Band Connectivity for Topological Quantum Chemistry: Band Structures As A Graph Theory Problem

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The conventional theory of solids is well suited to describing band structures locally near isolated points in momentum space, but struggles to capture the full, global picture necessary for understanding topological phenomena. In part of a recent paper [Bradlyn et al., Nature 547, 298–305 (2017)], we have introduced, the way to overcome this difficulty by formulating the problem of sewing together many disconnected local $k \cdot p$ band structures across the Brillouin zone in terms of graph theory. In the current manuscript we give the details of our full theoretical construction. We show that crystal symmetries strongly constrain the allowed connectivities of energy bands, and we employ graph-theoretic techniques such as graph connectivity to enumerate all the solutions to these constraints. The tools of graph theory allow us to identify disconnected groups of bands in these solutions, and so identify topologically distinct insulating phases.

I. INTRODUCTION

The fundamental assumption of the textbook approach to the theory of solids is that one reaps enormous benefit by trading the dependence of wavefunctions on real-space position for a dependence on crystal momentum $k$. Through Bloch’s theorem, this has led to the development of the $k \cdot p$ expansion, allowing for tractable approximations to material Hamiltonians near isolated points in momentum space. This approach found extraordinary successes throughout the twentieth century in the understanding and predicting the behavior of semiconductors, metals (and their Fermi surfaces), and insulators.

However, an essential shortcoming of this approach is that it is local in momentum space. This renders obscure the global properties of wavefunctions in momentum space. It is for precisely this reason that topological insulators appear foreign in the $k \cdot p$ approach to solid state physics. Isolated $k \cdot p$ Hamiltonians at various points in the Brillouin zone (BZ) are a priori completely independent, and so extracting global (topological) data from such a local momentum space picture seems hopeless at first sight. Such global data could, if existent, be used to characterize all possible band structures (materials) in nature. In particular, one useful outcome of such a characterization would be that it contains both topologically trivial and topologically non-trivial band-structures.

In a part of a recent paper[11], we have introduced, amongst several other concepts, a new way of providing such a global classification of band structures using graph theory, thereby updating band theory with its last missing ingredient. In this paper, we fill in the mathematical details necessary for a full and complete description of our theory. We map the problem at hand – patching together isolated $k \cdot p$ expansions into consistent global band structures – to a tractable problem in graph theory. We first re-interpret the $k \cdot p$ expansion as a representation subducing technique, and identify representations with the nodes of a ”band” graph. We then show that band structures consistent with the symmetries of a crystal can be put in one-to-one correspondence with graphs that incorporate the symmetries as constraints on how different edges can be joined together. We show that there are often multiple distinct allowed graphs, with different numbers of connected components corresponding to (disconnected) groups of energy bands. In tandem with results of Refs. [11] and [2] we will argue that these different connectivities correspond to topologically distinct phases. Thus, we will show that the physics of topological insulators can be captured by the connectivity of band structures, without the need to invoke the tools of differential geometry. This generalizes the known eigenvalue- and $K$-theory based approaches to computing topological indices[8,9], unifying them all through the lens of graph theory.
We will be interested in the behavior of \( H \) near some high-symmetry point \( k_0 \) in the BZ. Let us introduce the little group \( G_{k_0} \) of \( k_0 \), defined as

\[
G_{k_0} \equiv \{ R[t] \in G | Rk_0 = k_0 + K \},
\]

where \( K \) is a reciprocal lattice vector. This is the subgroup \( G_{k_0} \subset G \) of the space group which leaves \( k_0 \) invariant up to a reciprocal lattice vector. The little group plays a privileged role in constraining the \( k \cdot p \) theory near \( k_0 \). Restricted to the little group, the representation \( \Delta \) subduces to

\[
\Delta_{k_0} \equiv \Delta \downarrow G_{k_0} = \bigoplus \rho_i,
\]

where the \( \rho_i \) are irreducible representations acting on vector spaces \( V_i \). Using Eq. (4) we see that

\[
[\Delta_{k_0}(g), H(k_0)] = 0
\]

for all \( g \in G_{k_0} \): the matrix representatives of the elements of \( G_{k_0} \) commute with \( H(k_0) \). Schur’s Lemma then tells us that the matrix \( H(k_0) \) in the representation space of \( \Delta_{k_0} \) is a sum of constants \( \epsilon_i \) for each irrep \( \rho_i \) in the decomposition Eq. (5).

\[
H(k_0) = \bigoplus_i \epsilon_i P_{\rho_i},
\]

where \( P_{\rho_i} \) is the projector onto the representation space \( V_i \) of \( \rho_i \). For those representations which occur with multiplicity larger than one, Schur’s Lemma implies only that the Hamiltonian is block diagonal; we assume here that we have carried out any additional diagonalization needed to put the Hamiltonian in the form Eq. (5). Thus, electronic states at \( k_0 \) come in degenerate sets of energies \( \epsilon_i \) and degeneracy given by the dimension of the representations \( \rho_i \).

We thus see that at the high-symmetry point \( k_0 \) the structure of the Hamiltonian is almost trivial. Things become more interesting, however, when we look slightly away from \( k_0 \). Here the representatives of the little group \( G_{k_0} \) no longer commute with \( H(k) \), although they still place strong constraints. Writing \( k = k_0 + \delta k \), we may expand \( H(k) \) in powers of \( \delta k \)

\[
H(k) = H(k_0) + \delta k^\mu H^{(1)}_{\mu} + \delta k^\mu \delta k^\nu H^{(2)}_{\mu \nu} + \ldots,
\]

where we have introduced the matrix-valued expansion coefficients \( H^{(i)} \). We use \( \mu, \nu = 1, 2, \ldots, D \) to index the primitive basis directions in the BZ, and repeated indices are summed; \( H(k_0) \) is given by Eq. (5). We now observe that, if the energy spacings \( \epsilon_i - \epsilon_j \) are the largest scales in the problem, which is always the case for small \( \delta k \), the matrices \( H^{(n)} \) appearing in the expansion Eq. (6) are approximately block diagonal in the carrier space \( \bigoplus_i V_i \) of irreducible representations of \( G_{k_0} \). More precisely, we may write in perturbation theory

