

# Dirac Nodal Line in Hourglass Semimetal $\text{Nb}_3\text{SiTe}_6$

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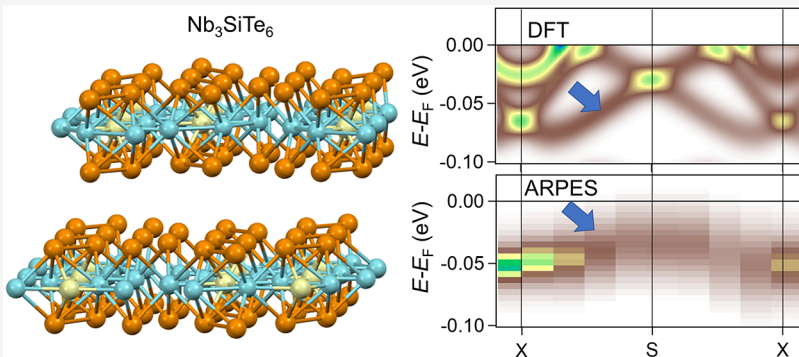
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**ABSTRACT:** Glide-mirror symmetry in nonsymmorphic crystals can foster the emergence of novel hourglass nodal loop states. Here, we present spectroscopic signatures from angle-resolved photoemission of a predicted topological hourglass semimetal phase in  $\text{Nb}_3\text{SiTe}_6$ . Linear band crossings are observed at the zone boundary of  $\text{Nb}_3\text{SiTe}_6$ , which could be the origin of the nontrivial Berry phase and are consistent with a predicted glide quantum spin Hall effect; such linear band crossings connect to form a nodal loop. Furthermore, the saddle-like Fermi surface of  $\text{Nb}_3\text{SiTe}_6$  observed in our results helps unveil linear band crossings that could be missed. *In situ* alkali-metal doping of  $\text{Nb}_3\text{SiTe}_6$  also facilitated the observation of other band crossings and parabolic bands at the zone center correlated with accidental nodal loop states. Overall, our results complete the system's band structure, help explain prior Hall measurements, and suggest the existence of a nodal loop at the zone center of  $\text{Nb}_3\text{SiTe}_6$ .

**KEYWORDS:** hourglass fermions, Dirac nodal loop, nonsymmorphic crystals, glide-mirror symmetry, photoemission, first-principles band structures

Topology in condensed-matter systems has emerged as a fundamental avenue for realizing exotic symmetry-protected Dirac fermions and emergent phases with unconventional but technologically relevant electronic properties, including topological insulators, topological crystalline insulators, and topological superconductors.<sup>1–6</sup> From initial theoretical discussions regarding the intimate connection between massless Dirac surface states and their topological protection due to time-reversal symmetry, particle-hole symmetry, etc.,<sup>7</sup> the theoretical and experimental focuses have now blossomed into considerations of unusual—albeit versatile—fermion states that arise due to certain crystal point group symmetries,<sup>5,6</sup> even those that are protected by a nonsymmorphic space group.<sup>8,9</sup> Indeed, in the presence of certain crystalline symmetries, the Dirac fermion states may cross to form a Dirac point or generate a manifold Dirac nodal point, a Dirac nodal curve, or a Dirac nodal loop in  $k$ -space,<sup>6,10–12</sup> and electrons' behaviors near a manifold Dirac point have been classified, for example, as Weyl fermion,<sup>6,10</sup> hourglass fermion,<sup>11</sup> and wallpaper fermion.<sup>12</sup> However,

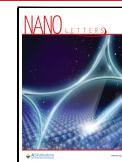
though numerous compounds have been predicted to host topological nodal lines or nodal loops,<sup>13</sup> recent experimental works via angle-resolved photoemission spectroscopy (ARPES) of these topological quasiparticle band dispersions have mainly focused on just a few isostructural families, such as transition metal monopnictides,<sup>14–16</sup> ZrSiS and others in the PbFCl-type family,<sup>17–22</sup> and PbTaSe<sub>2</sub>,<sup>13</sup> rendering the physical picture of most topological nodal line semimetals largely incomplete.

Particularly, hourglass fermions are among the newest types of nodal fermions expected to emerge in topological nodal line semimetals possessing glide-mirror symmetry.<sup>11,23</sup> In such

