Linear bands, zero-momentum Weyl semimetal, and topological transition in skutterudite-structure pnictides

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It was reported earlier [Phys. Rev. Lett. 106, 056401 (2011)] that the skutterudite structure compound CoSb3 displays a unique band structure with a topological transition versus a symmetry-preserving sublattice (Sb) displacement very near the structural ground state. The transition is through a massless Dirac-Weyl semimetal, point Fermi surface phase which is unique in that (1) it appears in a three-dimensional crystal, (2) the band critical point occurs at $k = 0$, and (3) linear bands are degenerate with conventional (massive) bands at the critical point (before inclusion of spin-orbit coupling). Further interest arises because the critical point separates a conventional (trivial) phase from a topological phase. In the native cubic structure this is a zero-gap topological semimetal; we show how spin-orbit coupling and uniaxial strain converts the system to a topological insulator (TI). We also analyze the origin of the linear band in this class of materials, which is the characteristic that makes them potentially useful in thermoelectric applications or possibly as transparent conductors. We characterize the formal charge as Co$^+$ $d^9$, consistent with the gap, with its 3 site symmetry, and with its lack of moment. The Sb states are characterized as $p_z$ (separately, $p_x$) $\sigma$ -bonded Sb$_2$ ring states occupied and the corresponding antibonding states empty. The remaining (locally) $p_z$ orbitals form molecular orbitals with definite parity centered on the empty 2a$_2$ site in the skutterudite structure. Eight such orbitals must be occupied; the one giving the linear band is an odd orbital singlet $A_{2u}$ at the zone center. We observe that the provocative linearity of the band within the gap is a consequence of the aforementioned near-degeneracy, which is also responsible for the small band gap.

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I. MOTIVATION AND INTRODUCTION

Recent years have seen an explosion of interest in two areas that, while distinct in themselves, have also found some areas of overlap. One is the situation where Dirac-Weyl linear bands emanate from a point Fermi surface (FS), the so-called Dirac point. While linear bands around symmetry points are not uncommon in crystal structures, it is uncommon that such points can determine the Fermi energy (EF), which is what happens in the celebrated case of graphene. It is necessary that there is a gap throughout the Brillouin zone except for the touching bands. Point FSs occur also in the case of conventional (massive) zero-gap semiconductors, and give rise to properties that have been studied in some detail. The case of the Dirac-Weyl semimetal (point Fermi surface) systems is quite different, leading to a great deal of new phenomenology.

The other new and active area relevant to this paper is that of topological insulators (TIs), in which the Brillouin zone integral of a certain “gauge field” derived from the $k$ dependence of the periodic part of the Bloch states of the crystal is nonvanishing. These integer-valued topological invariants delineate TIs from conventional (“trivial”) insulators, with distinctive properties that are manifested primarily in edge states. Identification of TIs, and study of their properties and the critical point that separates the phases, is highlighting new basic physics and several potential applications.

Recently we reported on the skutterudite structure compound CoSb3, which combines and interrelates several of these properties (Dirac-Weyl semimetal at $k = 0$; critical point of degeneracy with massive bands; transition to topological insulator) in a unique way. The skutterudite structure, which is illustrated in Fig. 1 and is discussed in Sec. II, is critical to the electronic behavior in CoSb3, though it may not be necessary for the unique type of critical point that arises. The relevant portion of the band structure, in a very small volume of the Brillouin zone (BZ) centered at Γ, is unusually simple compared to most of the TIs that have been discovered, viz. the Bi$_2$Se$_3$ class, the HgTe class, and the three-dimensional system HgCr$_2$Se$_4$.

