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(Article begins on next page)

Title: Spin-Berry curvature enhanced orbital Zeeman effect in a Kagome metal

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Abstract:

Berry phases and the related concept of Berry curvature can give rise to many unconventional phenomena in solids. Here, we discover colossal orbital Zeeman effect of topological origin in a bilayer kagome metal TbV₆Sn₆. Using spectroscopic-imaging scanning tunneling microscopy, we reveal that the magnetic field leads to the splitting of the gapped Dirac dispersion into two branches with enhanced momentum-dependent g factors, resulting in the substantial renormalization of the Dirac band. These measurements provide a direct observation of the magnetic field controlled orbital Zeeman coupling to the orbital magnetic moments of up to 200 Bohr magnetons near the gapped Dirac points. Our work provides the direct insight into the momentum-dependent nature of topological orbital moments and their tunability by magnetic field concomitant with the evolution of the spin-Berry curvature. These results can be extended to explore large orbital magnetic moments driven by the Berry curvature governed by other quantum numbers beyond spin, such as the valley in certain graphene-based structures.

Main text:

The Berry phase of electron wave functions can strongly influence the low-energy physics of solid state systems and give rise to various unconventional phenomena such as ferroelectricity, the topological Hall effect, and orbital magnetism ¹⁻¹¹. Materials with flat bands and Dirac crossings are generally believed to be ideal places where such topological phenomena can be realized, in large part due to finite Berry curvature of the bands amplified at particular momentum-space positions. For example in Dirac systems, a small Dirac mass gap can be generated by the spin-orbit coupling, which in turn produces enormous Berry curvature concentrated at the momentum of the Dirac crossing. Lifting the degeneracy of the electronic bands characterized by large Berry curvature can lead to unusual magnetism originating from purely orbital contributions, which has so far been detected in certain exfoliated Moire structures ³. Although Berry curvature effects permeate much of the physics of topological materials in condensed matter physics ^{1,2,4-11}, spectroscopic insights into these phenomena have been challenging. In particular, while Berry curvature is a quantity intrinsically dependent on crystal momenta, experimental detection of ensuing phenomena with momentum resolution is yet to be achieved.

Kagome metals have emerged as a highly tunable platform that is ideal for the explorations of exotic electronic phenomena intertwined with non-trivial band topology. The experimental efforts thus far have been focused on two different kagome metal sub-families. On the one side, magnetic kagome systems with Fe-¹²⁻¹⁹, Mn-²⁰⁻²⁴ or Co-^{8,25,26} kagome nets, in which magnetic order of the transition metals in the kagome plane gives rise to various forms of magnetism, have been explored to study Dirac fermions ^{12,13}, topological flat bands ^{8,27}, Weyl fermions ²⁶ and Fermi arcs ^{25,26}. These magnetic metals typically show a strongly spin-polarized electronic band structure, favorable for the realization of massive Dirac fermions and the Chern magnet physics ²⁸ due to the strong time-reversal symmetry breaking associated with the magnetic order. Dirac fermion bands in these systems can be tuned by an external magnetic field, albeit the process is generally intertwined with spin canting effects such as that in Fe₃Sn₂¹³ and YMn₆Sn₆²⁴. On the other side, an entirely non-magnetic V-based AV₃Sb₅ (A=Cs, K, Rb) kagome metal family ^{29–31} with a spin-degenerate band structure attracted a tremendous attention ^{32–44} due to the emergence of various density waves, exotic Fermi surface instabilities and unusual superconductivity.

The recently discovered family of V-based bilayer kagome metals in the RV_6Sn_6 (R = rare earth ion) structure offers a tunable physical regime between these two existing classes of layered kagome metals – magnetism can still be induced by the choice of the rare-earth element R, but the vanadium atom in the kagome layers remains non-magnetic ^{45–48}. Moreover, the R-induced magnetism does not seem to significantly affect the low-energy vanadium orbital derived band structure. Measurements have already demonstrated the robustness of the canonical features of the kagome systems including Dirac cones and flat bands ^{45,49,50}, thus setting a fresh stage for the search for unconventional topological phenomena. In this work, we use spectroscopic-imaging scanning tunneling microscopy in combination with first-principles electronic structure calculations to discover colossal orbital Zeeman effect of topological origin in TbV₆Sn₆. The observed orbital Zeeman splitting of the Dirac bands is fueled by the large spin-Berry curvature induced orbital magnetic moments coupling to external magnetic field. We find that the effect is surprisingly non-linear and that it changes concomitant with the evolution of the Berry curvature.