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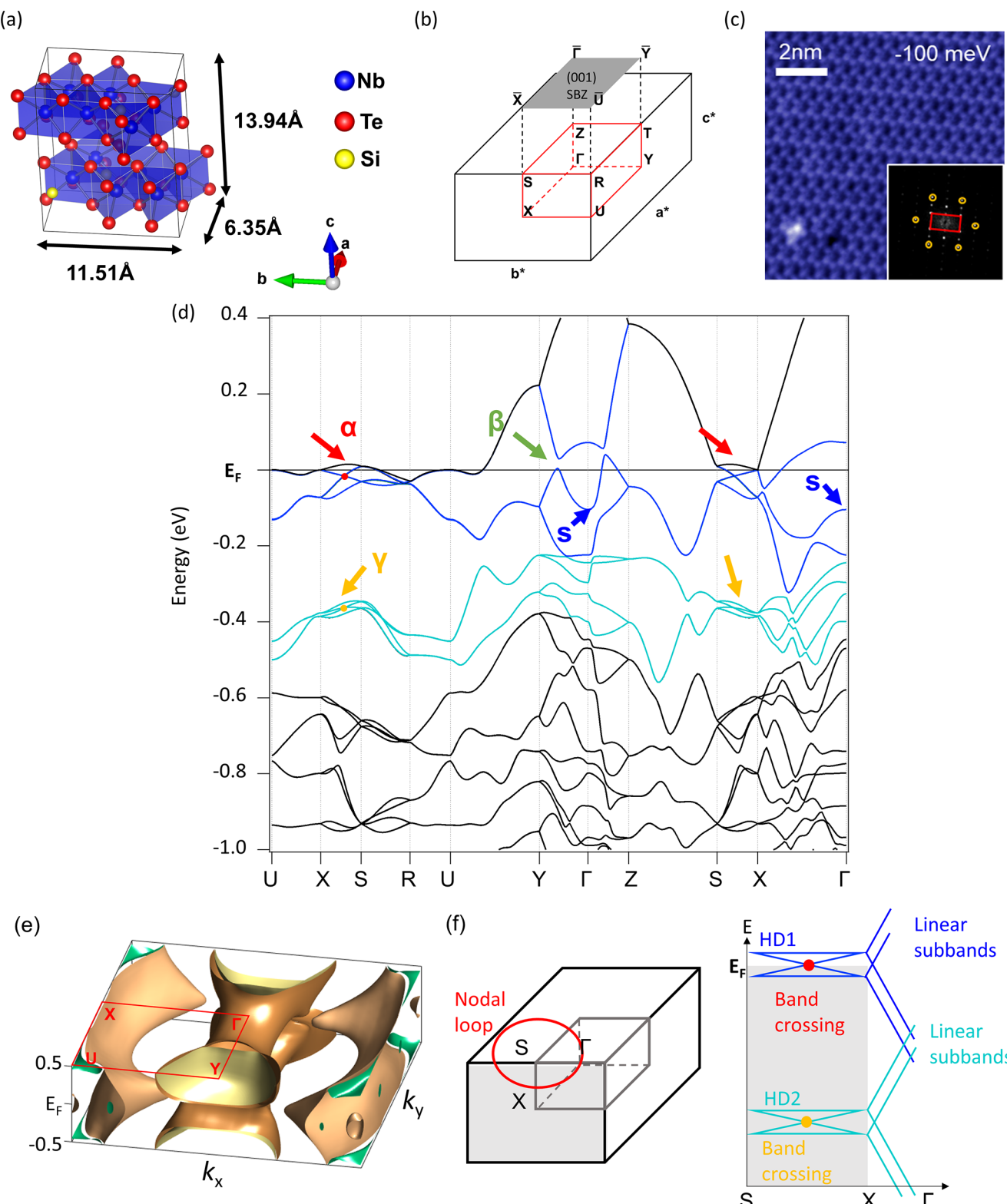
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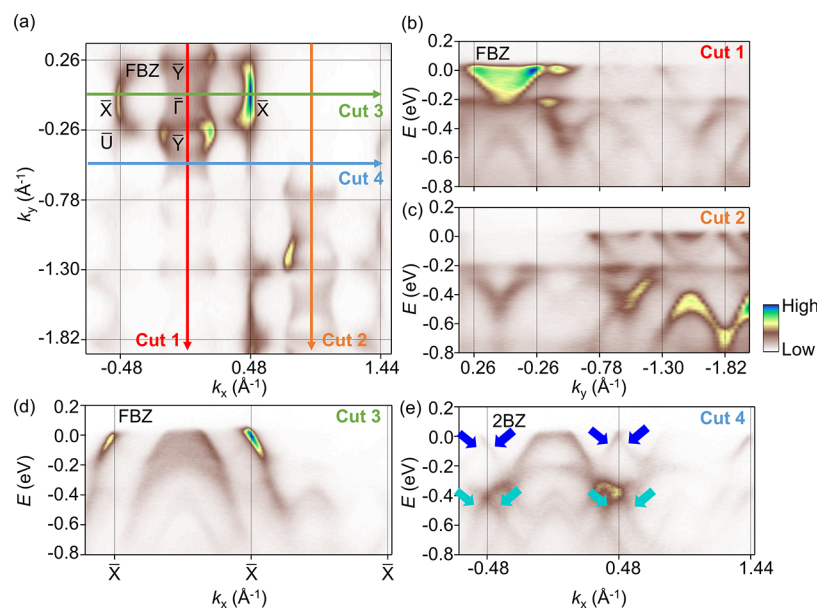


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**Figure 1. Atomic lattice structure and characterizing hourglass bands in  $\text{Nb}_3\text{SiTe}_6$ .** (a) Atomic structure of  $\text{Nb}_3\text{SiTe}_6$ . (b) Bulk BZ and (001)-projected surface BZ (gray rectangle). (c) Topographic image of an *in situ* cleaved  $\text{Nb}_3\text{SiTe}_6$  surface ( $V_s = -100 \text{ mV}$ ,  $I_t = 1500 \text{ pA}$ ,  $10 \times 10 \text{ nm}^2$ ,  $T = 4.2 \text{ K}$ ). The inset is the corresponding fast Fourier transform (FFT) of the topographic image. In the inset, the red rectangle marks the [001]-projected SBZ, while yellow circles indicate Bragg peaks from Te atoms in the topmost layer. (d) DFT band structure of  $\text{Nb}_3\text{SiTe}_6$ . Here, the label  $\alpha$  and red arrows identify hourglass band dispersions along the XS  $k$ -path near the Fermi level, while hourglass bands gapped by SOC along YTZ close to the Fermi level are marked by the label  $\beta$  and a green arrow; the label  $\gamma$  and yellow arrows indicate hourglass bands at deeper binding energies, and the label  $s$  and blue arrows flag bands that form a saddle-shaped surface centered about the  $\Gamma$  point in energy-momentum space. (e) Panoramic view of the constant energy contours of  $\text{Nb}_3\text{SiTe}_6$  in the first BZ (FBZ). (f) Schematic diagrams for the nodal loop in the XSRU plane (left, red oval) and band structures of the hourglass dispersion 1 (HD1) and hourglass dispersion 2 (HD2) along SX $\Gamma$  (right) that give rise to the nodal loop. The nodal loop is formed by the band crossing points (red and yellow points in the right panel).