The electronic structure of skutterudites was studied early on by Singh and Pickett, who focused on the peculiar valence band which was linear (except exceedingly near Γ) and whose linearity extended surprisingly far out into the BZ. It became clear that the linearity of the band was responsible for the large thermopower that is potentially useful in thermoelectric applications. The origin of the linearity was however not identified, but it leads to peculiar consequences: the density of states varies as $k^2$ near the band edge rather than the usual three-dimensional (3D) form $\sqrt{\mathbf{E}}$; as a consequence the carrier density scales differently with Fermi energy $E_F$; the inverse mass tensor $\nabla \nabla E_k$ is entirely off-diagonal corresponding to an “infinite” transport mass; the cyclotron mass differs from conventional 3D behavior, etc. As the Sb sublattice is shifted in a symmetry-preserving way, the small semiconducting gap of CoSb3 closes at a critical point, giving rise to an unusual occurrence: a Dirac-Weyl point in a 3D solid at the Γ point. This transition point sets...
FIG. 1. (Color online) Crystal structure of skutterudite minerals (viz. CoSb3) with space group $Im\bar{3}$ (#204), which includes inversion. The Co site [small pink (dark gray) sphere] is octahedrally coordinated to Sb atoms [small yellow (light gray) spheres]. Each Sb atom connects two octahedra, as in the perovskite structure which has the same connectivity of octahedra. The large (blue) sphere denotes a large open site that is unoccupied in CoSb3 but is occupied in filled skutterudites (see for example Ref. 19). The geometrical solid (center of figure) provides an indication of the volume and shape of the empty region.

the stage for a conventional insulator to topological insulator transition, although the specific behavior of the system at this point was not spelled out in our earlier work. The only other topological insulator phases in skutterudite materials that we are aware of are the examples by Jung et al.18 in CeOs4Pn12 filled skutterudites.

This paper is organized as follows. In Sec. II the skutterudite structure and its relation to the symmetric perovskite (ReO3) structure is reviewed. The computational methods are outlined in Sec. III. Section IV discusses the evolution of the band structure during a perovskite-to-skutterudite transformation, including the development of the gap and the appearance of the linear band. Section V is devoted to constructing a microscopic but transparent understanding of the electronic structure, including the development of the energy gap and the character of the peculiar linear band and the lower conduction band triplet that it interacts with near, at, and beyond the critical point. In Sec. VI we provide details of the band inversion leading to the topological transition, and analyze the anisotropy at the critical point. A brief summary is provided in Sec. VII.

II. STRUCTURE AND RELATION TO PEROVSKITE

It is useful for the purposes of this paper to consider the skutterudite structure to be a strongly distorted perovskite ATPn3 structure in which the A atom is missing (i.e., the ReO3 structure) and the interconnected TPa6 octahedra are rotated substantially ($T =$ transition metal; Pn = pnictogen). The skutterudite structure,20–22 pictured in Fig. 1, has space group $Im\bar{3}$ (#204) and a body-centered cubic (bcc) Bravais lattice, and is comprised of a bcc repetition of four formula units (f.u.) when expressed as TPa3. The pnictide (Pn) atoms form bonded units (nearly square but commonly designated as “rings”) which are not required by local environment or overall symmetry to be truly square. The three Pn4 squares in the primitive cell are oriented perpendicular to the coordinate axes. Transition metal (T) atoms (usually in the Co column) lie in six of the subcubes of the large cube of lattice constant $a$; the other two subcubes (octants) are empty. The structure has inversion symmetry and is symmorphic, with 24 point group operations. The cubic operation that is missing is reflection in (110) planes. The related filled skutterudites XT4Pn12 have an atom X incorporated into the large 2$a$ site of 3$m\bar{m}$ symmetry23 that remains empty in the compounds that we discuss.

Figure 2 illustrates two aspects of the structure. The top panel reveals that, although the Sb6 octahedra are rotated considerably (and strained) compared to the perovskite structure, a great deal of regularity is retained. The bottom panel gives a picture of the relationship between the Co atoms and the six Sb4 rings that it is coordinated with (only three mutually
orthogonal rings are pictured). A natural local coordinate system for describing an Sb₄ unit is to let it lie in the x-y plane; then the pₓ, pᵧ orbitals we will discuss will be oriented perpendicular to the plane of the Sb unit. The pₓ, pᵧ orbitals are nearly symmetry related and naturally lead to bonding and antibonding molecular orbitals, and hence are very similar in character, while the nonbonding pᵧ orbitals have distinctive character.