We focus on the bilayer kagome metal TbV₆Sn₆, which is a low-temperature soft ferromagnet ($T_c \approx 4.3$ K) due to ordering of the Tb³⁺ magnetic moments ^{46,47}. The crystal structure of TbV₆Sn₆ consists of a V atom kagome net inserted between alternating Sn¹-Sn²-Sn¹ slabs and TbSn³ layers (Fig. 1a,b). We cleave bulk single crystals of TbV₆Sn₆ in ultra-high vacuum and immediately insert them into the STM head (Methods). STM topographs reveal a layered structure with *ab*-plane terraces on the surface, and step heights consistent with a single unit cell of TbV₆Sn₆ (Fig. 1c). We identify the termination predominantly observed in Fig. 1b,c as the Sn² termination due to the clearly resolved individual atoms forming a hexagonal lattice (top inset in Fig. 1d) and its similarity with the equivalent Sn² surface imaged in YMn₆Sn₆²⁴.

Average dI/dV spectra over a wide energy range show a spectral peak and a sharp upturn in conductance around 200 meV above the Fermi level (bottom inset in Fig. 1d). Temperature-dependent dI/dVmeasurements reveal that spectra remain indistinguishable upon raising the temperature above the Curie temperature T_c (Fig. 2a) and well below T_c (Extended Data Figure 1), suggesting that the electronic structure in this energy range is not affected by the ferromagnetic ordering. Interestingly however, the spectra show a pronounced change as magnetic field is applied perpendicular to the sample surface (Fig. 2b,c). In particular, the spectral peak appears to split into two peaks that shift in opposite directions, i.e., one towards lower energy and the other conversely to higher energy (Fig. 2b,c). Magnetic field-induced spectral splitting is also robustly observed well inside the ferromagnetic state and well above T_c (Fig. 2d), which again points to the evolution of electronic band structure largely unaffected by the localized moments of Tb atoms that saturate below $0.3-0.4 \text{ T}^{46,47}$ (Supplementary Information Note 1).

To gain further insight into the low-energy electronic band structure in momentum-space (k-space), we proceed to use spectroscopic-imaging scanning tunneling microscopy. The technique, also referred to as quasiparticle interference (QPI) imaging, relies on scattering and interference of electrons detectable as periodic conductance modulations in $dI/dV(\mathbf{r}, V)$ maps to provide an insight into the momentum-space electronic band structure. To obtain a high momentum-space resolution, performing measurements over large regions of the sample is desirable (Fig. 3a). We first focus on the spectroscopic mapping at zero magnetic field. $dI/dV(\mathbf{r}, V)$ maps show strong "ripples" in real-space centered at defect sites (Fig. 3b), reflected in an approximately isotropic wave vector \mathbf{q}_1 seen around the center of associated Fourier transforms (Fig. 3c). The wave vector magnitude changes with energy, which can be visualized in the radially-averaged dispersion shown in Fig. 3d. From this plot, we can observe that the imaged electronic band exhibits a nearly linear dispersion with velocity 1315 \pm 64 meV·Å, starting at about 170 meV and extending over a large energy range to above 380 meV, where the signal disappears. We attribute the existence and the dispersion of q₁ to intra-band scattering arising from a gapped Dirac fermion dispersion at K (inset in Fig. 3d) for the following reasons. First, the dispersion velocity obtained from our data shows excellent agreement with the Dirac dispersion at K point appearing in a similar energy range in our theoretical calculations (Fig. 3d). Second, the morphology and the dispersion of q_1 is qualitatively similar to that measured in helical antiferromagnet YMn₆Sn₆ that has also been attributed to a Dirac dispersion at K²⁴. Third, the QPI bandwidth of about 200 meV matches closely to the width of the upper part of the same Dirac dispersion in density functional theory calculations ⁵¹. We note that the Dirac point imaged here at about +150 meV is different from that reported in Ref. ⁴⁷, which focuses on the lower Dirac crossing positioned at -200 meV. Based on density functional theory calculations, the lower Dirac crossing is separated from the upper one by about 400 meV, which is nicely consistent with the two different Dirac dispersions imaged by angle-resolved photoemission spectroscopy ⁴⁷ and STM (Fig. 3).