**Figure 2.** Fermi surface and band mappings of  $\text{Nb}_3\text{SiTe}_6$ . (a) Fermi surface map of  $\text{Nb}_3\text{SiTe}_6$ . Here, the result is obtained by integrating the ARPES intensity within  $\pm 10$  meV of the Fermi level. Solid black lines correspond to BZ boundaries, where the lengths of  $\overline{\Gamma}\overline{X}$  and  $\overline{\Gamma}\overline{Y}$  are  $0.48$  and  $0.26 \text{ \AA}^{-1}$ , respectively; the first BZ is at the top left corner (labeled FBZ), while strategically chosen momentum cuts used for obtaining ARPES band mappings are indicated by colored arrows. (b,c) Measured band maps along cuts 1 and 2 in the Fermi surface map of (a), respectively [red and orange arrows in (a)]; both momentum cuts are parallel to  $\overline{\Gamma}\overline{Y}$ . (d,e) Experimental ARPES spectra along cuts 3 and 4 in the Fermi surface map of (a), respectively [green and blue arrows in (a)]; both momentum cuts are parallel to  $\overline{\Gamma}\overline{X}$ , and the fat blue and aqua arrows in (e) identify two sets of Dirac-like, linear band crossings. In all mappings here, the incident photon energy is  $60 \text{ eV}$ .

materials, the partner switching of eigenstates between certain time-reversal invariant momenta (TRIM), as required by mirror and glide-mirror symmetries, causes so-called “hourglass-like” band dispersions to arise with a fourfold-degenerate nodal point, which connects with other nodal points to form a closed and topologically protected Dirac loop in  $k$ -space.<sup>23</sup> Prior experimental results have confirmed the existence of hourglass fermions in  $\text{KHgSb}$ ,<sup>24</sup> but their presence has also been predicted in the orthorhombic family of  $\text{X}_3\text{SiTe}_6$  ( $\text{X} = \text{Ta}, \text{Nb}$ ) crystals,<sup>23,25</sup> wherein the hourglass bands are robust under the spin–orbit interaction.<sup>23</sup> Nevertheless, while prior ARPES studies have focused on probing one-dimensional massless Dirac fermions in the related  $\text{NbSi}_{0.45}\text{Te}_2$  compound<sup>26</sup> as well as nodal lines and their relation to hourglass fermions in  $\text{Ta}_3\text{SiTe}_6$ ,<sup>27</sup> a comprehensive survey of the electronic band structure of  $\text{Nb}_3\text{SiTe}_6$ —one that considers hourglass bands that disperse with respect to the momentum  $k_z$  along the direction normal to the crystal’s surface—is still lacking.

In this work, by combining density functional theory (DFT) calculations together with high-resolution ARPES, we have carefully performed a systematic study of the electronic band structure of the hourglass semimetal candidate  $\text{Nb}_3\text{SiTe}_6$ . By conducting mappings with respect to the momentum  $k_z$  parallel to the crystal’s surface normal, we have observed a downward dispersing band near the Fermi level that is the lower part of hourglass fermion bands, as inferred from our DFT calculations. The observed band shapes strongly support  $\text{Nb}_3\text{SiTe}_6$  as an hourglass semimetal candidate. Moreover, our careful  $k_z$  band mappings indicate that the nodal line associated with these hourglass dispersions is periodic every two Brillouin zones (BZs), as verified by our DFT calculations of the spectral function. Also, a saddle-like Fermi surface has been mapped, which supports prior evidence from transport<sup>28–31</sup> for the dominance of hole carriers in the system’s electronic

properties. Lastly, *in situ* K doping of  $\text{Nb}_3\text{SiTe}_6$  during ARPES facilitated the examination of other band crossings and parabolic bands at the zone center that are associated with nodal loop dispersions. Our results not only provide compelling indications of hourglass fermions in  $\text{Nb}_3\text{SiTe}_6$  but also suggest that these hourglass fermions must figure prominently in the physics of  $\text{Nb}_3\text{SiTe}_6$  by virtue of their proximity to the system’s Fermi level, rendering  $\text{Nb}_3\text{SiTe}_6$  an intriguing candidate for topological device applications.

