Inherited weak topological insulator signatures in the topological hourglass semimetal Nb$_3$XTe$_6$ (X = Si, Ge)

Q. Wan,1,* T. Y. Yang,1,* S. Li,2 M. Yang,3,4 Z. Zhu,5 C. L. Wu,1 C. Peng,1 S. K. Mo,1 W. Wu2, Z. H. Chen,6 Y. B. Huang,6 L. L. Lev,7,8 V. N. Strocov,7 J. Hu,9 Z. Q. Mao,10 Hao Zheng,5 J. F. Jia,5 Y. G. Shi,3,4,11 Shengyuan A. Yang2 and N. Xu1,†

1 Institute of Advanced Studies, Wuhan University, Wuhan 430072, China
2 Research Laboratory for Quantum Materials, Singapore University of Technology and Design, Singapore 487372, Singapore
3 Beijing National Laboratory for Condensed Matter Physics and Institute of Physics, Chinese Academy of Sciences, Beijing 100190, China
4 School of Physical Sciences, University of Chinese Academy of Sciences, Beijing 100190, China
5 Key Laboratory of Artificial Structures and Quantum Control (Ministry of Education), Shenyang National Laboratory for Materials Science, School of Physics and Astronomy, Shanghai Jiao Tong University, 800 Dongchuan Road, Shanghai 200240, China
6 Shanghai Synchrotron Radiation Facility, Shanghai Advanced Research Institute, Chinese Academy of Sciences, Shanghai 201204, China
7 Swiss Light Source, Paul Scherrer Institut, CH-5232 Villigen PSI, Switzerland
8 Moscow Institute of Physics and Technology, 9, Institutskiy lane, Dolgoprudny, Moscow region 141701, Russia
9 Department of Physics, University of Arkansas, Fayetteville, Arkansas 72701, USA
10 Department of Physics, Pennsylvania State University, University Park, Pennsylvania 16803, USA
11 Songshan Lake Materials Laboratory, Dongguan, Guangdong 523808, China

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Using spin-resolved and angle-resolved photoemission spectroscopy and first-principles calculations, we have identified bulk band inversion and the spin-polarized surface state evolved from a weak topological insulator (TI) phase in van der Waals materials Nb$_3$XTe$_6$ (X = Si, Ge). The fingerprints of weak TIs homologically emerge with hourglass fermions as multinodal chains composed by the same pair of valence and conduction bands gapped by spin-orbit coupling. The unique topological state, with a pair of valence and conduction bands encoding both weak TI and hourglass semimetal nature, is essential and guaranteed by nonsymmorphic symmetry. It is distinct from TIs studied previously based on band inversions without symmetry protections.

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I. INTRODUCTION

The topological property of wave functions is a key characterization of quantum materials [1–5]. Novel boundary states provide an observable signature of the nontrivial topological invariant of wave functions, and topological phase transitions are accompanied with modifications of boundary states. From the band theory point of view, band inversion [Figs. 1(a) and 1(b)] plays a critical role in the formation of a topological state. In a topological insulator (TI) [6–16], spin-orbit coupling (SOC) opens a bulk hybridization gap between the inverted bands, and Dirac surface states emerge in the gap with spin-momentum locking [Fig. 1(c)]. Benefiting from specific crystalline symmetries, nodes can survive within the bulk hybridization gap [Fig. 1(d)] and define topological semimetals (TSMs) [17–37]. Tuning the hopping term, SOC, on-site, and long-range Coulomb interactions, etc., topological phase transitions can be realized by removing the band inversion without changing symmetry [38–40], as described in Fig. 1(b). However, the wave function simultaneously embedding multiple types of topological phases is elusive and the properties of such state are unknown.

Here, we unveil a unique topological phase hosting both weak TI and hourglass TSM fingerprints in van der Waals (vdW) materials, Nb$_3$XTe$_6$ (X = Si, Ge). It is evolved from SOC gapped multinodal chains composed by the same pair of valence and conduction bands gapped by spin-orbit coupling. The unique topological state, with a pair of valence and conduction bands encoding both weak TI and hourglass semimetal nature, is essential and guaranteed by nonsymmorphic symmetry. It is distinct from TIs studied previously based on band inversions without symmetry protections.