Next, we apply magnetic field parallel to the *c*-axis and measure the concomitant response of the Dirac dispersion at 4.2 K. Our field-dependent QPI measurements reveal that the Dirac band dispersion exhibits a pronounced field-induced change. Notably, the band appears to split into two branches – one shifting upwards to higher energy and the other one shifting to lower energy (Fig. 3d-h, Extended Data Figure 2, Extended Data Figure 3). This observation suggests that the Dirac band at zero magnetic field is spin-degenerate, consistent with the combined action of inversion and time-reversal symmetries. Similarly to the spectral peak saturation at magnetic fields $B \ge 4$ T in Fig. 2, the Dirac dispersion measured in the same range of magnetic fields appears indistinguishable (Fig. 3g,h), pointing to an intimate connection between the two observables. As the spectral peak emerges at approximately the same energy as the bottom of the massive Dirac band and shifts concomitantly with the band evolution (Fig. 4a,b), the spectral peak in Fig. 2 is likely a consequence of the large density of states near the Dirac band bottom, as also hypothesized to be the case in the Mn-based cousin, TbMn₆Sn₆²⁸. This interpretation is further confirmed by examining the field-induced shift of d//dV peaks and the QPI bands, which both exhibit a nearly identical behavior (Fig. 4e,f).

We stress that the field-induced band structure evolution observed here cannot be explained by spin canting, as reported for example, in kagome magnet Fe_3Sn_2 ^{13,18}. From magnetization measurements, it is clear that the magnetization in TbV₆Sn₆ reaches a plateau at a magnetic field of less than 0.3-0.4 T ^{46,47}, so the saturation at much higher fields reported here is rather peculiar and cannot be explained in the

framework of localized Tb magnetic moments that are already saturated at low fields. Further supporting the notion that the role of ferromagnetism from Tb moment ordering driving our observations is negligible, we point that the electronic band structure in the relevant energy range is indistinguishable in the ferromagnetic and paramagnetic states (Figure 2, Extended Data Figure 1, Supplementary Information Note 1).

This brings up an intriguing question of the origin of our observations. As the effect of local Tb magnetic moments can be largely excluded (Supplementary Information Note 1), we inspect itinerant orbital magnetic moments rooted in Berry curvature that can in some cases reach enormous levels ^{1,9,52}. We explicitly calculate orbital magnetic moments in this system associated with the massive Dirac dispersion imaged in our experiments. The large spin Berry curvature of the Dirac bands around the K point (see right inset of Fig. 1e for the spin up channel) prompts orbital moments of several meV/T (Fig. 4j, Extended Data Fig. 4). In addition, the orbital moment inversely depends upon the size of the Dirac masses, which can be theoretically adjusted by an artificial tuning of the local spin-orbit term. One can see that the maximum values of the orbital moment in the band region of interest change considerably, increasing as the Dirac masses decrease. Interestingly, the experimental band evolution is also not rigid – visual comparison near the band bottom reveals substantial rounding of the band curvature at higher fields (Fig. 4g). Our theoretical calculations suggest that both the gradual change in the effective g factor with magnetic field and the anomalously large magnitude can be explained by the non-trivial Berry curvature effects associated with the Dirac band, giving rise to orbital magnetic moments. The non-linearity of the effect (Fig. 4e,f) may suggest that second-order corrections may be necessary to fully capture the underlying physics.

It is also worth noting that the magnitude of the band shift in our work is enormous (up to 11.20 meV/T), amounting to the magnetic moment on the order of 200 Bohr magnetons. This is more than two orders of magnitude higher than the conventional spin Zeeman effect *g*-factor $g_s \approx 2$ that would lead to 0.058 meV/T. It is also much larger than what has been detected from spectral peak shifts in Co₃Sn₂S₂ (0.078 meV/T ⁵³ and 0.174 meV/T ⁸), which were also attributed to orbital moments. Aside from the much larger magnitude reported here, our momentum-space evolution allows us to unambiguously determine the origin of these moments to arise from the Dirac band. At the quantitative level of comparison with our theoretical analysis, the experimental orbital moment is larger than the calculated one. Nonetheless, we stress here that the precise size of Dirac gaps and the strength of spin-orbit coupling might also be not well captured by density functional-based calculations, specifically when electronic correlations could be relevant ^{54,55}.