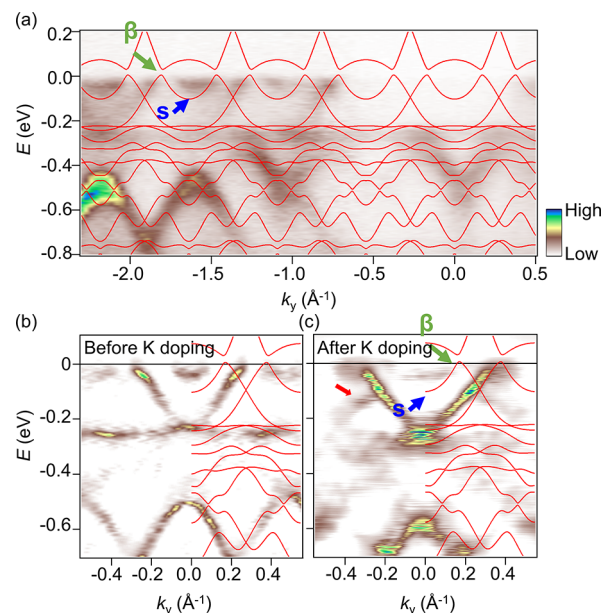
As shown in Figure 1(a), the orthorhombic unit cell for  $\text{Nb}_3\text{SiTe}_6$  consists of two van der Waals-bonded layers, each of which contains a Nb–Si atomic sheet sandwiched in between two puckered-hexagonal atomic layers of Te. The stacking between the unit cell’s component layers is of the AB-type. That is, the four Te layers are in an AABB stacking pattern along the (001) direction of the unit cell, suggestive of the system’s glide-mirror symmetry. The bulk lattice is assembled by vertically stacking many unit cells on top of one another with van der Waals bonding between layers. For reference, Figure 1(b) displays the orthorhombic bulk BZ along with its (001)-projected surface BZ with key high-symmetry points indicated. Topographic images of *in situ* cleaved  $\text{Nb}_3\text{SiTe}_6$  single crystals, as obtained from scanning tunneling microscopy (STM), confirmed both the atomic structure and high crystallinity of the probed surfaces [Figure 1(c)]. First-principles calculations of the bulk band structure for  $\text{Nb}_3\text{SiTe}_6$  [see Figure 1(d)] yield two hourglass-like band dispersions near  $E = 0 \text{ eV}$  and  $E = -0.4 \text{ eV}$  [marked as  $\alpha$  and  $\gamma$ , respectively, in Figure 1(d)], which correspond to two pairs of hourglass fermions. Near the zone center, another set of bands with a nodal loop dispersion is evident along  $\overline{\Gamma}\overline{Y}$  in the theoretical band structure [identified as  $\beta$  in Figure 1(d)], whose lower half converges at the zone center to form a saddle-like surface in energy-momentum space [labeled as  $s$  in Figure

1(d)]; the gap opening in these hourglass bands is due to glide-mirror symmetry breaking. The Fermi surface of  $\text{Nb}_3\text{SiTe}_6$  itself is generally saddle-shaped with a local minimum along  $\Gamma\text{Y}$  and a local maximum along  $\Gamma\text{X}$  [Figure 1(e)], and the hourglass fermion band crossings along  $\text{XS}$  [identified as  $\alpha$  and  $\gamma$  in Figure 1(d)] connect to form a Dirac nodal loop in  $k$ -space in the XSRU plane, centered about the zone boundary point S [see Figure 1(f)].

Representative ARPES results for the valence band structure of  $\text{Nb}_3\text{SiTe}_6$  are summarized in Figure 2. In Figure 2(a), the Fermi surface contour map, taken over multiple BZs, displays a dumbbell-shaped contour centered about the zone center and with its long axis oriented along  $\overline{\Gamma}\text{Y}$ , while another open contour centered about the  $\bar{X}$  point arches along the zone boundary; the latter contour should be associated with a topological nodal line, as in  $\text{Ta}_3\text{SiTe}_6$ .<sup>27</sup> Clearly, though the intensities of Fermi surface contours decrease markedly outside of the first BZ (FBZ) and are likely modulated by complex matrix element effects and photon-energy-dependent ( $k_z$ -dependent) cross sections,<sup>32</sup> such contours are nevertheless sharp and display a clear periodic pattern in  $k$  space over multiple BZs, indicative of the high single crystallinity of the measured  $\text{Nb}_3\text{SiTe}_6$  sample [Figure 2(a)], as confirmed by our X-ray diffraction results (see Supporting Information). Particularly, the dumbbell-shaped contour in the FBZ continuously connects with that of the next BZ along  $\overline{\Gamma}\text{Y}$ .

