data as a function of magnetic field of Pb2BiS3AuTe2 with that of Bi2Te3 and BiTe at T = 2 K. For purposes of comparison, this data has been normalized to unity (see supporting
information). The similarities among these various systems, especially between [Pb2BiS3][AuTe2] and Bi2Te3, is surprising. The topological insulator (TI) Bi2Te3 and semi-metal Bi are known for having unusually strong SOC among materials systems. This striking resemblance indicates that aspects of the SOC coupling in [Pb2BiS3][AuTe2] may be unusual. In order to gain a deeper understanding, we quantitatively investigated Bi2Te3 into the unconventional class of systems and the rest such as Au28 into the conventional class. Our [Pb2BiS3][AuTe2] sample falls between these two classes, suggesting the strong SOC, heterostructured systems may have exceptional due to strong SOC.

Figure 2. τe/τso versus effective atomic number (Z) for different systems. Weighted arithmetic mean is used to calculate the effective atomic number of materials composed of multiple elements. For [Pb2BiS3][AuTe2], only the elements Au and Te are accounted for the Z because the density of states at the Fermi surface are mainly contributed by Au 3d electrons and Te p electrons34. The red solid line is a simulation using the relation Z4. TI Bi2Te3 violates the Abrikosov and Gorkov’s prediction τe/τso ~ Z4 due to the topology-related Berry’s phase and the extremely large SOC. [Pb2BiS3][AuTe2] does not follow the Z4 relation. The τso and τe of Al, InP/InGaAs, InO3, Hg0.76Cd0.24Te, Au and Ag are from references28–38.
The large SOC can be understood by our \textit{ab initio} calculations. The electronic structure near the Fermi level of \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\) purely originates from the \([\text{AuTe}_2]\) layer. Figure 3a is a contour plot of the local density of state (LDOS) of \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\) near the Fermi level, which clearly shows that the electronic states are tightly confined to the \([\text{AuTe}_2]\) layer. Figure 3b exhibits the energy band dispersions of the \([\text{AuTe}_2]\) layer and the single-unit-thick films. The vertical arrow marks the gap opening due to strong SOC. Figure 3c shows the detailed information of the gap opening in 3D reciprocal space. Without SOC in the \([\text{AuTe}_2]\) layer, a Dirac-like gapless state appears at the hole-band between the \(\Gamma\) and \(X\) points. SOC opens the gapless Dirac-like state to form a spin-orbit gap of \(\varepsilon_{\text{SO}} = 0.21\) eV. This value is comparable to the SOC-induced band inversion energy in TIs and the spin-orbit gap of the topology-nontrivial Sb-bilayer system\(^{26,29}\). The SOC gap in the \([\text{AuTe}_2]\) layer also occurs at the \(\Gamma\) point near the Fermi Energy (see the top panel of Figure S6 in supporting information). The large spin-orbit gap elucidates the large SOC and the violation of the Abrikosov-Gorkov scattering model. Since \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\) is not a TI, our studies thus suggest that materials can violate the Abrikosov-Gorkov scattering model as long as their SOC strengths are sufficiently large.

The large SOC, combined with the unique heterostructure in \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\), may lead to helical spin textures that have been observed in topological insulators and semiconductor heterostructure films\(^{27,30}\). We carried out theoretical investigations on the single-unit-cell film \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\) films where the twofold spin-degeneracy is lifted due to inversion symmetry breaking. Figure 4a presents the Fermi surfaces of the electron and hole bands and their projections at the Fermi level. The dual projection sheets are induced by broken inversion symmetry. Both electron bands and hole bands are found to have spin structures whose direction changes as a function of crystal momentum \(k\). The spin textures of the electron and hole pockets exhibit different topological properties. For the electron pockets, the spin does not undergo a full winding as one traverses around the Fermi surface while for the hole pockets the spin winds around 3 times. The helical-like spin texture in the hole pocket can be attributed to the strong SOC and the inversion-asymmetry induced spin splitting in a heterostructure system.