Our experiments provide a direct visualization of orbital Zeeman splitting of topological origin and the momentum-space insight into the surprisingly large orbital magnetic moments fueled by the enormous spin-Berry curvature at the gapped Dirac point. Our observations directly demonstrate that the opening of the Dirac gap in this system is not due to the time-reversal symmetry breaking magnetic order of the Tb atoms. It rather arises from strong spin-orbit coupling, which in turn leads to a non-zero spin Berry curvature. The latter generates the orbital magnetic moment $\pm m(k)$, opposite for the two spin species but pointing in the *c*-axis. In a time-reversal symmetric situation, the opposite orbital magnetic moments for the two spin species would exactly cancel out; however, when time-reversal symmetry is broken by magnetic field, the two branches can become decoupled and split in external magnetic field. The applied magnetic field **B** along the *c*-axis couples to the dominating orbital magnetic moments by orbital Zeeman coupling $\pm m(k)B$ (ignoring the much smaller spin-Zeeman coupling) and lifts the degeneracy of the

approximately spin-degenerate Dirac band. This allows the Dirac branches with opposite spins and opposite sign of the orbital moments to shift in opposite directions, as seen in our STM experiments.

We stress that our results here unveil a different physical mechanism compared to previous experiments imaging field-tunable electronic band structure ^{13,24}. In for example Fe₃Sn₂ ¹³ and YMn₆Sn₆ ²⁴, the observed electronic band structure change inevitably happens in a regime where spins are continuously tilting with field, which makes it impossible to disentangle spin effects from orbital magnetic moment contributions, if any. Fe₃Sn₂ and YMn₆Sn₆ also exhibit a strongly spin-polarized electronic band structure with singlydegenerate Dirac dispersions composed of magnetic Fe or Mn d orbitals even in the absence of magnetic field. In our experiment on TbV₆Sn₆ here, we can conclude that the effects of Tb magnetic moments are minimal (Supplementary Information Note 1), which enables the momentum-resolved visualization of the elusive orbital physics and orbital Zeeman effect. Interestingly, the anomalously large magnitude of the orbital magnetic moments due to spin-Berry curvature in TbV₆Sn₆ may be attributed to the combination of a high Dirac band velocity and the relatively small spin-orbit coupling in a simple effective theory description (Supplementary Information Note 2). This could also provide a guiding principle to look for topological moments of even larger magnitudes in other materials, by focusing on maximizing the Dirac band velocity while minimizing the spin-orbit coupling induced Dirac gap. The striking evolution of effective magnetic moments from enormous to vanishing provides a surprising experimental insight into the phenomenological behavior of topological orbital magnetic moments strongly dependent on the Dirac band curvature and associated Dirac gap.

It is interesting to note that orbital magnetic moments that arise from Berry curvature of the Bloch electronic states can be generated by other quantum numbers. For instance, graphene-based systems are expected to also show orbital magnetic moments tunable by magnetic field. In single layer graphene, inversion symmetry breaking due to the attachment to the substrate creates a Dirac gap. However in contrast to TbV₆Sn₆, the relevant degree of freedom is the valley degree of freedom (K vs K'), not spin ⁵⁶. So in single layer graphene, there will also be Berry curvature induced orbital magnetic field could in principle couple to these, shifting one higher and the other one lower in energy, but there would be no orbital Zeeman splitting or spin Berry curvature. To visualize similar Berry curvature induced phenomena in graphene and Moire heterostructures, momentum-resolved magnetic field dependent imaging of the electronic band structure we presented in our work would be highly desirable.

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Author Contributions Statement:

H.L. performed the STM experiments. H.L analyzed the STM data with the help from S.C. G.P. synthesized and characterized the samples under the supervision of S.D.W. P.E., G.S. and D.D.S. performed density functional theory calculations. Z.W. provided theoretical input on the interpretation of STM data. C.B. and F.M. provided insights on angle-resolved photoemission spectroscopy data in connection to STM data. H.L., G.S., D.D.S, S.D.W., Z.W. and I.Z. wrote the paper with the input from all authors. I.Z. supervised the project.

Competing Interests Statement:

The Authors declare no Competing Financial or Non-Financial Interests.