\[
H^{(n)}_{\mu_1 \mu_2 \ldots \mu_n} = \bigoplus_i H^{(n)}_{\mu_1 \mu_2 \ldots \mu_n} P_{\rho_i} + \mathcal{O}(\delta \epsilon^{-1}),
\]
where $\delta \epsilon = \min_{i \neq j} (|\epsilon_i - \epsilon_j|)$. Note that although each $H^{(\alpha)}$ acts only within the space $V_\alpha$, they are not diagonal matrices within this space; rather from Eq. (1) we deduce that for each $g \in G_{k_0}$

$$
\rho_i(g)H^{(\alpha)}_{\mu_1\mu_2...\mu_n} \rho_i(g^{-1})\delta k_{\mu_1} \delta k_{\mu_2} ... \delta k_{\mu_n} = H^{(\alpha)}_{\mu_1\mu_2...\mu_n}(g\delta k)_{\mu_1}(g\delta k)_{\mu_2} ... (g\delta k)_{\mu_n},
$$

The practical consequence of this is that the momentum representations (those in which lattice translations are pure lattice translations) are no longer diagonal in the $k$-space, but rather from representations of $G_{k_0}$ (as we will explore further in Sec. III A). Because the representations $\rho_i$ of the little group $G_{k_0}$ play a special role in the theory of little group representations, we introduce the double groups, which are subduced from representations of $G_{k_0}$, (as we will see in the example in Sec. II C). The double groups are central extensions of the space group, and for spinless particles, a rotation by $2\pi$ should leave the wavefunction invariant, while for spin-1/2 particles, a rotation by $2\pi$ multiplies the wavefunction by $-1$. To accomodate this in the theory of little group representations, we introduce the double space groups and their representations.

The double groups are central extensions of the space group, obtained by adjoining an element $E$ which commutes with all elements of $G_k$. This element signifies a $2\pi$ rotation, and so we extend the groups so that every $n$-fold rotation, when raised to the $n$-th power, yields $E$. Representations of the double groups are termed single- or double-valued based on their value on $E$. In particular, we have

$$
\rho(E|\{R|d\}) = \begin{cases} \rho(|R|d), & \text{\sim single-valued} \\ -\rho(|R|d), & \text{\sim double-valued}. \end{cases}
$$

For the remainder of this work, we will focus solely on the double groups; we will follow the accepted convention of distinguishing double-valued representations with the use of an overbar. Finally, we will often be interested in systems with time-reversal (TR) symmetry. Space groups with time reversal symmetry contain an additional antiunitary element $T$, which commutes with all other elements of the space group, leaves real-space (direct space) invariant, and maps $k$ to $-k$. With TR, it is important to distinguish between three different types of $k$-vectors and little groups.

First, there are those vectors $k_0$ such that $-k_0 \equiv k_0$. In this case, $T \in G_{k_0}$. In spinless representations $\rho$ of
As an example of the above ideas, let us examine the $\mathbf{k} \cdot \mathbf{p}$ Hamiltonian for two closely related cubic crystals: the symmorphic group $I432$ (211), and the related nonsymmorphic group $I4132$ (214), which share the same point group. We take as a basis for the BCC lattice the three vectors

$$
\mathbf{e}_1 = \frac{1}{2}(-\hat{x} + \hat{y} + \hat{z})
$$

$$
\mathbf{e}_2 = \frac{1}{2}(\hat{x} - \hat{y} + \hat{z})
$$

$$
\mathbf{e}_3 = \frac{1}{2}(\hat{x} + \hat{y} - \hat{z}).
$$

(11)

Both space groups have point group $O$ (432), the octahedral group, consisting of all orientation-preserving symmetries of the cube. However, in $I4132$, the fourfold rotation $C_{4z}$ about the $z$-axis is embedded into the space group as a screw rotation $C_{4z}[00\frac{1}{2}]$, a difference

$$
H^{(211)}(\delta \mathbf{k}) = H^{(214)}(\delta \mathbf{k}) \approx c_0 \mathbb{I} + \begin{pmatrix}
\null & \null & 0 \\
0 & -a \delta k_z & -b \delta k_z \\
\frac{\sqrt{3}}{4} (a-b) \delta k_+ & \frac{\sqrt{3}}{4} (a-b) \delta k_- & 0 \\
\end{pmatrix},
$$

(13)

in terms of two real-valued phenomenological parameters $a$ and $b$, and where we have defined $\delta k_{\pm} = \delta k_x \pm i \delta k_y$. This describes a fourfold degeneracy at $\Gamma$ that disperses linearly, and corresponds to the “spin-3/2” fermion introduced in Ref. [35].

Looking next at the $P$ point, we find a very different situation. Without TR, the little co-group $\overline{G}_P \approx T$ of $P$ in both cases is generated by a $C_3$ rotation about the cubic body-diagonal, and a twofold rotation $C_{2z}$. With TR, in both cases the little co-group is augmented by the antiunitary operation $\mathcal{A} = C_{2z} g T$, which maps the $P$ point to itself. The little co-group $\overline{G}_P$ both with and without $\mathcal{A}$ has only two-dimensional spinful representations, since it contains the Pauli group of $C_{2z}, C_{2y}, C_{2z}, E$, and their squares (a fourfold rotation is needed to make the spin-3/2 representation irreducible). For the following, we will consider a TR-invariant system, so that the little co-group contains $\mathcal{A}$. For SG $I432$ the representations of the little co-group determine uniquely the representations of the little group acting on the $\mathbf{k} \cdot \mathbf{p}$ Hamiltonian. Focusing still on $J = 3/2$ states, we then have for the irreps of the $4 \times 4$ representation, which is conventionally denoted $\Gamma_8$ in both space groups. 

In both of these SGs, the little co-group $\overline{G}_P$ of the $\Gamma$ point is given by the point group $O$, while the little co-group $\overline{G}_P$ of the $P$ point is the tetrahedral group $T$, the subgroup of $O$ obtained by removing all fourfold rotations, as well as removing the twofold rotations along the diagonals.