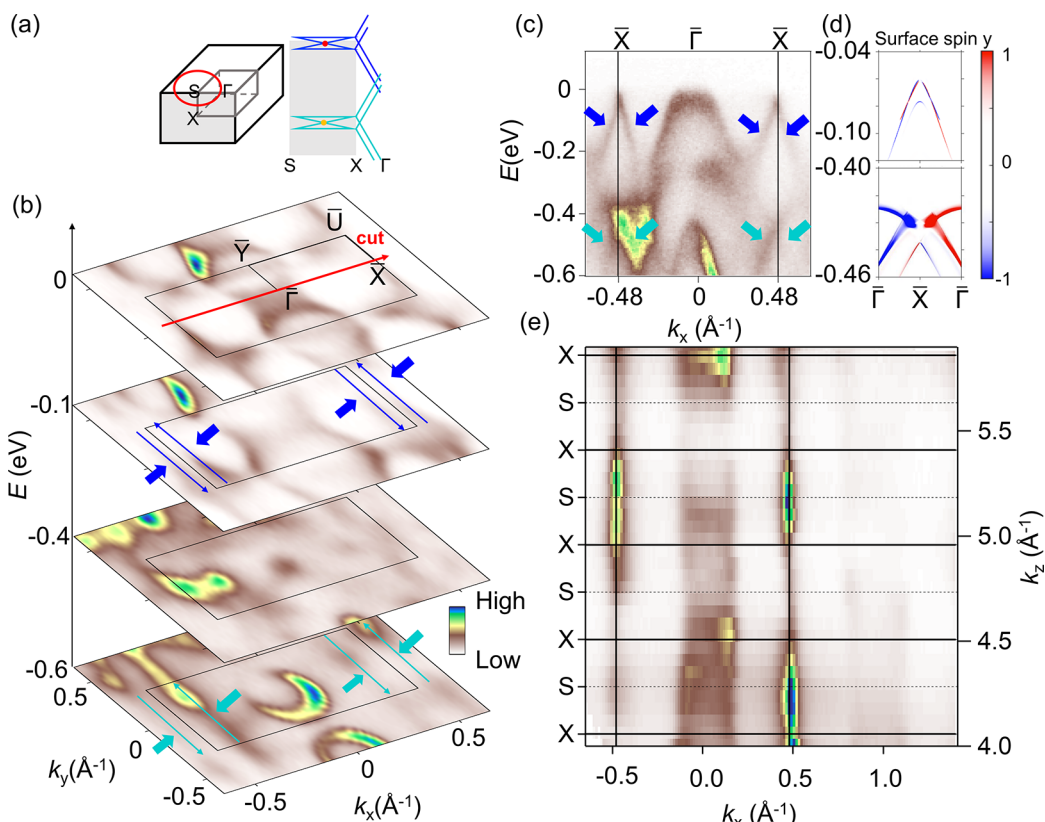
Select momentum cuts through these dumbbell-shaped Fermi surface contours [Figure 2(a)] at strategically chosen  $k_x$  and  $k_y$ , where  $k_x$  and  $k_y$  are momentum components parallel to  $\overline{\Gamma}\text{X}$  and  $\overline{\Gamma}\text{Y}$ , respectively, provide further support for both the periodicity and anisotropy of the system's electronic band structure while illustrating rich photoemission matrix element effects in the measured band dispersions [Figures 2(b–e) and 3(a)]. Indeed, though ARPES intensities of states near the Fermi level in the FBZ are too intense for resolving band features [cuts 1 and 3, e.g.,  $k_x = k_y = 0 \text{ \AA}^{-1}$ , in Figure 2(b,d)],  $k$ -space cuts taken away from the sample's normal in other Brillouin zones [cuts 2 and 4, e.g.,  $k_x = 0.96 \text{ \AA}^{-1}$  and  $k_y = -0.52 \text{ \AA}^{-1}$ , in Figure 2(c,e)] are highly illuminating. For cut 2 parallel to  $\overline{\Gamma}\text{Y}$  [Figure 2(c)], two parabolic bands opening upward—a small one with its minimum at  $E \approx -0.02 \text{ eV}$  and another spanning the energy range  $E = 0$  to  $-0.2 \text{ eV}$ —are repeatedly observable at the zone centers of multiple Brillouin zones [see Figure 2(c) from  $k_y = -0.78$  to  $-1.82 \text{ \AA}^{-1}$ ]; however, for cut 4 parallel to  $\overline{\Gamma}\text{X}$  [Figure 2(e)], a large, hole-like band centered about the zone center and with its apex at  $E \approx -0.02 \text{ eV}$  is instead discernible. These two results thus reveal a saddle-like surface in energy-momentum space, with its saddle point at the zone center at  $E \approx -0.02 \text{ eV}$ , as well as the dominance of hole-like carriers to the crystal's electronic properties, consistent with the theoretical band structure [refer to bands marked by blue arrows in Figure 1(d)] and prior observations from transport and the system's thermoelectric properties.<sup>28–31</sup> Nevertheless, the predicted energy position of the saddle point is deeper in energy (at  $E \approx -0.1 \text{ eV}$ ) than that observed in these measurements (at  $E \approx -0.02 \text{ eV}$ ), likely attributable to slight doping of the  $\text{Nb}_3\text{SiTe}_6$  crystal during its synthesis.

Furthermore, the observation of this saddle-like surface is inherently tied to potential signatures of hourglass dispersions gapped by spin-orbit coupling (SOC) near the zone center of  $\text{Nb}_3\text{SiTe}_6$ ,<sup>23</sup> per our theoretical and experimental band structure comparisons [Figures 1(d) and 3(a–c)]. Like Figure



**Figure 3. Candidate nodal loop bands before and after alkali-metal doping along  $\overline{\Gamma}\text{Y}$ .** (a) Band structure along cut 2 in Figure 2(a) obtained using 60 eV photons; the mapping is overlaid with DFT results (red solid lines) and a nodal loop band structure (green arrow and label  $\beta$ ) and the saddle-shaped surface (blue arrow and label  $s$ ) are identified. (b,c) 2D curvature results for band mappings along  $\overline{\Gamma}\text{Y}$  before and after K doping, respectively, each taken using an incident photon energy of 100 eV and superimposed with DFT band structures (red solid lines). In (c), a band whose intensity is enhanced after K doping is identified (red arrow), and bands corresponding to the saddle-shape surface (blue arrow and label  $s$ ) and the hourglass bands gapped by SOC (green arrow and label  $\beta$ ) are indicated.