*These authors contributed equally to this work.
†nxu@whu.edu.cn
directions, there are two glide mirror planes perpendicular to the Brillouin zone with the high-symmetry points indicated. (g) Bulk band structure of Nb3SiTe6. The red symbols are extracted band dispersions.

symmetry. Our results not only realize a unique topological phase hosting distinctions of both weak TI and TSM phases by the same group of valence and conduction bands, but also provide a striking vdW material platform for fine-tuning the electronic and topological properties. Few-layer Nb3SiTe6 flakes have been successfully fabricated and enhanced electron coherency was reported [45]. By fine-tuning stoichiometry, directional massless Dirac fermions have been achieved in NbSi0.45Te2 with one-dimensional (1D) long-range order [46].

II. EXPERIMENTAL AND COMPUTATIONAL DETAILS

The Nb3SiTe6 single crystals were synthesized with a mixture of Nb, Si, and Te at a molar ratio of 3:1:6 using chemical vapor transport. During growth of Nb3SiTe6 the temperature was set at 950 °C and 850 °C, respectively, for the hot and cold ends of the double zone tube furnace. Single crystals of Nb3GeTe6 were grown by a solid-state reaction method. A mixture of Nb, Ge and Te at a molar ratio of 3:1:6 was pulverized, pressed into a pellet, placed in an alumina crucible, and then sealed in a highly evacuated quartz tube. After that, the tube was heated to 1100 °C for 10 h and slowly cooled down to 650 °C at a rate of 2 °C/h.

Clean surfaces for ARPES measurements were obtained by cleaving samples in situ in a vacuum greater than 5 × 10−11 Torr. VUV and SX-ARPES experiments were performed at Dreamline of the Shanghai Synchrotron Radiation Facility and SX-ARPES end station of the ADRESS beamline at the Swiss Light Source, Paul Scherrer Institute, Switzerland [47], respectively. Spin-resolved ARPES measurements were done with a home-designed ARPES facility equipped a very low energy electron diffraction (VLEED) spin detector, with a Sherman function of S = 0.27 used to generate the measured spin polarizations.

First-principles calculations were performed within the framework of density functional theory using the Perdew-Burke-Ernzerhof type [48] generalized gradient approximation for the exchange correlation functional as implemented.
in the Vienna *ab initio* simulation package [49–51]. The BZ was sampled with Γ-centered \( k \) mesh of size \( 8 \times 5 \times 4 \), and the cutoff energy was set as 350 eV. The energy and force convergence criteria were set to be \( 10^{-5} \text{eV} \) and 0.01 eV/Å, respectively. The van der Waals (vdW) corrections have been taken into account with the approach of Dion *et al.* [52]. The surface states were calculated by constructing the maximally localized Wannier functions [53,54] and by using the iterative Green’s function method [55,56] as implemented in the WANNIERTOOLS package [57].

III. RESULTS

Figure 1(g) shows the ARPES spectra of Nb₃SiTe₆ acquired with \( hν = 300–570 \) eV. A \( k_z \) dispersive feature confirms the bulk origin of the photoelectron intensity in SX-ARPES measurements, due to the increase of the photoelectron mean free path compared to the VUV energy range [58]. A heavy band (named as \( \alpha \)) and a light band (named as \( \beta \)) are resolved along the Γ–Z direction. Compared to calculations, they are visible in every other BZ due to a photoemission selection rule [59–61] related to nonsymmorphic symmetry \( M_\gamma \) in Nb₃SiTe₆. Similarly, we observe a single branch of Dirac-cone-like dispersion along the Z–S direction in Fig. 1(h) due to \( M_\gamma \). Off of the photoemission scattering plan, the selection rule is not strictly applied and both branches of Dirac-cone-like dispersions are visible along the T–R direction. Our results suggest a quasi-1D Dirac-cone-like dispersion along the S–R direction, consistent with the nodal line along the S–R high-symmetry line (\( NLSR \)) protected by the interplay of \( M_\gamma \), \( T \)–\( M_\gamma \), and \( PT \), as indicated by the calculations without SOC (black line in Fig. 1(h)).