Figure Legends:



Figure 1. TbV₆**Sn**₆ **crystal structure and electronic band structure. (a)** Ball model representing the layer stacking order. (b) A 3D ball model showing the ferromagnetic structure of TbV₆Sn₆ (blue arrows represent the Tb spins pointing along the *c*-axis). (c) STM topograph of a 180 nm square region spanning multiple consecutive unit cell steps between Sn² hexagonal layers. The inset is the topographic line profile extracted along the black dashed line. Each step height is 9.1 Å, which equals the *c*-axis lattice constant. (d) STM topograph of Sn² termination. The upper right inset shows a zoom-in view of a small region and a hexagonal Sn lattice. The bottom left inset is a spatially averaged dI/dV spectrum taken in region in (c). (e) Density functional theory calculations of TbV₆Sn₆ electronic band structure along the Γ-K-M-Γ line. The thick pink band is a gapped Dirac cone above the Fermi level, which carries a large spin Berry curvature for each spin channel (depicted in the right inset for spin up). More details of the relevant portion of the band structure with and without spin-orbit coupling can be found in Extended Data Figure 4. The left inset is a constant energy contour (CEC) around the K point at 120 meV. The pink circle is the CEC of the Dirac band at K. STM setup conditions: *I*_{set} = 10 pA, *V*_{sample} = 1V (b); *I*_{set} = 400 pA, *V*_{sample} = 100 mV (c).



Figure 2. The evolution of electronic density of states as a function of magnetic field and temperature. (a) Spatially-averaged dI/dV spectrum as a function of temperature taken on the Sn₂ termination at 0 T. (b) Spatially averaged dI/dV spectrum and (c) associated second derivative of dI/dV as a function of magnetic field taken on the Sn₂ termination at 4.2 K. It can be seen that the dominant spectral feature at 0 T splits into two branches, at energies labeled as E^* and E^{**} . Their dispersion saturates at higher fields. (d) Magnetization curve of TbV₆Sn₆ at 2 K (top panel) and evolution of E^* as a function of magnetic field at 2 K (ferromagnetic, FM), at 4.2 K (near the transition point between FM and paramagnetic, PM phase) and at 10 K (PM) respectively. E^* data points at 10 K were acquired over a different microscopic region of the sample tens of micrometers away compared to the 2 K and 4.2 K data; the 10 meV difference between E^* at 10 K and those at lower temperatures is likely due to small chemical potential shift between different regions. Blue, purple and red dashed curves are visual guides denoting trends in the data. STM setup conditions: $I_{set} = 1$ nA, $V_{sample} = 300$ mV, $V_{exc} = 5$ mV (a); $I_{set} = 1$ nA, $V_{sample} = 300$ mV, $V_{exc} = 2$ mV (b,c).



Figure 3. Momentum-space visualization of the Dirac band using spectroscopic imaging scanning tunneling microscopy. (a) STM topograph of the Sn² hexagonal termination. (b) Representative $d/dV(\mathbf{r}, V)$ maps of the region in (a) at biases 300 mV, 250 mV and 200 mV (from left to right panel) and (c) their Fourier transforms. Pink arrow denotes the dominant QPI wave vector **q** and the atomic Bragg peaks are circled in blue. (d-h) Energy-dependent radially-averaged linecuts S(q, E), where E denotes energy, starting from the center of the Fourier transforms in (c) at different magnetic fields applied perpendicular to the sample. The bottom right insets in (d-h) show the associated second derivatives of (d-h): $\partial_a \partial_E S(q, E)$. The bottom left inset in (d) refers to the density functional theory calculated Dirac band at K point slightly above Fermi level. By shifting the chemical potential down by 130 meV likely due to chemical doping by accidental defects, the Dirac band (dashed pink line in (d)) matches the STM QPI dispersion (gray background) very well. In addition to the Dirac bands, we also observe another wave vector at very low q that is nearly dispersionless but shifts with magnetic field (arrow in (f)), which we hypothesize it may originate from another band. Before S(q, E) maps in (d-h) are generated, each dI/dVmap is processed by subtracting the average value to artificially suppress the bright center in the resulting Fourier transform and smoothed on 1 pixel length scale. We note that we apply the 1:2 conversion between momentum-transfer space (q-space) and momentum space (k-space) to superimpose the density functional theory calculated band (pink dashed lined) on top of our STM data in (d). This conversion is applied because QPI wave vector q is a back-scattering vector, connecting k and -k points on the near-circular constant energy contour of the Dirac cone at K in momentum space. STM setup conditions: I_{set} = 1 nA, V_{sample} = 300 mV (a); I_{set} = 1.5 nA, V_{sample} = 300 mV, V_{exc} = 2 mV (b). Data were taken at 4.2 K.