Let us first examine the Hamiltonian near the $\Gamma$ point. As remarked above, by Eq. (9), the representations matrices for symmetry elements corresponding to the same point group operation in the little group of $\Gamma$ are identical for both space groups $I432$ and $I4132$, and follows trivially from the representation theory of the little co-group $\overline{G}_P \approx O$. The spinful irreps of the octahedral group are obtained directly from subduction of the half-integer spin-$J$ irreps of $SU(2)$, and the representation spaces are spanned by $\{\ell = 0, J = \frac{1}{2}\}, \{\ell = 1, J = \frac{1}{2}\}$ or $\{\ell = 1, J = \frac{3}{2}\}$ basis functions. We focus on four bands transforming in the $J = \frac{3}{2}$ representation, which is conventionally denoted $\overline{\Gamma}_8$ in both space groups.

Enforcing the symmetries on the four-band $\mathbf{k} \cdot \mathbf{p}$ Hamiltonian in this representation yields in both cases

$$
\begin{align*}
\mathbf{g}_1 &= \frac{2\pi}{a}(0, 1, 1), \quad \mathbf{g}_2 = \frac{2\pi}{a}(1, 0, 1), \quad \mathbf{g}_3 = \frac{2\pi}{a}(1, 1, 0),
\end{align*}
$$

(12)

where

$$
\begin{align*}
\mathbf{g}_1 &= \frac{2\pi}{a}(0, 1, 1), \quad \mathbf{g}_2 = \frac{2\pi}{a}(1, 0, 1), \quad \mathbf{g}_3 = \frac{2\pi}{a}(1, 1, 0).
\end{align*}
$$
\[H^{(211)}(k_P + \delta k) \approx \begin{pmatrix} 
\epsilon_1 + v_1 \delta k \cdot \sigma & 0 \\
0 & \epsilon_2 + v_2 \delta k \cdot \sigma 
\end{pmatrix},\]

where \(\sigma\) is the vector of Pauli matrices. This describes a pair of twofold degeneracies at \(k_P\) with different energies that disperse linearly away from \(k_P\), where \(\epsilon\) is a pair of Weyl fermions at energies \(\epsilon_1\) and \(\epsilon_2\).

Turning now to SG \(I4_132\), we find a completely different phenomenon. Although the little co-group \(G_P\) is unchanged, the non-symmorphic character of the space group manifests itself in the little group \(G_P\): this group is generated by the rotation \(\{C_3[000]\}\) about the cubic body diagonal, and also the twofold screw rotation \(\{C_{2x}[\frac{1}{2} 0 \, 0]\}\); these two generators fully determine the extension of \(G_P\) by the group of lattice translations. In any representation \(\rho\) of \(G_P\) we must have

\[\rho(C_2x[\frac{1}{2} 0 0]^2) = \rho(E[011]) = e^{-i\pi} \rho(E[000]).\]

We recognize the three-band lower-right block of the Hamiltonian as the “spin-1/2” Weyl Hamiltonian of Ref. 38. From this equation, we see that for spin-1/2 systems, the representative of the \(C_{2x}\) screw in the little group \(G_P\) squares to \(+I\). In effect, double-valued representations of the little group \(G_P\) are isomorphic to single-valued representations of the little co-group \(G_P\). Thus, as was discussed extensively in Sec. I of the Supplementary Material of Ref. 38, bands at the \(P\) point in SG \(I4_132\) transform in the sum of a one dimensional representation \(\mathcal{P}_4\) and a three dimensional representation \(\mathcal{P}_7^{(\text{NS})}\) of \(G_P\); we have added the superscript “(NS)” to distinguish this representation from the 2D representation in space group \(I4_32\). For the \(k \cdot p\) expansion for the four bands considered here, this yields to linear order

\[H^{(214)}(k_P + \delta k) \approx \begin{pmatrix} 
\epsilon_1 & 0 & 0 & 0 \\
0 & \epsilon_2 & a\delta k_x & \ast \delta k_y \\
0 & \ast \delta k_x & \epsilon_2 & 0 \\
0 & \ast \delta k_y & 0 & \epsilon_2 
\end{pmatrix}.\]

We can see from this example that the degeneracies and structure of local Bloch Hamiltonians depends strongly on the symmorphicity of the crystal symmetry group. This is also expected to be the case at points in between \(\Gamma\) and \(P\). At this point, however, we have no way of connecting the expanded \(k \cdot p\) Hamiltonians at \(\Gamma\) and \(P\) to form a consistent global band structure. In the next section, we will develop the tools necessary to do so.

### III. Patching Together \(k \cdot p\) Hamiltonians: Global Band Structures

In order to patch together the local band structures obtained from the \(k \cdot p\) expansions at various points, we will make use of the constraints imposed by crystal symmetry. Having just reformulated the traditional \(k \cdot p\) theory in terms of group representations, we will be able to show how the compatibility between little group representations along high-symmetry lines and planes strongly constrains the allowed connections between bands near different \(k\) points. In order to solve these constraints, we will map the problem to a problem in graph theory.

#### A. Compatibility relations

To begin, we examine the symmetry constraints on energy bands at different \(k\)-vectors \(k_1\) and \(k_t\), where the little group \(G_{k_t}\) is a proper subgroup of the little group \(G_{k_1}\). This occurs, for instance, when \(k_1\) is a high-symmetry point in the BZ, and \(k_t\) is a point on a high-symmetry line with endpoint \(k_1\). For instance, in the example of SG \(I4_32\) in Sec. [1C] we could take \(k_1\) as the \(\Gamma\) point, and \(k_t = (k_1, k_t, k_t)\), which lies on the high-symmetry line \(\Lambda\), with little co-group generated only by the \(C_3\) rotation about the cubic body diagonal. In all such cases, the relation \(G_{k_t} \subset G_{k_1}\) implies that each irrep of \(G_{k_t} \subset G_{k_1}\) restricts to (subduces) a direct sum of (generally more than one) irreps \(\bigoplus_i \sigma_i\) of \(G_{k_t}\), which we denote by

\[\rho \downarrow G_{k_t} \approx \bigoplus_i \sigma_i.\]

The restriction is obtained by removing from \(\rho\) the matrices for those elements not in \(G_{k_t}\), and viewing the group representation formed by the remaining matrices as a (in general reducible) representation of \(G_{k_t}\).