Furthermore, we observe multiple band inversions between the \( \alpha \) and \( \beta \) bands, forming two kinds of nodal chains with finite and infinite length, respectively, if ignoring SOC. Along the Γ–Y direction [Figs. 1(i) and 1(j)], besides the \( \alpha \) and \( \beta \) bands in the first BZ [red marks in Fig. 1(j)], the \( \alpha' \) and \( \beta' \) bands are expected in the second BZ [blue marks in Fig. 1(j)] with zero spectra weight due to selection rule related to \( M_\gamma \). Because both the \( \alpha \) and \( \beta \) bands cross \( E_F \) along the Γ–Y direction, the bottom (top) of the \( \alpha \) (\( \beta \)) band locates below (above) \( E_F \) at the Γ (\( \Gamma' \)) point. Therefore, the \( \alpha \) and \( \beta' \) bands have to overlap with each other at the Γ point as reproduced by first-principles calculations [Fig. 1(j)]. The \( \alpha \) band bottom is \( \sim 100 \) meV below \( E_F \) in the same order of energy resolution of SX-ARPES measurements. Although we cannot clearly resolve the parabolic dispersion within this small energy window, the DFT calculations show an overall agreement with SX-ARPES results [Figs. 1(i) and 1(j)]. The band inversion between the \( \alpha \) and \( \beta' \) bands around the Γ point leads to a nodal ring in the \( k_z = 0 \) plane (labeled as \( NR_\gamma \)), which is protected by \( M_\gamma \) when SOC is not included (as seen from Figs. S1 and S2 in the Supplemental Material [62]).

Along the Γ–X direction, the \( \alpha' \) and \( \beta' \) bands can be resolved in the second BZ; they are quite close to each other near \( E_F \) and well separated at \( E_F > 0.2 \text{eV} \). Similarly, because both the \( \alpha' \) and \( \beta' \) bands cross \( E_F \), we can ascertain that the \( \beta' \) band crosses with the \( \alpha \) band with missing spectra weight near the \( X \) point, forming a nodal chain touching at the \( S \) points in the \( k_y = 0 \) plane (labeled as \( NC_\gamma \)), which is guaranteed by nonsymmorphic symmetries as discussed in detail later.

Figure 2(a) summarizes the nodes composed by the \( \alpha/\alpha' \) and \( \beta/\beta' \) bands when SOC is not included. First-principles calculations further unveil two more nodal rings touching with \( NR_\gamma \) and forming a nodal chain with a finite length (\( NC_\gamma \)). Topological drumhead surface states are shown in the calculated spectra function on the (001) surface [Fig. 2(b)], with the boundary bonding to surface projections of the nodes. The drumhead states are fourfold degenerated (including the spin degeneracy), because each point on the nodal chain projections line corresponds to two nodes with ±\( k_z \) in the three-dimensional (3D) BZ. This is consistent with previous theoretical study on topological nodal chain semimetal IrF₄ [63].

In the presence of SOC, the nodal line and chains are gapped, however, in two distinct ways. SOC gaps \( NLR_\gamma \) as described in Fig. 1(d), with hourglass nodes forming rings \( NR_{HG} \) surrounding the \( S \) points in Fig. 2(c) (more details discussed in Figs. S3 and S4 in the Supplemental Material [62]). For \( NC_\gamma \) and \( NC_\gamma \), SOC fully gaps the nodes and results in a TI scenario as described in Fig. 1(e). Parity calculations indicate \( Z_2 \) indices of \( (0;110) \) in a weak TI configuration with band inversion happening at the \( \Gamma \) and \( U \) points [Fig. 2(c)]. Topological surface states related to the weak TI phase emerge on the (001) surface [Fig. 2(d)]. Compared to the calculation without SOC, the bulk band projections open a gap at the \( \Gamma \) point, and a pair of Dirac cones appears inside the gap with Dirac points close to the bulk valence and conduction bands, respectively [Fig. 2(e)]. We note that \( M_\gamma \) is preserved on the \( \Gamma–X \) path near the (001) surface region; therefore along the \( \Gamma–X \) direction such a dual Dirac-cone structure is an effective hourglass dispersion as imposed by \( M_\gamma \).
The spin-polarized topological surface states is directly observed in our spin-resolved VUV-ARPES measurements on Nb3GeTe6, which shares a similar electronic structure with Nb3SiTe6 (Fig. S5 in the Supplemental Material [62]). In Fig. 3(a), we clearly identify a band near the \( \bar{X} \) point showing no dispersion with photon energy in the range of 24–90 eV, which conforms the two-dimensional (2D) surface-state nature. We plot the Fermi surface (FS) mapped in the surface BZ [Fig. 3(b)]. Figures 3(c) and 3(d) display the surface band structure along the \( \bar{\Gamma} - \bar{X} \) and \( \bar{\Gamma} - \bar{U} \) directions, respectively, in which additional spectra weight appears near \( E_F \) around the \( \bar{X} \) and \( \bar{U} \) points, in good agreement with the topological surface state indicated by calculations in Fig. 2(d).