If the origin is chosen so the large empty sites are centered at \((\frac{1}{4}, \frac{1}{4}, \frac{1}{4})\) and \((\frac{3}{4}, \frac{3}{4}, \frac{3}{4})\), then the rings are centered in each of the other subcubes \((\frac{1}{4}, \frac{1}{4}, \frac{3}{4}), \frac{3}{4}, \frac{3}{4}, \frac{3}{4})\). A ring oriented perpendicular to \(\hat{z}\) is neighbored by rings above and below \((\pm \hat{z})\) directions) oriented perpendicular to (say) \(\hat{y}\), by rings in the \(\pm \hat{x}\) directions perpendicular to \(\hat{x}\), and neighbored by the empty sites along the \(\pm \hat{y}\) directions. A symmetric combination of the four pₓ orbitals on a ring, call it \(Pₓ\), is orthogonal to the \(Pᵧ\) orbitals on neighboring rings, so if dispersion is governed by interring hopping (rather than through Co atoms) some rather flat Sb pₓ bands should result. Such behavior is seen in fatbands plots (see below).

As mentioned, the skutterudite structure is related to the perovskite structure \(\square TPn₃\) (\(\square\) denotes an empty A site). Beginning from perovskite, a rotation of the octahedra keeping the Pn atoms along the cube faces results in the formation of the (nearly square) Pn₄ rings, and the Pn octahedra become distorted and less identifiable as a structural feature. The transformation is, in terms of the internal coordinates \(u\) and \(v\),

\[
u'(s) = \frac{1}{4} + s (u - \frac{1}{4}); \quad v'(s) = \frac{1}{4} + s (v - \frac{1}{4})\].

The transformation path, from perovskite for \(s = 0\) to the observed structure for \(s = 1\), is pictured in Fig. 1 of Ref. 24. Below we make use of this transformation to follow the opening of the (pseudo)gap between occupied and unoccupied states.

III. COMPUTATIONAL METHODS

Two all-electron full-potential codes, FPLO-9 25 and WIEN2k 26 based on the augmented plane wave+local orbitals (APW+lo) method 27 have been used in these calculations, with consistent results. The Brillouin zone was sampled with regular \(12 \times 12 \times 12 \ k\) mesh during self-consistency. For WIEN2k, the basis size was determined by \(R_{\text{mt}}K_{\max} = 7\). Atomic radii used were 2.50 a.u. for Co and 2.23 a.u. for Sb, and the Perdew-Wang form 28 of exchange-correlation functional was used. The experimental lattice parameters and atomic positions were taken from Refs. 20–22.

IV. PEROVS KITE TO SKUTTERUDITE TRANSFORMATION

The development of the electronic spectrum as the crystal is distorted from perovskite to the skutterudite structure is pictured in the three panels of Fig. 3. For the ideal perovskite the result is a highly metallic state, with no gap or pseudogap near the Fermi level. The important features only arise near the end of the distortion path. For \(s = 0.75\) (top panel of Fig. 3) the valence bands have just become disjointed from the conduction band and gap formation is imminent. For \(s = 0.90\) the gap is well formed and the minimum direct gap is emerging at the \(\Gamma\) point. The unusual high velocity band arising from the \(\Gamma\) point, already striking at \(s = 0.75\), remains unchanged at this point. The valence bands are a mass of indecipherable spaghetti.

By \(s = 0.95\) (bottom panel) a similarly high velocity band is emerging from the valence bands to the maximum at the \(\Gamma\) point. At \(s = 1.0\) it becomes clear that it is a partner...
of the high velocity conduction band, as discussed in our earlier paper\textsuperscript{10} and to which we return in the next section. No doubt the formation of the gap, that is, formation of occupied bonding bands and unoccupied antibonding bands, is behind the stability of the skutterudite structure in the Co pnictides class of materials. The fact that the gap only opens as the Sb $p_z$ and $p_x$ bonding-antibonding splitting becomes strong supports the picture that this $t_{ppz}$ interaction takes precedence over the Co-Sb interactions within the CoSb\textsubscript{6} unit; however, these latter interactions (i.e., the presence of the Co atom) are necessary for the gap formation.