At this point in the argument, we remark that there exists a whole manifold (line, or plane) of \(k\)-vectors with little group \(G_{k_t}\), which we denote \(\{k_t\}\). In the example of SG \(I4_32\) for instance, all \(k\) points of the form \(\{k_t\} = \{\frac{1}{2} (g_1 + g_2 + g_3), t \in [0,1]\}\) have the same little group \(G_{\Lambda}\) (the endpoints have the larger little groups \(G_{\Gamma}\) and \(G_P\), both of which contain \(G_{\Lambda}\)). In these situations, the restriction \(\rho \downarrow G_{k_t}\) of little group representations holds along the entire manifold of points \(\{k_t\}\). This has two main consequences for \(k \cdot p\) Hamiltonians. First, we know that the \(k \cdot p\) Hamiltonian \(H(k + \delta k)\) around point \(k_t\) transforms according to the same representation (and therefore takes the same form) along the entire manifold \(\{k_t\}\). Second, we know that if we focus on a Hamiltonian near \(k_1\) transforming in some representation \(\rho\) of \(G_{k_1}\), we know that for \(k_1 + \delta k_t \in \{k_t\}\), the Hamiltonian \(H(k + \delta k_t)\) must transform in the restricted representation \(\rho \downarrow G_{k_t}\). Thus, the compatibility relations Eq. (17) between representations of the little
groups $G_{k_1}$ and $G_{k_2}$ constrains the representation space of Bloch functions that may appear in a band structure.

B. Global band structures

The compatibility of $k \cdot p$ band structures gives us our first clue on how to piece spectra at different $k$ points together: along lower-symmetry manifolds emanating from high-symmetry $k$-vectors, the representations of electronic states appearing in the Hamiltonian must be compatible. We can complete the picture by noting that the closure of manifolds $\{k_1\}$ of points with little group $G_{k_1}$ contain more than one high-symmetry $k$-vector $k_1$ and $k_2$, namely the points at the boundary of $\{k_1\}$. For example, in our cubic crystal, with $G_{k_1} = G_1$, we have that $k_1$ is the $\Gamma$ point, and $k_2$ is the $P$ point. Not only do we have $G_{k_1} \subset G_{k_2}$, but also $G_{k_2} \subset G_{k_1}$. This allows us to patch together $k \cdot p$ expansions about the points $k_1$ and $k_2$, along the manifold $\{k_i\}$. If we have a Hamiltonian near $k_1$ that transforms according to a representation $\rho_1$ of $G_{k_1}$, and a Hamiltonian near $k_2$ that transforms according to a representations $\rho_2$ of $G_{k_2}$, then these can be consistently connected only if the representations

$$\rho_1 \downarrow G_{k_1} \approx \rho_2 \downarrow G_{k_2} \approx \bigoplus_i \sigma_i$$

restricted to the manifold $\{k_i\}$ are equivalent. Furthermore, along the manifold $\{k_i\}$ we can trace bands carrying each representation $\sigma_i$ from an irrep of $G_{k_1}$ at $k_1$ all the way to an irrep of $G_{k_2}$ at $k_2$.

Extending this logic to all $k$-vectors in the BZ, we can say that patching together $k \cdot p$ Hamiltonians consistently requires ensuring that the set of little group representations appearing in the spectrum at each $k$-vector manifold are compatible. Even once this is ensured, however, there still may be an ambiguity on how bands join together. This happens when there is a representation $\sigma_n$ that appears in the subductions $\rho_1 \downarrow G_{k_1}$ and $\rho_2 \downarrow G_{k_2}$ with multiplicity $m_n > 1$. In this case, multiple bands along the the manifold $\{k_1\}$ carry the representation $\sigma_n$. If these bands are nondegenerate at both $k_1$ and $k_2$, then there are different inequivalent ways of connecting the band structure consistent with the symmetries. We illustrate this schematically in Fig. 1 where the representation $\sigma_n$ appears with multiplicity 2, allowing for the two distinct connectivities shown in (a) and (b). More complicated examples quickly arise in other space groups.

We thus seek a method to find all distinct ways of patching together local band structures to form global spectra, consistent with the compatibility constraints of crystal symmetry. In order to do this efficiently, we will map this to a graph theory problem: vertices in the graph will correspond to irreps of the little group at each symmetry distinct (manifold of) $k$-vector(s), and two vertices can be joined by an edge only if the corresponding bands connect (implying that the representations are compatible). We will formalize this in the following section, and show how it allows us to fully classify and enumerate allowed band structures.

IV. CONSTRUCTING CONNECTIVITY GRAPHS

In order to map the problem of finding global band structures to graph theory, we first introduce some basic terminology and concepts in Sec. IV A. We then define in Def. 1 the precise mapping from band structures to “connectivity graphs”, and show how global band structures can be constructed by analyzing these graphs one subgraph at a time. We then revisit our cubic crystal example from Sec. IV C and show how to patch together the $k \cdot p$ band structures at the $\Gamma$ and $P$ points in both the symmorphic (I432) and non-symmorphic (I4132) cases. Finally, we show how spectral graph theory allows us to analyze the connectivity of global band structures directly in graph-theoretic language.

A. Review of graph theory

We start by reviewing some basic notions about graphs which will be useful throughout the text. A complete treatment of graph theory can be found in Refs. 40 and 41.