\begin{figure}
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\includegraphics[width=\textwidth]{figure3}
\caption{(a) Contour plot of the local density of state (LDOS) of \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\) near the Fermi level. The \([\text{AuTe}_2]\) layer dominates the electronic structure. (b) Energy bands dispersion of one-unit-cell \([\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]\) film (light blue), and \([\text{AuTe}_2]\) single layer (light red). A hole pocket lies at \(\Gamma\) and an electron pocket is formed between \(\Gamma\)-\(Y\). (c) A Dirac-like gapless state appears at the hole-band without SOC (upper panel). SOC opens the Dirac-like gapless state with a spin-orbit gap of 0.21 eV (lower panel). The gap opening is denoted in (b) by the black arrow.}
\end{figure}
For example the Ga$_{1-x}$In$_x$As/InP quantum well has shown helical spin texture in the presence of SOC. The spin textures in the hole pocket exhibit spin-flip between the hole carriers with opposite momentum $k$ and $-k$, as a consequence of time-reversal symmetry. The shape of the Fermi surface reflects the orthorhombic crystalline symmetry of [Pb$_2$BiS$_3$][AuTe$_2$]. In addition, the spin-flip in the hole pocket may indicate a non-vanishing Berry’s phase. Direct detection of the Berry’s phase is beyond the scope of this study. However, the WAL in our sample is consistent with a nontrivial Berry’s phase which gives opposite signs to time-reversed electron paths. This type of destructive quantum interference can also result in WAL. Our modeled helical spin textures in single-unit-cell films could be experimentally pursued as we demonstrate that single-unit-cell films naturally occur on the cleaved surfaces of a bulk crystal (Figure S1 in supporting information). Further experiments such as spin-resolved photoemission spectroscopy are needed to confirm our theoretical predictions of the helical spin textures.

**Conclusions**

To summarize, large SOC strength was discovered in the 2D heterostructure [Pb$_2$BiS$_3$][AuTe$_2$]. This large SOC is induced by a large spin-orbit gap $\varepsilon_{SO} = 0.21$ eV. More broadly, our work suggests that materials can violate the Abrikosov-Gorkov scattering model for sufficiently strong spin-orbit coupling. *Ab initio* calculations reveal helical-like spin textures and spin-flips at the Fermi surfaces. These predictions can be attributed to the effect of strong SOC and inversion symmetry breaking in the heterostructure as well as the constraint of crystalline symmetry in the AuTe$_2$ layer. More generally, our work points out that naturally forming heterostructures made of heavy atoms provide a new direction for exploring novel quantum phenomena at the atomic scale. Given the large number of naturally formed organic and inorganic heterostructures and their hybrids, more discoveries can be expected in this direction.

**Methods**

Bulk crystal synthesis follows a self-flux method by melting stoichiometric compositions of [Pb$_2$BiS$_3$][AuTe$_2$]. Details of the crystal synthesis were reported in our previous paper. Crystals were exfoliated on SiO$_2$/Si wafers using the so-called scotch tape method and were post-annealed at 350°C in argon gas to remove chemical residuals on the cleaved surface. The crystals’ thickness is determined by AFM. Morphology studies on cleaved surfaces were conducted by a MultiMode scanning probe microscope (Vecco) operated in a peak-force tapping mode that has vertical spatial resolution up to 50 pm. Cleaved crystals with thickness 40 nm were selected for transport characterization. Standard photolithography (LaserWirtier, MICROTECH) and magnetron sputtering were employed to pattern contacts on cleaved crystals. In-plane electrical characterization was conducted using the standard four probe method in a liquid $^4$He variable temperature cryostat equipped with a triple-axis vector magnet system (AMI).
Electronic structures of $[\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]$ bulk, one-unit-cell thick $[\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]$ film and the $[\text{AuTe}_2]$ single layer were calculated within density functional theory using Projector Augmented Wave method implemented in the VASP code. Perdew-Burke-Ernzerhof type generalized gradient approximation was utilized for the exchange correlation functional. Spin-orbit coupling was included in the non-collinear form. Experimentally resolved crystal structure was employed for $[\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]$, and crystal structure of $[\text{AuTe}_2]$ single layer was prepared from the bulk crystal structure of $[\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]$ by removing $[\text{Pb}_2\text{BiS}_3]$ layers. The crystal structure of the monolayer $[\text{Pb}_2\text{BiS}_3][\text{AuTe}_2]$ is calculated assuming a vacuum layer of 3 nm.

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Author Contributions
L.F. and M.G.K. synthesized the crystals. L.F., Y.J. and W.K.K. conducted the experiments. J.I. performed first-principles calculations. L.F., J.I., Y.J., W.D., A.G., K.M., M.G.K., W.K.K. and G.W.C. analyzed the data and wrote the manuscript.

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