V. ELECTRONIC STRUCTURE

A. The Zintl viewpoint

Although in the “parent” perovskite system the structure is that of vertex-connected CoSb\textsubscript{6} octahedra, in the strongly distorted skutterudite structure the Sb\textsubscript{4} rings form a basic structural motif. These compounds, viz. CoSb\textsubscript{3}, are sometimes characterized as Zintl materials in which the Sb\textsubscript{4} unit balances the charge of the Co unit. Within the Zintl picture, the charge of the Co\textsuperscript{3+} ion must be balanced by a (Sb\textsubscript{4})\textsuperscript{4+/3−} unit: there are only 3/4 as many Sb\textsubscript{4} rings as Co atoms. Thus there is tension between the Zintl picture, which does not lead to integral formal charges on both of the primary units, and the presence of the gap that specifies an integral number of occupied bonding and antibonding states. Several papers\textsuperscript{14,23,24,29-35} have presented results and some analysis for empty and for a few filled skutterudites. We have benefited from the previous work as we proceed on a deeper analysis.

We first follow the commonly held line of reasoning that the short Sb-Sb distance in the Sb\textsubscript{4} ring is the most fundamental aspect of the electronic structure.\textsuperscript{35} The $p_x$-$p_x$ bonding along an $x$ axis is characterized by a hopping amplitude $t_{ppz}$ $\approx$ 3 eV, so the corresponding bonding and antibonding states lie at $\varepsilon_p \pm t_{ppz}$. The separation of Sa atoms and the $\sigma$ bonding between $p_x$ orbitals along the $y$ direction is indistinguishable from that of the $p_x$, $\sigma$ case. Indeed, the corresponding $p_x$ bonding and antibonding character (not shown in figures) is centered roughly $\pm$ 3–4 eV below and above the gap. The on-site energies $\varepsilon_p \approx 3\varepsilon_p = \varepsilon_p$ therefore lie in the vicinity of the gap. This picture leads to half-filled $p_x$ and $p_z$ orbitals.

The $p_z$ orbitals within an Sb\textsubscript{4} unit are orthogonal to the $p_x$, $p_y$ orbitals and couple by a $t_{ppz}$ hopping amplitude. This coupling leads to doubly degenerate levels at $\pm t_{ppz}$ relative to $\varepsilon_p$. The evidence is that there is negligible crystal field splitting of the three on-site $p$ orbitals. However, interunit $p_x$-$p_z$ interaction will be stronger than this intrunit coupling, and the $p_z$ orbitals also couple to the Co orbitals. The $p_z$ character, provided by the fatbands representation in Fig. 4 is in fact distributed throughout the $-6$ to $+5$ eV region and is not reproducible by a simple model due to other couplings. The $p_z$ projected density of states (DOS) indicates the $p_z$ orbitals are at least half-filled, perhaps slightly more.