A graph $\mathcal{G}$ consists of two sets: a set of nodes (or vertices) $N(\mathcal{G})$, and a set of edges $E(\mathcal{G})$. Each edge $e \in E(\mathcal{G})$ connects two distinct nodes $n_1, n_2 \in N(\mathcal{G})$. If two nodes $n_1$ and $n_2$ are connected by $\ell > 0$ edges, we label the edges as $(n_1, n_2, \alpha)$, where $\alpha \in \{1, \ldots, \ell_{n_1n_2}\}$.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{Two distinct allowed global band structures along the line $k_1 \leftrightarrow k_2$. $\rho_1$ and $\rho_1^2$ are two distinct representations of $G_{k_1}$, and $\rho_2$ and $\rho_2^2$ are two distinct representations of $G_{k_2}$. All four of these representations subduce the representation $\sigma_n$ of the little group $G_{k_n}$ of the line $\{k_i\}$. Because of this, global energy bands may be connected in two ways. In the band structure (a), the states transforming in representation $\rho_1^1$ at $k_1$ connect to the states transforming in the representation $\rho_2^1$ at $k_2$. Contrarily, in (b), the states transforming in representation $\rho_1^2$ at $k_1$ connect to the states transforming in the representation $\rho_2^2$ at $k_2$.}
\end{figure}
(note that if \(\ell_{n_1n_2} > 1\), the graph contains \(\ell_{n_1n_2} - 1\) loops connecting nodes \(n_1\) and \(n_2\)); furthermore, since \(n_1\) and \(n_2\) are by hypothesis distinct, our graphs do not contain self-loops. We will be exclusively concerned with undirected graphs, meaning that the edges \((n_1, n_2, \alpha)\) and \((n_2, n_1, \alpha)\) are not distinguished (the edges do not have directional "arrows" associated to them). In Fig. 2 we show a pictorial representation of a graph with eight nodes, labelled \(n_1\) through \(n_8\), and denoted by blue diamonds, black circles, and a red square. This graph also has eight edges. In particular, nodes \(n_7\) and \(n_8\) are connected by two edges labelled \((n_7, n_8, \alpha)\), with \(\alpha \in \{1, \ell_{n_7n_8} = 2\}\).

A partition of a graph is a subset \(P_1 \subset N(\mathcal{G})\) of nodes of \(\mathcal{G}\) such that for all \(p, q \in P_1\), \((p, q, \alpha) \notin E(\mathcal{G})\). In words, this says that a partition is a subset of nodes such that no two nodes in the subset are connected by edges. Note that in any graph, each node lies in a trivial partition containing itself only (and so the decomposition of a graph into partitions is not unique).

Finally, a subgraph \(H \subset \mathcal{G}\) of a graph \(\mathcal{G}\) is a graph such that the set of nodes \(N(H) \subset N(\mathcal{G})\), the set of edges \(E(H) \subset E(\mathcal{G})\), subject to the restriction that for each edge \((p, q, \alpha) \in E(H)\), we have \(p, q \in N(H)\). In words, a subgraph of a graph consists of a subset of nodes, and a subset of edges connecting those nodes only. In the graph shown in Fig. 2 for example, one subgraph \(H\) has \(N(H) = \{n_1, n_2, n_9\}, E(H) = \{(n_1, n_4, 1), (n_2, n_4, 1)\}\). This subgraph is called a 2-partite (or bipartite), since all nodes can be placed into the two partitions \(P_1 = \{n_1, n_2\}\) and \(P_2 = \{n_4\}\). Note also that in general a subgraph containing nodes \(n_1\) and \(n_2\) need not contain all the edges connecting these two nodes. However, for our purposes, we will only make use of subgraphs \(H \subset \mathcal{G}\) which contain all edges from \(E(\mathcal{G})\) connecting nodes in \(N(H)\).

We now show that band structures in momentum space map to N-partite graphs, where the \(N\) partitions correspond to high-symmetry \(k\) - vectors in the BZ. Groups of bands isolated in energy from all others will map to connected components of a graph \(\mathcal{G}\) - subgraphs whose points are not connected to other nodes in the graph by any edges. Mathematically, we define subgraphs \(H \subset \mathcal{G}\) such that for all \(h_i \in N(H)\) and all \(g_i \in N(\mathcal{G}) \setminus N(H), (h_i, g_i, \alpha) \notin E(\mathcal{G})\). For example, the graph shown in Fig. 2 has two connected components \(H_1\) and \(H_2\), where \(N(H_1) = \{n_6, n_7, n_8\}\) and \(E(H_1) = \{(n_6, n_7, 1), (n_6, n_8, 1), (n_7, n_8, 1), (n_7, n_8, 2)\}\); \(H_2\) contains all the remaining nodes and edges. We will refer to each connected component of a graph as a disconnected subgraph.

**B. Mapping to graph theory**

We are now in a position to map the problem of forming global band structures to a graph theory problem. We start with a space group, and a set of little group representations at high-symmetry \(k\)-vectors. We will define a connectivity graph, such that each valid connectivity graph yields a solution of the compatibility relations. First, we identify all symmetry-distinct \(k\)-vector manifolds throughout the BZ. For symmorphic groups this list corresponds with the list of \(k\)-vectors found on the Bilbao Crystallographic Server\(^{33}\). For non-symmorphic groups, there is the additional subtlety that the little group representation according to which a Bloch function transforms may change upon following an energy band smoothly through a full cycle across the BZ\(^{43}\). The details of how we identify \(k\)-manifolds in the non-symmorphic case can be found in Ref. 43. Each \(k\)-manifold will label a partition of our connectivity graph. In each partition \(k_i\) (where the index \(i\) runs over different \(k\)-manifolds in the BZ), we will place a node labelled by the irreps \(\rho_i^a\) of the little group \(\Gamma_{k_i}\) which appear in the \(k \cdot p\) expansion. We allow edges between nodes in partitions \(k_i\) and \(k_j\) only if either \(\Gamma_{k_i} \subset \Gamma_{k_j}\), or \(\Gamma_{k_j} \subset \Gamma_{k_i}\); these edges represent the continuation of energy bands from a high-symmetry \(k\)-vector to a lower symmetry \(k\)-manifold. Finally, we would like to enforce the compatibility relations as a constraint on allowed edges. In particular, given a pair of partitions \(k_i\) and \(k_i\) such that \(\Gamma_{k_i} \subset \Gamma_{k_j}\), we look at each node \(\rho_i^a\) in the partition \(k_i\). We would like to demand that the restriction,

\[
\rho_i^a \downarrow \Gamma_{k_i} \approx \bigoplus_{b=1}^{N} \sigma_i^b
\]

FIG. 2. Example of a graph. There are eight nodes labelled \(n_1\) through \(n_8\), indicated by blue diamonds, black circles, and red squares. There are eight edges, shown as solid black lines connecting pairs of nodes. Nodes in the graph can be placed (amongst other options) in following three partitions: \{\(n_1, n_2, n_3, n_8\), \{\(n_4, n_5, n_6\), and \{\(n_7\), since there are no edges connecting nodes within each of these sets; the different colors and symbols for the nodes correspond to this partitioning. This graph has two connected components: the first is the subgraph on the left consisting of the nodes \{\(n_1, n_2, n_3, n_4, n_5\), and the four edges connected to them. The second connected component consists of the three nodes \{\(n_6, n_7, n_8\) and the four edges connected to them.
is respected by the edges connecting $\rho_i^k$ to nodes in the partition $\mathbf{k}_t$.