We performed spin-resolved ARPES measurements to study the spin texture of the surface states. We note that the upper branch of the lower Dirac cone (DC\(_L\)) and lower branch of the upper Dirac cone (DC\(_U\)) are quite close to each other near the \( \bar{X} \) point [Fig. 2(d)] and the spin signals would be mixed by limited experimental resolution. To avoid this complexity, we focus on the surface states along the \( \bar{\Gamma} - \bar{X} \) direction in which only single Dirac cone dispersion has nonzero spectra weight in an ARPES experiment, due to the selection rule related to \( \hat{M}_y \). Figures 3(e) and 3(f) show the spin-resolved energy distribution curve (EDC) intensity \( I_{\gamma\gamma}^{1\pm} \) in the \( x \) and \( y \) directions, respectively, measured at the \( E \) point labeled in Figs. 3(b) and 3(c). While \( I_{\gamma}^1 \) is almost equal to \( I_{\gamma}^1 \) [Fig. 3(e)], there is a clear difference in \( I_{\gamma}^1 \) and \( I_{\gamma}^1 \) at the EDC’s peak that corresponds to surface states [Fig. 3(f)], indicating that the observed surface state is spin polarized along the \( y \) direction. At the time-reversal symmetric \( F \) point labeled in Figs. 3(b) and 3(c), the surface state is spin polarized along the \( y \) direction [Fig. 3(g)], however, in the opposite direction to that at the \( E \) point. Our spin-resolved VUV ARPES clearly reveals the topological surface states with spin-momentum locking as evidence of inherited weak TI phase in Nb3XTe6.

The bulk bands of Nb3XTe6 are spin degenerated as required by time-reversal and inversion symmetries. The “hidden” spin signals from the bulk projection bands, as observed in systems with both time-reversal and inversion symmetries [64,65], are forbidden along the \( \bar{\Gamma} - \bar{X} \) direction by symmetries of Nb3XTe6 monolayer (Fig. S6 in the Supplemental Material [62]). Therefore, spin polarization observed in Figs. 3(e)–3(g) is from topological surface states, without contribution from bulk band projections.

**IV. DISCUSSIONS**

Figure 4 schematically summarizes our main finding of the electronic structure and topological properties of Nb3XTe6. From SX-ARPES results in Fig. 1, we directly observe the bulk \( \alpha \) and \( \beta \) bands overlap with each other near \( E_F \) and form nodal line NL\(_{\text{SR}}\) and nodal chains NC\(_x\)/NC\(_y\) if ignoring SOC [Fig. 4(a)]. A topological drumhead state emerges on the (001) surface, bonding to the surface projections of nodes [Fig. 4(b)]. In contrast to the nodal line/chain induced by band overlap without symmetry protection as depicted in Figs. 1(a) and 1(b), NC\(_x\) is essential and based on a band inversion [Fig. 1(k)] guaranteed by nonsymmorphic symmetries, as illustrated by Figs. 4(c) and 4(d). Because the \( k_y = 0 \) plane, which hosts the nodal chain NC\(_x\) [yellow plane in Figs. 4(a) and (c)], is the invariant plane of \( \hat{M}_y \), for each band \( |u \) we have

\[
\hat{M}_y |u\rangle = g_{\tilde{M}_y} |u\rangle.
\]

Because of nonprimitive translation operations, \( \hat{M}_y \) takes eigenvalues of

\[
g_{\tilde{M}_y} = \pm e^{-\frac{2\pi i}{\tilde{Q}}},
\]

when SOC is not included. We would consider band structure along the path of \( K - \bar{\Gamma} - Q \) in Fig. 4(c), where \( K = (\pi, 0, k_z) \) and \( Q = (k_x, 0, \pi) \) are arbitrary points sitting on the \( S - S \) and \( S - Z \) high-symmetry lines, respectively. The solid and dashed lines in Fig. 4(d) distinguish the additional phase term \((-1)\) in \( g_{\tilde{M}_y} \). Because the \( K \) point on the \( S - S \) high-symmetry line is invariant under \( \hat{M}_y \), and the commutation relationship between \( \hat{M}_x \) and \( \hat{M}_y \) is

\[
\hat{M}_y \hat{M}_x = T_{110} \hat{M}_x \hat{M}_y,
\]

where \( T_{110}(x, y, z) = (x + a, y - b, z) \),

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the bands with additional phase term have to degenerate at the $K$ point with the opposite eigenvalues of $\pm i$ in Fig. 4(d):

$$\tilde{M}_x\tilde{M}_y|u_k\rangle_K = e^{-i k_x a} M_x M_y |u_k\rangle_K = -g_{\tilde{M}_x\tilde{M}_y}|u_k\rangle_K.$$ 