B. Complications with the Zintl viewpoint

It is readily demonstrated that an analysis focusing primarily on the Sb\textsubscript{4} rings has strong limitations. We have performed calculations with the Co atom removed, \textsquare Sb\textsubscript{3}. The Sb $p$ band complex is broad (10 eV) and featureless. There is no hint of a gap or even pseudogap, indicating that the $p_x$ and $p_z$ bonding-antibonding feature described above is far from a dominant feature. Only when Co is reinserted into the structure does the remarkably clean 1 eV gap become carved out around the Fermi level, except of course for the peculiar linear band near the zone center. The gap therefore arises from the Co 3$d$ mixing (evidently strongly) with Sb 5$p$ orbitals, with the linear band emerging from the valence bands being of strong Sb character. Figure 5 illustrates the Co 3$d$ character. It lies mostly below the gap, in the very narrow $−1$ to $−2$ eV range. A much smaller amount of character, roughly something like four bands, lies immediately above the gap in the 0–1 eV range. These two parts of the 3$d$ projected DOS are neatly and impressively split by the 1 eV gap. The “charge state” (or formal valence, or oxidation state) of the Co atom is a question that can be asked, since this is an insulating material. If there are four unoccupied $d$ bands (though this cannot be claimed very conclusively due to the strong hybridization) then considering there are four Co atoms in the cell, the occupation would be $d^8$ Co\textsuperscript{1+}. A nonmagnetic ion with this filling is natural for the Co 3 site symmetry. As the CoSb\textsubscript{6} octahedron is distorted strongly in progressing from the cubic perovskite structure to the skutterudite one, the crystal field levels reduce as $t_{2g} \rightarrow e + b_2$, the former being the doublet, and $e_g \rightarrow a_1 + b_1$. One unoccupied orbital singlet (both spins) gives the $d^8$ occupation. The magnetic quantum number $m = 0$ orbital with respect to the local threefold axis is the natural one for the 3$d$ holes to occupy.

Charge balance will then leave the rings as (Sb\textsubscript{4})$^{−4/3}$, a very unsatisfactory result for accounting for the gap: each Sb\textsubscript{4}...
ring would not have an integer number of (fully) occupied bands (or molecular orbitals). Within the picture that the \( p_z \) and \( p_x \) orbitals are half-filled due to the large \( t_{pp} \) bonding-antibonding splitting, this Co charge state would suggest \( p_z \) orbitals to be 1/6 more than half-filled, a peculiar result and one that would only be compatible with a metallic, rather than semiconducting, state.

The integrated DOS for the Co 3d states in fact gives a clear \( d^8 \) occupation. However, with near-neutral atoms there might be significant Co 4s, 4p occupation, which could bring Co nearer to a neutral \( d^8s^1 \) configuration. Some Co sp character is in fact found in the upper valence bands, though it is possible that this represents tails of Sb \( p \) orbitals. The Co \( d \) occupation \( d^8 \) is clear, however.

C. Sb 6p-Co 3d mixing

It was noted above that when Co is removed from the structure, there is no gap nor any indication of one, nor any candidate for a potential linear band. The most direct interaction of the Sb \( p_z \) orbitals on the ring is between the \( p_z \) orbital on the nearest Sb in the ring with the rotated \( xz \) and \( yz \) orbitals on the eight Co neighbors. Sb\( p \) molecular orbitals (MOs) formed by linear combinations of \( p_z \) orbitals can be sorted according to whether they are even or odd parity with respect to the center of inversion at the Co site. The even parity MOs mix with the Co 3d orbitals, forming bonding and antibonding pairs tending to open a gap; the odd parity MOs do not mix with the 3d orbitals. As a result of this coupling, four of the five 3d orbitals are lowered and occupied, while one is raised and unoccupied, giving the 1 eV gap and the \( d^8 \) occupation.

D. Band character near the gap

The fatbands representation of Sb \( p_z \) character in Fig. 4 shows the substantial \( p_z \) character of the linear band in the upper valence bands near the zone center. This \( p_z \) character is in fact the dominant character, although various other small contributions from other valence orbitals are mixed in by the low Sb site symmetry. The character of the threefold degenerate conduction band minimum state is more complex. These bands have primarily Co \( \sim t_{2g} \) character; the quotes arise from the fact that the degeneracies of the cubic \( t_{2g} \) and \( e_g \) subshells are broken by the distortion of the octahedra in the skutterudite structure. There is a strongly \( d_{z^2} \) flat band just above (+0.2 eV) the gap, and additional strong \( d_{x^2−y^2} \) character in the 0.2–0.6 eV region.

The band figures we show below demonstrate that the linear band couples to only one of the triplet of conduction bands at and near \( \Gamma \), that is, couples to only one linear combination of the triplet states. It does not couple to the Co 3d orbitals.