Formalizing the preceding discussion, we define

**Definition 1.** Given a collection of little group representations, $\mathcal{M}$, (i.e. bands) forming a (physical) band representation for a space group $G$, we construct the **connectivity graph** $C_M$ as follows: we associate a node, $p_k^a \in C_M$, in the graph to each representation $p_k^a \in \mathcal{M}$ of the little group $G_k$ of (the closure of) every high-symmetry manifold (point, line, plane, and volume), $\mathbf{k}_i$. If an irrep occurs multiple times in $\mathcal{M}$, there is a separate node for each occurrence.

The degree of each node, $p_k^a$, is $P_k \cdot \dim(p_k^a)$, where $P_k$ is the number of high-symmetry manifolds connected to the point $\mathbf{k}_i$: $\dim(p_k^a)$ edges lead to each of these other $\mathbf{k}$-manifolds in the graph, one for each energy band. When the manifold corresponding to $\mathbf{k}_i$ is contained within the manifold corresponding to $\mathbf{k}_j$, as in a high-symmetry point that lies on a high-symmetry line, their little groups satisfy $G_{\mathbf{k}_j} \subset G_{\mathbf{k}_i}$. For each node $p_k^a$, we compute

$$\rho_i^a \downarrow G_{\mathbf{k}_j} \approx \bigoplus_b \rho_b^j.$$  \hspace{1cm} (20)

We then connect each node $p_k^a$ to the node $p_k^b$ with $\dim(p_k^a)$ edges.

We have thus reduced the problem of constructing all globally consistent band structures consistent with space group symmetries to the problem of constructing all valid connectivity graphs. This is an enormous simplification: instead of looking at the whole continuum of $\mathbf{k}$-vectors, we need here only look at the finite set (typically around 20) of symmetry distinct $\mathbf{k}$-manifolds. We have explained how to tabulate the minimal set of $\mathbf{k}$-manifolds for each of the 230 space groups in Ref. 14. The data is available through the BANDREP program on the Bilbao Crystallographic Server.15 It is convenient here to make a distinction between maximal and non-maximal $\mathbf{k}$-manifolds. We call a $\mathbf{k}$-vector $\mathbf{k}_0$ maximal if the little co-group $G_{\mathbf{k}_0}$ (the point group of the little group $G_{\mathbf{k}_0}$) of the $\mathbf{k}$-manifold $\{\mathbf{k}_0\}$ containing $\mathbf{k}_0$ is not a proper subgroup of the little co-group of any $\mathbf{k}$-manifold connected to $\{\mathbf{k}_0\}$. While we must consider in our connectivity graphs all partitions corresponding to maximal $\mathbf{k}$-manifolds, it happens that some non-maximal $\mathbf{k}$-manifolds give only redundant constraints on the connectivity graph. For instance, all space groups have a $\mathbf{k}$-manifold labelled $G_P$, the general position with lowest possible symmetry. In centrosymmetric crystals with time-reversal symmetry, the little co-group of $G_P$ contains the composition of inversion and time-reversal symmetries, while in all other cases it is the trivial group. In either case, the little group of a generic $\mathbf{k}$-point in $G_P$ has exactly one single-valued, and one double-valued representation. Furthermore, in every space group, we have that $G_{GP} \subseteq G_k$ for every $\mathbf{k}$-manifold $\{\mathbf{k}\}$. Thus, every $\mathbf{k}$-manifold is compatible with the general position, and the compatibility relations are entirely trivial. As such, we can consistently remove the manifold $G_P$ from our connectivity graphs without loss of generality.

A similar redundancy comes from considering closed cycles between compatible $\mathbf{k}$-manifolds, as depicted in Fig. 3. Suppose that we have six $\mathbf{k}$-manifolds $\mathbf{k}_1, \ldots, \mathbf{k}_6$. Suppose also that the little groups of these $\mathbf{k}$ manifolds satisfy the following group-subgroup relations:

$$G_{\mathbf{k}_i} \subseteq G_{\mathbf{k}_j}, \forall i$$ \hspace{1cm} (21)

$$G_{\mathbf{k}_5} \subseteq G_{\mathbf{k}_3}, G_{\mathbf{k}_5} \subseteq G_{\mathbf{k}_2}$$ \hspace{1cm} (22)

$$G_{\mathbf{k}_4} \subseteq G_{\mathbf{k}_5}, G_{\mathbf{k}_4} \subseteq G_{\mathbf{k}_2}$$ \hspace{1cm} (23)

Visually in Fig. 3 this has the interpretation that $\mathbf{k}_1, \mathbf{k}_2,$ and $\mathbf{k}_3$ are high-symmetry points. $\mathbf{k}_4$ is the symmetry line connecting $\mathbf{k}_1$ and $\mathbf{k}_2$, and similarly $\mathbf{k}_5$ is the symmetry line connecting $\mathbf{k}_2$ and $\mathbf{k}_3$. Finally, $\mathbf{k}_6$ is the symmetry plane containing all of these. In this configuration, it is easy to see that the compatibility relations along $\mathbf{k}_6$ are trivially satisfied if they are satisfied along $\mathbf{k}_4$ and $\mathbf{k}_5$. As such, we can remove the partition corresponding to $\mathbf{k}_6$ from our connectivity graphs. We summarize all such redundancies, and the algorithms we use to remove them, for every space group in Ref. 15.