2(c) but now overlaid with theoretical band structures, Figure 3(a) displays the band mapping for cut 2 indicated in the Fermi surface map of Figure 2(a) (with  $k_x = 0.96 \text{ \AA}^{-1}$ ), which is the cleanest cut parallel to  $\overline{\Gamma}\text{Y}$  for observing and confirming the periodicity of one of the saddle-shaped surface's bands over multiple BZs [blue arrow in Figure 3(a)]. Figure 3(b,c) summarizes two-dimensional (2D) curvature results for ARPES spectra measured along  $\overline{\Gamma}\text{Y}$  in the FBZ before and after *in situ* alkali doping, each superimposed with the associated DFT calculations of pure  $\text{Nb}_3\text{SiTe}_6$ .<sup>33</sup> The results before and after alkali-metal doping are quite clear [compare Figures 3(b) and 3(c)]: aside from effectively shifting all bands to deeper binding energies, alkali-metal doping of the  $\text{Nb}_3\text{SiTe}_6$  surface facilitates further identification of band features likely related to nodal loop bands [green arrow in each of Figure 3(a,c)], namely, one giving rise to a band crossing at the  $\bar{Y}$  point [red arrow in Figure 3(c)] and also the upward opening parabolic band of the saddle-like surface [blue arrow in each of Figure 3(a,c)]. The photoemission intensity of the former is largely suppressed in the pristine case [Figure 3(b)] but is visible in the alkali-doped system [red arrow in Figure 3(c)], likely due to complex photoelectron diffraction effects and cross section variations of the probed bands. Overall, though the complete band dispersions of the hourglass fermions gapped by SOC are inaccessible through photoemission based upon our theoretical and experimental analysis, there is nevertheless evidence for nodal loop band dispersions located very close to the Fermi level [Figure 3(a–c)],<sup>23</sup> which



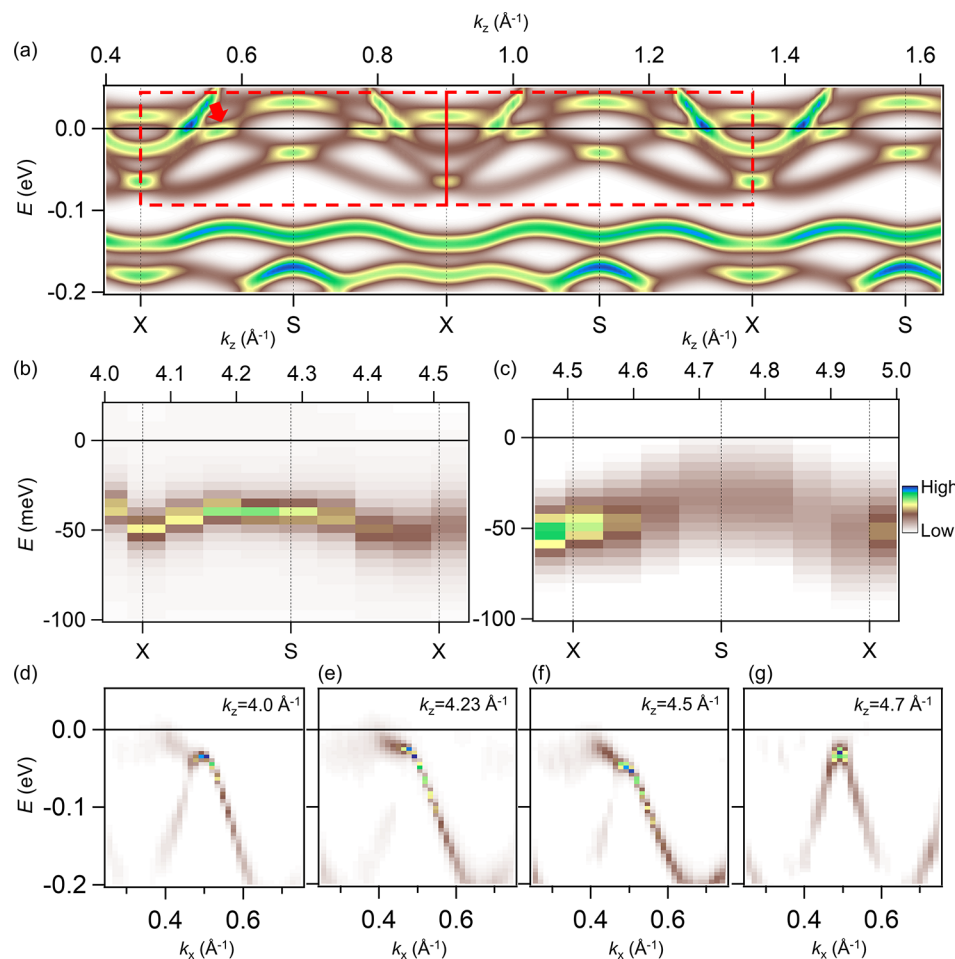
**Figure 4. Constant energy contour maps and potential double QSH state in  $\text{Nb}_3\text{SiTe}_6$ .** (a) BZ and the schematic band structures (blue and aqua lines) of hourglass fermions and their partner linear subbands along the  $\text{SX}\Gamma$   $k$ -path. (b,c) Constant energy contour maps in the FBZ and band mapping along  $\overline{\Gamma}\text{X}$ , respectively, all taken using 70 eV photons. The integrated energy range in each map is 25 meV; in (b,c), aqua and blue arrows indicate two sets of linear bands near each of the  $\overline{\text{X}}$  points, while in (b), the red solid arrow indicates a momentum cut corresponding to the ARPES spectra in (c). (d) Spin-resolved DFT band structures for the two linear band crossings at the  $\overline{\text{X}}$  point with spin-polarization projection taken along the  $k_y$ -direction or  $\overline{\Gamma}\text{Y}$ . (e) Fermi surface mapping in the  $k_x$ – $k_z$  plane obtained by integrating the ARPES intensity within  $\pm 25$  meV of the Fermi level, where  $k_x$  and  $k_z$  are momentum components parallel to the  $\overline{\Gamma}\text{X}$  and [001] directions, respectively; BZ boundaries are labeled along the left axis and/or identified by black lines.