E. A consistent viewpoint: An empty-site molecular orbital

A generalized viewpoint of the most physical orbitals in CoSb\( _3 \) is that of four Co\( ^{1+} \) \( d^8 \) atoms (as discussed above), three Sb\( _8 \) rings with \( p_{z^2} \) and \( p_{x^2−y^2} \) bonding states filled and antibonding states unoccupied, and molecular orbitals formed from the 12 \( p_z \) orbitals and centered on the empty 2\( a \) site. This site is an inversion center, so the MOs can be classified as parity even or odd. (Note that these are different MOs than were discussed in Sec. V C.) Twelve MOs can be formed, which can be classified also by the tetrahedral 2\( a \) site symmetry. Charge counting requires that eight of these MOs must be occupied (with both spins).

We will not attempt to identify the positions and dispersions of these MOs, except to note that linear band is one of these MOs. Jung et al.\(^{35} \) identified this MO as \( A_{2u} \), an orbitally nondegenerate state with odd parity at \( k = 0 \), and therefore not mixing with the Co 3d orbitals at (or near) \( \Gamma \). The density arising from this orbital is of interest. We had earlier investigated the possibility that an electron may reside in the large empty 2\( a \) site in the skutterudite structure. This type of “electride” configuration, in which an interstitial electron without a nucleus becomes an anion, is well established in molecular solids.\(^{36,37} \) However, the density at the 2\( a \) site is very small, ruling out this possibility.

An isosurface plot of the density arising from the linear band near \( \Gamma \) is shown in Fig. 6. The closed surface in the density encloses a region of low density centered on the unoccupied site. Exploring the density with isosurfaces at various values of density reveals no local maximum at this large interstitial site, thus no electride-like character. Lefebvre-Devos et al. have presented\(^{30} \) complementary isosurface density plots of this band, with the high density regions of two types. One region surrounds the empty site (with rather complex shape) consistent with its origin as a MO comprised of \( p_z \) Sb\( _8 \) ring states, with the tetrahedral symmetry of the 2\( a \) site. The second region is within each Co\( _8 \) octahedron, centered on either side of the Co atom along the local \( \bar{3} \) axis, appearing as a three-bladed propeller that arises from the lobes of the \( p_z \) orbitals that lie closer to the Co atom.
Interaction of the $p_z$ states with Co $3d$ orbitals appears to cause a majority of the density of this state to border the large empty site. The result is not an electrode state but rather a large molecular orbital (1 of 12 in total) centered on the empty site, with one of them ($A_{2u}$) per primitive cell. We have previously reported that this state at $\Gamma$ is parity-odd.\textsuperscript{10}

Such an orbital has the same hopping amplitude $t$ along each nearest neighbor direction on a $bcc$ lattice, hence its dispersion looks like that of an $s$ orbital, $E_k = 8t \cos(k, a/2) \cos(k, a/2) \cos(k, a/2)$, and does not lead to a linear band at $k = 0$. This should not be surprising, as the true linearity at $\Gamma$ is accidental, due to a degeneracy that has to be tuned. In our previous paper\textsuperscript{10} this tuning was accomplished by making two on-site energies in the tight-binding model identical.

The “linear band” aspect of the skutterudites may have been overinterpreted, because it arose in the most common member (the mineral skutterudite CoAs$_3$) which was studied for small Sb sublattice displacement (see Sec. II for the linear distortion, characterized by the parameter $s$), for this band to remain linear all the way to $k = 0$, marking a critical point. In that work, the progression of the band crossing at the critical point was plotted. Due to the threefold degeneracy of the conducting band edge, the progression versus $s$ was from semiconductor for $s \leq 1.19$, to the critical point at $s = 1.020$ with linear valence and conduction bands degenerate with two quadratic bands, to a zero-gap semiconductor for $s \geq 1.021$ due to a symmetry-determined degeneracy, even when spin-orbit coupling (SOC) is included. Due to the band inversion and the character of the states at $\Gamma$, the transition is from trivial insulator to one with topological character. Because the degeneracy at the point Fermi surface (at $\Gamma$) remains, the latter state is a topological zero-gap semiconductor rather than a topological insulator.