Figure 4. Dirac band evolution and the saturation of the effective q factor. (a,b) The second derivative of radially averaged linecuts in the Fourier transforms of the dI/dV maps at 0 T (a) and 6 T (b). The right sides of the panels in (a, b) show the corresponding spatially averaged dI/dV spectra. The blue dashed lines denote the features in spectra likely corresponding to the band bottom. (c) The linecut taken along the red solid line in (a,b), as well as in equivalent datasets at 1 T, 2 T and 4 T. Purple squares and red triangles denote the band bottom positions in the energy axis. (d) Second derivative of dI/dV spectra from 0 T to 6 T. Orange circles and black triangles denote the peaks in the spectra. (e) Evolution of spectral markers with magnetic field extracted from (c,d) (color and symbol-coded to match the corresponding peaks in (c,d)). (f) Band splitting and spectral peak splitting extracted from (e). Brown line is extracted from QPI measurements representing band splitting, and the gray line is extracted from $-d^3I/dV^3$ curves representing the spectral peak splitting. (g) Radially averaged Fourier transform linecut starting at the center of the Fourier transform at 0 T and 6T (same as in insets in Fig. 3d-h), and the extracted band points denoted as $E_0(q)$ for the Dirac band at OT, $E_1(q)$ for top Dirac band at 6 T and $E_2(q)$ for the bottom band at 6 T. The analysis of raw Fourier transform data can be found in Extended Data Figure 5. (h,i) The difference between energies of equivalent momentum-space states at different fields, obtained by the subtraction of $E_1(q)$ and $E_0(q)$ (h), and $E_2(q)$ and $E_0(q)$ (i). (j) Theoretically calculated evolution of the orbital moment m at the Dirac point (k=0) for the topmost Dirac cone upon a continuous change of the Dirac gap Δ . The gap is artificially tuned by acting on the strength of the local spin-orbit coupling in our first-principles calculations (Methods, Extended Data Figure 4). STM setup conditions: Iset = 1.5 nA, Vsample = 300 mV, Vexc = 2 mV (a,b).

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Methods

Bulk single crystal synthesis. Single crystals of TbV_6Sn_6 were synthesized from Tb (pieces, 99.9%), Sn (shot, 99.99%), V (pieces, 99.7%) using a conventional flux-based growth technique. The flux mixtures of Tb, V, Sn were loaded inside a crucible with a molar ratio of 1:6:20 and then sealed in a quartz tube filled with Argon atmosphere. The sealed tube was heated at 1125 °C for 15 hours and then slowly cooled down at a rate of 2 °C/h to 780 °C. Thin millimeter-sized shiny plate-like single crystals were separated from the molten flux via centrifuging at 780 °C.

STM experiments. Single crystals of TbV_6Sn_6 were cleaved in ultrahigh vacuum and immediately inserted into the STM head, where they were held at ~4.5 K during measurements. STM data were acquired using a customized Unisoku USM1300 STM system. Spectroscopic measurements were made using a standard lock-in technique with 910-Hz frequency and bias excitation as detailed in the figure captions. STM tips used were home-made electrochemically etched tungsten tips, annealed in UHV to a bright orange color before STM experiments.

Theoretical calculations. We employed first-principles calculations based on the density functional theory as implemented in the Vienna ab-initio simulation package (VASP) ⁵⁷, within the projector-augmented plane-wave (PAW) method ⁵⁸. The generalized gradient approximation as parametrized by the PBE-GGA functional for the exchange-correlation potential is used ⁵⁹ by expanding the Kohn–Sham wave functions into plane waves up to an energy cutoff of 400 eV. We sampled the Brillouin zone on an 12x12x6 regular mesh by including SOC self-consistently. For the calculation of the spin Berry curvature and orbital magnetic moments, the Kohn–Sham wave functions were projected onto a Tb *d*, V *d* and Sn *s*, *p*-type basis ⁶⁰, and the quantities were computed by using our in-house post-wan library [https://github.com/philipp-eck/post_wan], consistent with Ref. ⁶¹.

Data Availability

The data supporting the findings of this study are available on Zenodo (https://zenodo.org/records/10800955) and upon request from the corresponding author.

Code availability

The computer code used for data analysis is available upon request from the corresponding authors.

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