With these simplifications in hand, we can now begin to systematically construct valid connectivity graphs, and hence valid global band structures. The combinatorics involved with directly carrying out this program are still quite daunting, however there are ways to make the task manageable. We present a complete set of algorithms for this task in Ref. 15 here we focus on their key conceptual insight – the reduction to subgraphs.

### C. Reduction to subgraphs

The fundamental building blocks of a connectivity graph are subgraphs consisting of three partitions corre-
responding to momenta $k_1, k_2,$ and $k_i$ such that $G_{k_i} \subset G_{k_1}$ and $G_{k_i} \subset G_{k_2}$. These subgraphs represent, for example, energy bands at two high-symmetry points $k_1$ and $k_2$ connected along a line $\{k_i\}$. The full connectivity graph is given by a union of permissible subgraphs of this type. As described in Ref. 45, by considering every permutation of valid three-partition subgraphs, we can assemble every allowed connectivity graph in this way. We must take care, however, that in piecing together the subgraphs we enumerate every connectivity graph only once. As alluded to in Sec. III B, when there are multiple nodes $p_k^{(a)}$ corresponding to the same representation $\rho_k$, then permuting these nodes in a subgraph does not result in a distinct graph, as we illustrate in Fig. 4. Furthermore, when we account for the fact that the energy bands represented by nodes in our graphs come associated with energy eigenvalues, we arrive at additional constraints from eigenvalue repulsion. Note that the inverse map from a connectivity graph to a band structure requires a certain choice of spatial embedding of the connectivity graph. In particular, all nodes in the partition labelled by $k_1$ map onto the manifold $k_i$ in the BZ. When this is done, edges of the connectivity graph may cross, corresponding to crossings of bands in the band structure. Generically, crossings along high-symmetry lines are only protected if the two bands carry different representations of the little group of the line. Accidental crossings of identical representations are not stable to perturbations: they will either gap, stay gapless over a 1D manifold of generic k-vectors (in the case of nodal lines with both inversion and TR symmetry) or in the case of Weyl nodes (which require broken inversion symmetry), they can be pushed away from high-symmetry lines. Because we are interested in classifying generic, stable band structures, we will discount connectivity graphs corresponding to accidental crossings. In Fig. 5 we show examples of a permissible and a non-permissible crossing of bands. In Ref. 45 we present a set of algorithms that we have developed to systematically rule out unstable crossings.

D. Example: the cubic crystal revisited

To see some of this machinery at work, we now revisit the $k \cdot p$ models for the cubic space groups $I 4 3 2$ (211) and $I 4 1 3 2$ (214) which we examined locally in Sec. II C. We will now construct the connectivity subgraph for these models along the $\Gamma \leftrightarrow \Lambda \leftrightarrow P$ partitions. Let us start with the symmorphic group $I 4 3 2$ (211). In the connectivity subgraph, we have partitions $\Gamma, P,$ and $\Lambda$. In the $\Gamma$ partition we have a single node corresponding to the four-dimensional $\Gamma_8$ little group representation, and in the $P$ partition we have two nodes corresponding to the representations $\overline{P}_6$ and $\overline{P}_7$. We would like to now find the compatibility relations for these representations along $\Lambda$. There are three distinct one-dimensional spinful representations distinguished by their (one-dimensional) matrix value for $\{C_3\{000\}$, and given by

$$\chi_4(\{C_3\{000\}) = -1$$

$$\chi_5(\{C_3\{000\}) = e^{-i\pi/3}$$

$$\chi_6(\{C_3\{000\}) = e^{i\pi/3}$$

(24)

From the compatibility relations computed in Ref. 46 we have

$$\Gamma_8 \downarrow G_\Lambda \approx \tilde{\chi}_4 \oplus \tilde{\chi}_5 \oplus \tilde{\chi}_6$$

(25)

and

$$\overline{P}_6 \downarrow G_\Lambda \approx \overline{\chi}_4 \oplus \overline{\chi}_6,$$

$$\overline{P}_7 \downarrow G_\Lambda \approx \overline{\chi}_4 \oplus \overline{\chi}_5,$$

(26)

Thus, in the partition labelled by $\Gamma$, we have a single node $\Gamma_8$; in the partition labelled by $P$ we have two nodes $\overline{P}_6$ and $\overline{P}_7$; lastly, in the partition labelled by $\Lambda$ we have four nodes, labelled $\overline{\chi}_4, \overline{\chi}_4^2, \overline{\chi}_5, \overline{\chi}_6$. Note that the nodes $\overline{\chi}_4^1$ and $\overline{\chi}_4^2$ correspond to the two copies of the $\overline{\chi}_4$ representation appearing in the compatibility relations Eqs. 25, 26. We deduce from the compatibility relations...
We draw this graph in Fig. 6b. As in the symmorphic example, between the Λ and Γ partitions, these two choices of connectivity are isomorphic. We illustrate this explicitly enumeration of node and edge sets. Let us first order and enumerate the m nodes of G. Using this ordering as a reference basis, we introduce the adjacency matrix \( A_\mathcal{G} \) of the graph \( \mathcal{G} \), an \( m \times m \) matrix where the \((i,j)\)’th entry is the number of edges of the form \((i,j,\alpha) \in E(\mathcal{G})\) connecting the nodes labelled by \( i \) and \( j \), or zero if no such edges exist. Symbolically, we may write

\[
(A_\mathcal{G})_{ij} = \begin{cases} \ell_{ij} & \text{if } (i,j,\alpha) \in E(\mathcal{G}), \ell_{ij} \geq 1 \\ 0 & \text{otherwise} \end{cases}
\]

Because our graphs contain no self-loops, the adjacency matrix is always purely off-diagonal. Furthermore, since our graphs are undirected, the adjacency matrix is symmetric. The sum of the elements in the \( i \)’th row of the adjacency matrix is known as the degree of the node \( i \), denoted by

\[
d_i = \sum_{j=1}^{\left|\mathcal{N}(\mathcal{G})\right|} (A_\mathcal{G})_{ij},
\]

and gives the total number of edges containing the node \( n_i \). This allows us to form the degree matrix

\[
(D_\mathcal{G})_{ij} = d_i \delta_{ij}
\]