We now demonstrate that a simple tetragonal distortion, such as might result from growth on a substrate with small lattice mismatch, does produce a topological insulator (TI) but with an unexpected placement of the gap. The progression of the band structure very near $\Gamma$ is shown in Fig. 7 for strain $c/a = 1.01$. The gap before the critical point is reached always has quadratic bands for sufficiently small $k$. At the critical point $s = 1.020$ (when the strain is imposed) leads, unexpectedly, to linear bands only in certain of the high symmetry directions (such as the $(111)$ direction, toward the $P$ point). In other high symmetry directions (from $\Gamma$ toward $H$ is pictured) the bands are quadratic and the upper one is quite flat (heavy mass). We return to this point in the next subsection.

Beyond the critical point, the four bands become nondegenerate once more; the lower (linear) band has crossed the lowest of the three conduction bands and acquired mixed character. The coupling between bands is clearly much smaller along the axes ($\Gamma$-$H$ direction) than along the other symmetry lines, and the gap occurs along this direction. Because of the heavy mass of the lower conduction bands, the gap is quite small, being of the order of 1 meV for the tetragonal strain that is pictured in Fig. 7.

B. Anisotropy at the critical point

Inspecting the bands along the three cubic high symmetry directions, a second peculiar feature becomes evident. At the critical point, the “linear band” disperses linearly along the $(111)$ directions, while they are quadratic (with distinct masses) along both the $(100)$ and $(110)$ directions. The directional dependence as well as the magnitude dependence of the wave vector is anomalous. The form of the dispersion for small $|k|$ can be constructed. A polynomial in the direction of $\hat{k}$ (\hat{k}) with cubic symmetry that vanishes along the $(100)$ directions can be constructed as

$$P_{100}(\hat{k}) = \sqrt{\prod_{j=1}^{6} (\hat{k} - \hat{k}_j)^2}$$

$$= \prod_{j=1}^{6} |\hat{k} - \hat{k}_j|,$$

(2)

where $\{\hat{k}_j\}$ are the six $(100)$ unit vectors. The denominator normalizes the polynomial to unity along each of the $(111)$ directions; $k_{111}$ lies along the $(111)$ direction (any of them).
C. Isovalent skutterudite antimonides

We have studied the band structures of antimonide sisters of CoSb$_3$, based on the transition metal atoms Rh and Ir. Band structures, or sometimes only gap values, of these compounds have been reported earlier.$^{25,31,38}$ The band structures around the gap at $k = 0$ differ somewhat in the literature, due to sensitivity to structure (and that some use relaxed coordinates while others use the measured structure), to the level of convergence of the calculation, and the exchange-correlation functional.

The difference in lattice constant is compensated very closely by the effect of difference in interatomic distances on hopping amplitudes, and the resulting bands are very similar. The single feature of interest is the magnitude of the gap (and whether positive or negative, the latter giving an inverted band structure) and the strength of SOC effects. As mentioned above, band inversion alone does not in itself give rise to a topological insulator state in these compounds because even after SOC is included, a twofold degeneracy at $\Gamma$ is lowest and the system is a point Fermi surface, zero-gap semiconductor. Such cases are referred to as topological semimetals. However as shown above, the topological semimetal can be driven into the TI phase by tetragonal distortion that lifts the final degeneracy.

$\text{RhSb}_3$. For a range of volumes around the experimental one ($a = 9.23$ Å), this compound has an inverted band structure, hence it is a point Fermi surface, zero-gap topological semimetal. In the inverted band structure before considering SOC, the singlet lies 23 meV above the triplet that pins the Fermi level. Inclusion of SOC leaves the Fermi energy pinned to a doublet. A tetragonal distortion raises the degeneracy; a topological insulator is obtained for $c/a > 1$ in the same way that happens for CoSb$_3$ although no sublattice strain is required in RhSb$_3$.