Because there are two layers in the unit cell of Nb$_3$XTe$_6$ [Fig. 1(e)] and the interlayer coupling strength is considerable [Fig. 1(g)], the interlayer splitting has to be taken into account in Fig. 4(d). The double degeneracy on the $S$-$Z$ high-symmetry line is guaranteed by $TC_{2z}$ symmetry, where $C_{2z}$ is a screw rotation along the $z$ direction; i.e.,

$$\tilde{C}_{2z}(x, y, z) \rightarrow (-x, -y, z + 1/2c).$$

At the $k_z = \pi$ plane which includes the $S$-$Z$ path and is invariant under $TC_{2z}$, $(T\tilde{C}_{2z})^2 = -1$ ensures each band to be Kramers doubly degenerate. Also, by noting

$$\tilde{M}_x(T\tilde{C}_{2z}) = T_{10}(T\tilde{C}_{2z})\tilde{M}_x = e^{-i(k_x a + k_y b)(T\tilde{C}_{2z})}\tilde{M}_x = e^{-i(k_x a)}(T\tilde{C}_{2z})\tilde{M}_x,$$

along the $S$-$Z$ line, each Kramers pair at the $Q$ point must share the same $\tilde{M}_x$ eigenvalue, as imposed

$$\tilde{M}_x(T\tilde{C}_{2z})|u_Q\rangle = e^{-i(k_x a)}(T\tilde{C}_{2z})\tilde{M}_x|u_Q\rangle = g_{\tilde{M}_x}|u_Q\rangle,$$

where $g_{\tilde{M}_x} = e^{-i k_x a}$ or $-e^{-i k_x a}$.

Thus, the band dispersion along the $K$-$\Gamma$-$Q$ path effectively forms an hourglass dispersion, with the hourglass node with opposite $g_{\tilde{M}_x}$ values guaranteed by nonsymmorphic symmetries. It is different from band inversion without symmetry protection as described in Fig. 1(b). By tuning hopping parameters, the band inversion along the $\Gamma$-$K$ direction can be lifted and the hourglass nodes will be shifted to the $\Gamma$-$Q$ direction but cannot be eliminated without breaking the nonsymmorphic symmetries (Fig. S7 in the Supplemental Material [62]).

The considerable strength of SOC in Nb$_3$XTe$_6$ eventually gaps the nodal line and chains in Fig. 4(a) in two ways. As protected by $\tilde{M}_x$, SOC turns NL$_{\text{SR}}$ into an hourglass nodal ring NR$_{\text{HG}}$ [Fig. 4(e)]. In contrast, SOC fully gaps $N_C$ and $N_C$ and results in a weak TI phase with $Z_2 = \{0; 110\}$. Topological drumhead surface states evolve into a pair of spin-split Dirac cones at the $\bar{Q}$ point in Fig. 4(f). Such a dual Dirac-cone structure is effectively an hourglass dispersion along the $\bar{Q}$-$\bar{X}$ direction, because the $\tilde{M}_x$ is preserved on the $\bar{Q}$-$\bar{X}$ path. The upper branches of the two Dirac cones have to be degenerated at the $\bar{X}$ point as required by time-reversal symmetry, through the bulk conduction bands. Similarly, the lower branches of the Dirac cones have to be degenerated at the $\bar{X}$ point through the bulk valence bands. An hourglass node is formed and guaranteed by $\tilde{M}_x$ [inset of Fig. 4(f)]. The spin-polarized topological surface state, as the signature of a weak TI phase, is observed in spin-resolved VUV-ARPES measurements in Fig. 3, coexisting with bulk hourglass fermions.

Therefore, we uncover a unique topological phase in Nb$_3$XTe$_6$, which shows a full set of fingerprints of weak TI and hourglass TSM phases near $E_F$, derived from the same group of valence and conduction bands in a single material. The inherited TI signatures are stable and guaranteed by nonsymmorphic symmetries, in contrast to that induced by band inversion without symmetry protection. The unique behaviors of the topological hourglass surface state on the (001) surface with a narrow bandwidth inherited from the weak TI phase and the interactions with homological bulk hourglass fermions call for further research activities. Considering the compositional-tunable nature of the layered vDW Nb$_3$X$_2$Te$_6$ family [66] and 1D massless Dirac fermions observed in NbSi$_{0.45}$Te$_2$ [46], our results also unveil a promising platform for tuning the hourglass fermions and topological phase transition with multiple parameters such as thickness and $X$ stoichiometry.

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