have yet to be considered in the physics of hourglass semimetals and related systems.<sup>24,26,27</sup>

Besides candidate gapped hourglass bands close to the zone center, two other species of hourglass subbands with linear, Dirac-like dispersions near the BZ boundaries also appear in our ARPES data—one set very close to the Fermi level, the other buried at deeper binding energies from  $E = -0.4$  to  $-0.6$  eV [Figures 1(d), 2(e), and 4(a–c)]. A schematic diagram showing the relationships of these linear dispersions (along  $\Gamma\text{X}$ ) to their partner hourglass band crossings (on the  $\text{XS}$   $k$ -path) is displayed in Figure 4(a). Evidently, like the presumed nodal loop bands of the saddle-like surface, the ARPES intensities of these linear bands exhibit marked photoemission matrix element effects, as captured by their modulations across the BZ in the constant energy contour maps [Figure 2(b)] and left–right intensity asymmetry in the band mappings [Figure 2(d,e) and 4(c)]. All linear dispersions here are spin-polarized [schematically indicated by thin aqua and blue arrows in Figure 4(b)], as shown in the related hourglass fermion system KHgSb<sup>11,24</sup> and further unveiled in our theoretical calculations of each linear band’s spin texture (with spin projection taken along the  $k_y$ -direction or parallel to  $\overline{\Gamma}\text{Y}$ ) [Figure 4(d)]. (See also Section 5 in the Supporting Information.) Specifically, for each Dirac-like point, the two partner bands possess opposite spin polarizations relative to one another, which all taken

together contribute no net current flow over the BZ.<sup>11</sup> Consequently, such hourglass systems have been dubbed glide-symmetric analogues of the quantum spin Hall (QSH) state—or more aptly, so-called double QSH systems per the two species of linear band crossings [Figure 4(b,c)].<sup>11,24,34</sup> Thus far, experimental signatures of a double QSH state have only appeared in transport measurements of KHgSb,<sup>34</sup> but clearly, in the case of  $\text{Nb}_3\text{SiTe}_6$ , our theoretical calculations of spin textures around the  $\overline{\text{X}}$  point now also suggest that  $\text{Nb}_3\text{SiTe}_6$  is a candidate double QSH system [Figure 4(d)].

However, in  $\text{Nb}_3\text{SiTe}_6$ , the two sets of linear bands also exhibit distinct, divergent behaviors in their spin polarizations [Figure 4(d)] (see Section 5 in the Supporting Information): for bands at deeper binding energies [aqua arrows in Figure 4(b,c)], the linear dispersions exhibit strong spin polarization, while for those nearest the Fermi level [blue arrows in Figure 4(b,c)], there exists two subbands with an energy splitting of  $\sim 20$  meV and very closely spaced spin-polarized bands near the Fermi level; though, all linear subbands here are more clearly spin-polarized at deeper binding energies (starting from  $E \approx -0.075$  eV). Therefore, due to the expected small energy splitting between subbands and the marked superpositions of spin polarizations near the Fermi level around the  $\overline{\text{X}}$  points, a spin-resolved transport measurement might be unable to identify a clear double QSH state in  $\text{Nb}_3\text{SiTe}_6$ . Nevertheless, this observation certainly does not discount the importance of



**Figure 5.**  $k_z$ -Dependent band mappings of hourglass fermion dispersions. (a) DFT band structure results along the XS  $k$ -path; the nodal point of the hourglass fermion in the middle of XS is marked by a red arrow, and the dashed red rectangles identify both the BZ boundaries and possible regions of interest for  $k_z$ -dependent mappings of hourglass bands near the Fermi level. (b) 2D curvature map corresponding to the  $k_z$ -dependent spectra taken at the  $\bar{X}$  point; here,  $k_z$  is varied from around 4.0 to 4.5  $\text{\AA}^{-1}$ . (c) Similar to (b) but for  $k_z$  taken from around 4.5 to 5.0  $\text{\AA}^{-1}$ . (d–g) 2D curvature band mappings along  $\bar{X}\bar{X}$  measured near  $\bar{X}$  for  $k_z = 4.0, 4.23, 4.5$ , and  $4.7 \text{\AA}^{-1}$ , respectively.

these linear band crossings to the system's nontrivial band topology.<sup>23</sup> Indeed, in a recent transport study of  $\text{Nb}_3\text{SiTe}_6$ , a nontrivial Berry phase has been found via Shubnikov-de Haas oscillation measurements,<sup>30</sup> which can be readily attributed to the linear subbands of the hourglass fermions and their unusual spin textures about the  $\bar{X}$  points. Spin-resolved ARPES measurements would be helpful for probing the spin textures of these linear subbands in the future.