Generally, reducing volume (applying pressure) in these Sb-based skutterudites leads to a larger gap. In the case of RhSb$_3$, our calculations indicate a gap never actually opens within cubic symmetry, for peripheral reasons. When the band inversion at $\Gamma$ is undone and a gap at $\Gamma$ appears at sufficiently high pressure ($a = 9.06$ Å), the material is metallic due to other (Rh 4$d$) bands having become lowered in energy, crossing the Fermi level along the $\Gamma$-$N$ direction.

$\text{IrSb}_3$. The larger SOC for heavier atoms led us to consider this 5$d$ compound. However, the linear band and several others have practically no metal atom character and are not affected by its spin-orbit strength. We obtain, using the same computational methods as for the other two compounds, that like CoSb$_3$, IrSb$_3$ has a $\sim$80 meV gap and therefore no topological character at the experimental equilibrium structure. Like CoSb$_3$, it can be driven to an inverted band structure and hence topological character by internal and tetragonal strains.

VII. SUMMARY

One goal of this work was to provide a simple but faithful picture of the electronic structure of CoSb$_3$: the origin of the gap and the character of the linear band being the most basic. The picture is this. Co is in a d$^8$ configuration, leaving four electrons in the Co$_4$Sb$_{12}$ unit cell to go into other bands. The
$p_z$ and $p_y$ orbitals on the Sb$_2$ ring form strongly bonding (occupied) and antibonding (unoccupied) states, accounting for two of the three $5p$ electrons of each Sb atom, for a total of 12 occupied $\sigma$-bonding states (of each spin). The 12 $\pi$-oriented $p_z$ orbitals couple between units as well as with the Co 3$d$ orbitals, and form “molecular orbitals” centered on the vacant 2$a$ sites in the skutterudite lattice. Eight of these MOs are occupied by 16 electrons, one from each Sb and one from each Co. The linear band is best pictured as arising from the coupling and dispersion of these MOs.

We have also justified our earlier statement$^{10}$ that CoSb$_3$ is very near a conventional-to-topological transition. A small symmetry-preserving internal strain, a small applied tetragonal strain, and spin-orbit coupling drives CoSb$_3$ into a topological insulator phase. This critical point also marks a point of accidental (but tunable, by the strains) degeneracy that gives rise to emergence of linear Dirac-Weyl bands emanating from $k = 0$, where they are degenerate with massive bands. At this critical point, the dispersion is nonanalytic at $k = 0$ and anisotropy is extreme, with the mass varying from normal to vanishing. In this respect CoSb$_3$ at its critical point bears several similarities to the semi-Dirac (Dirac-Weyl) behavior encountered$^{39–41}$ in ultrathin layers of VO$_2$.

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12occupied and antibonding (unoccupied) states, accounting for two of the three $5p$ electrons of each Sb atom, for a total of 12 occupied $\sigma$-bonding states (of each spin). The 12 $\pi$-oriented $p_z$ orbitals couple between units as well as with the Co 3$d$ orbitals, and form “molecular orbitals” centered on the vacant 2$a$ sites in the skutterudite lattice. Eight of these MOs are occupied by 16 electrons, one from each Sb and one from each Co. The linear band is best pictured as arising from the coupling and dispersion of these MOs.

We have also justified our earlier statement$^{10}$ that CoSb$_3$ is very near a conventional-to-topological transition. A small symmetry-preserving internal strain, a small applied tetragonal strain, and spin-orbit coupling drives CoSb$_3$ into a topological insulator phase. This critical point also marks a point of accidental (but tunable, by the strains) degeneracy that gives rise to emergence of linear Dirac-Weyl bands emanating from $k = 0$, where they are degenerate with massive bands. At this critical point, the dispersion is nonanalytic at $k = 0$ and anisotropy is extreme, with the mass varying from normal to vanishing. In this respect CoSb$_3$ at its critical point bears several similarities to the semi-Dirac (Dirac-Weyl) behavior encountered$^{39–41}$ in ultrathin layers of VO$_2$.

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