Lastly, because the band structure of  $\text{Nb}_3\text{SiTe}_6$  should exhibit dispersion along the  $k_z$ -direction or the surface normal direction, as evidenced by the predicted hourglass bands along the XS  $k$ -path [Figure 1(f)], band mappings under varying incident photon energy were undertaken to directly probe hourglass fermions' band dispersions in  $\text{Nb}_3\text{SiTe}_6$ . As revealed in the Fermi surface mapping in the  $k_x-k_z$  plane [Figure 4(e)], where  $k_x$  and  $k_z$  are momentum components parallel to the  $\bar{X}\bar{X}$  and [001] directions, respectively, the linear subbands nearest the Fermi level at the  $\bar{X}$  point [Figure 4(a–d)] give rise to a continuous line along XS, though the photoemission intensity itself varies but peaks every two BZs [Figure 4(e)]. These periodic intensity variations are attributable to the weak van der Waals bonding between neighboring  $\text{Nb}_3\text{SiTe}_6$  layers, consistent with prior findings from ARPES mappings of the related hourglass semimetal  $\text{Ta}_3\text{SiTe}_6$ .<sup>27</sup> Moreover, this

periodicity is readily captured in our theoretical spectral function of  $\text{Nb}_3\text{SiTe}_6$  along XS calculated over multiple BZs, once DFT results are unfolded according to the BZ of 1T  $\text{NbTe}_2$  [Figure 5(a)]. Such experimental and theoretical results suggest that the spectral weights of hourglass bands near the Fermi level [nodal point indicated by a red arrow in Figure 5(a)] should vary periodically every two BZs [Figures 4(e) and 5(a)].

Furthermore, since the energy splitting between hourglass bands near the Fermi level is often very small [ $\sim 20$ –50 meV, per Figures 1(d) and 5(a)], 2D curvature data processing<sup>33</sup> of  $k_z$ -dependent band mappings around the  $\bar{X}$  point was employed to search for hourglass bands with characteristic dispersions along the XS  $k$ -path [Figure 5(b–g)]. As shown in Figure 5(b,c), 2D curvature band mappings along XS exhibit a downward dispersing band with an apex at the S point, which corresponds quite well with an hourglass band found in the theoretical data [Figure 5(a)]; the experimental results confirm that only half of the hourglass-like structure is accessible by photoemission [Figure 5(a–c)]. Even so, although the hourglass nodal point near  $E \approx 0$  eV is not discernible in the measured mappings [compare Figure 5(a) to Figures 5(b,c)], the observed downward band is certainly direct evidence for hourglass physics [Figure 5(a–c)] and

importantly a Dirac nodal curve in  $\text{Nb}_3\text{SiTe}_6$ , as confirmed by 2D curvature mappings around the  $\bar{X}$  point taken along  $\overline{\Gamma}\bar{X}$  at judiciously chosen  $k_z$  [Figure 5(d–g)]. Undoubtedly, the apex of the Dirac-like band crossing structure near the Fermi level varies as a function of  $k_z$ , shifting by about  $\sim+20$  meV ( $\sim-20$  meV) to lower (higher) binding energies as  $k_z$  is continuously tuned from the X to S (S to X) points [Figure 5(b–g)]. While  $\text{Ta}_3\text{SiTe}_6$  and the related system  $\text{NbSi}_{0.45}\text{Te}_2$  have been measured very recently by ARPES,<sup>26,27</sup> this Dirac nodal curve along XS and more generally all dispersions along XS of  $\text{X}_3\text{SiTe}_6$  ( $\text{X} = \text{Ta}, \text{Nb}$ ) have not been reported yet.<sup>26,27</sup> However, the strongest and most direct evidence for hourglass physics in  $\text{Nb}_3\text{SiTe}_6$  is provided by studying bands along XS, as in Figures 4(e)–5(g).

Altogether, our comprehensive photoemission and theoretical study of a bulk  $\text{Nb}_3\text{SiTe}_6$  single crystal provides direct evidence for the existence of hourglass fermions as well as unconventional spin textures in  $\text{Nb}_3\text{SiTe}_6$ . Specifically, the observation of a saddle-shaped surface in the measured band mappings and Fermi surface supports the existence of nodal loop dispersions near the zone center, in corroboration with prior transport works.<sup>28</sup> Additionally, the identification of two species of Dirac-like, linear band crossings at the  $\bar{X}$  point, together with their theoretical spin-resolved band structures, support a scenario for an exotic double QSH state in  $\text{Nb}_3\text{SiTe}_6$ , previously only discovered in the hourglass semimetal  $\text{KHgSb}$ .<sup>34</sup> Furthermore, our photon-energy-dependent or  $k_z$ -dependent mappings along XS—never undertaken in any ARPES work on the orthorhombic family of  $\text{X}_3\text{SiTe}_6$  ( $\text{X} = \text{Ta}, \text{Nb}$ ) single crystals<sup>26,27</sup>—have unveiled an hourglass band that is a Dirac nodal curve, based upon our DFT results and experimental data. Though only half of an hourglass-like structure is accessible through photoemission, the  $k_z$ -dependent band maps nonetheless suggest that hourglass fermions must contribute to the electronic properties of  $\text{Nb}_3\text{SiTe}_6$  due to the proximity of their band structures near the Fermi level. All in all, our work not only provides a future impetus in the identification of even more unusual topological materials but also identifies  $\text{Nb}_3\text{SiTe}_6$  as a candidate system harboring unconventional spin textures accessible to spintronic applications.

## ■ ASSOCIATED CONTENT

### SI Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.nanolett.2c03293>.

Additional information about the sample preparation, the experimental setup, the DFT calculation details, XRD measurement, XPS measurement, and spin-resolved DFT calculations (PDF)

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### Author Contributions

RYL, MKL, PC, and JAH performed ARPES measurements at the ALS or NSRRC, where JD and CMC provided technical help during beamline operations. AH and HTJ performed theoretical calculations. RS synthesized the single crystal. SCW performed XRD measurements. CCS and TMC performed STM measurements. RYL and TCC analyzed the data, interpreted the results, and wrote the first draft. All coauthors contributed to discussions and improvements that led to the final version of the manuscript.

**Notes**

The authors declare no competing financial interest.

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