For example, the graph \( \mathcal{G} \) in Fig. 2 has adjacency matrix

\[
A_\mathcal{G} = \begin{pmatrix}
n_1 & n_2 & n_3 & n_4 & n_5 & n_6 & n_7 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 2 \\
1 & 1 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 1 \\
0 & 0 & 0 & 2 & 0 & 0 & 1 \\
\end{pmatrix}
\]

where – as we will do in all cases where it is not readily apparent – we have labelled the rows and columns.
of the matrix by the corresponding node in our chosen ordering. We have ordered the rows and columns of $A_G$ here to correspond with our 3-partitioning of the graph $G$. The horizontal and vertical lines delineate the distinct partitions. This 3-partite structure of the graph immediately guarantees that $A_G$ to be block-off-diagonal, and allows us to build $A_G$ iteratively by considering the sub-blocks one-by-one. From this adjacency matrix Eq. (33) we compute the degree matrix

$$D_G = \begin{pmatrix} n_1 & n_2 & n_3 & n_4 & n_5 & n_6 & n_7 \\ 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 2 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 3 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 2 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 2 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 3 \\ \end{pmatrix}, \quad (34)$$

Next, let us introduce the **Laplacian matrix** for a graph $G$, defined in terms of the adjacency matrix (30) and the degree matrix (32) as

$$L_G = D_G - A_G. \quad (35)$$

As we remind the reader in Appendix [A] the zero eigenvectors of the Laplacian $L_G$ give the connected components of the graph $G$. To see this, we note that zero eigenvectors $\vec{f}$ of $L_G$ satisfy

$$\sum_c (A_G)_{cb} f_b = 0 \quad (36)$$

$$\sum_c (A_G)_{cb} f_a \delta_{ab} - (A_G)_{ab} f_b = 0. \quad (37)$$

For fixed $a$, we see that this equation is satisfied if $f_b = 1$ whenever $(A_G)_{ab} \neq 0$. By induction, this allows us to deduce that the vector

$$f_b = \begin{cases} 1, & \text{a and b lie in the same connected component} \\ 0, & \text{otherwise} \end{cases} \quad (38)$$

is a zero eigenvector of $L_G$, as supposed. As a corollary, the multiplicity of the zero eigenvalue of $L_G$ gives the number of connected components of the graph $G$.

As an example, let us consider the Laplacian matrix for the graph given in Fig. (2). Subtracting the adjacency matrix Eq. (33) from the degree matrix Eq. (34), we find

$$L_G = \begin{pmatrix} n_1 & n_2 & n_3 & n_4 & n_5 & n_6 & n_7 \\ 1 & 0 & 0 & 0 & -1 & 0 & 0 \\ 0 & 2 & 0 & 0 & -1 & -1 & 0 \\ 0 & 0 & 1 & 0 & 0 & -1 & 0 \\ 0 & 0 & 0 & 3 & 0 & 0 & -1 \\ -1 & -1 & 0 & 0 & 2 & 0 & 0 \\ 0 & -1 & -1 & 0 & 0 & 2 & 0 \\ 0 & 0 & 0 & -1 & 0 & 2 & -1 \\ 0 & 0 & 0 & 0 & 0 & -1 & 3 \\ \end{pmatrix}, \quad (39)$$

From Fig. [4] we see that the vectors

$$\vec{x}_1^T = (n_1 \ n_2 \ n_3 \ n_4 \ n_5 \ n_6 \ n_7)^T = (1 \ 1 \ 1 \ 0 \ 1 \ 1 \ 0 \ 0) \quad (40)$$

and

$$\vec{x}_2^T = (n_1 \ n_2 \ n_3 \ n_4 \ n_5 \ n_6 \ n_7)^T = (0 \ 0 \ 0 \ 1 \ 0 \ 0 \ 1 \ 1) \quad (41)$$

take the value 1 on exactly one of the connected components of $G$, and zero on the other connected component. It is straightforward to verify that

$$L_G \vec{x}_1 = L_G \vec{x}_2 = 0. \quad (42)$$

Finally, since the characteristic polynomial $p_{L_G}(\lambda)$ of the Laplacian matrix is

$$p_{L_G}(\lambda) = \lambda^2(\lambda-5)(\lambda-3)(\lambda^2-5\lambda+5)(\lambda^2-3\lambda+1), \quad (43)$$

we see immediately that there are only two zero-eigenvectors. Thus two vectors $\vec{x}_1^T$ and $\vec{x}_2^T$ span the entire null space of $L_G$ in accordance with our claim.

### VI. GRAPHS AND BAND REPRESENTATIONS

We can now apply the tools of spectral graph theory to the connectivity graphs of Sec. [IV]. This will allow us to identify all of the disconnected components of a connectivity graph, and hence under the inverse of our graph theory mapping, all of the disconnected groups of bands in a global band structure. As such, we can in principle use these tools to identify all possible insulating band structures allowed in each space group. We are primarily interested, however, in topological insulators. To determine whether a given disconnected component of a connectivity graph corresponds to a topological group of bands, we will combine the graph theory mapping presented here with the theory of band representations of Refs. [1] and [2]. After a brief review of the theory of band representations, we will show how we can enumerate topological band structures by finding the allowed disconnected connectivity graphs corresponding to the so called “elementary band representations” (EBRs). We will then see how this works in the example of a two-dimensional inversion symmetric topological insulator, and so demonstrate how our theory contains the standard theory of eigenvalue-based topological invariants. Finally, we will show how our graph theory method allows us to predict not only topological insulators, but topological semimetals as well.

#### A. Review of band representations

Band representations, first introduced by Zak [20] and applied to spin-orbit coupled and topological materials in a series of papers by the present authors [11][12][13][14] relate the...
real-space localized electronic orbitals in a crystal to the momentum-space band structure. Roughly speaking, orbitals located at a position in the unit cell of a crystal transform under a representation of the symmetry group of the local crystal field (the site-symmetry group) of that position. By applying to this orbital the remaining crystal symmetry operations (including translation), we induce (in the sense of representation theory, c.f. Ref. [3]) an infinite-dimensional representation of the full space group. The Fourier transform of this representation determines the little group representations at every \( k \)-vector of the Brillouin zone in the energy bands coming from these orbitals. For the mathematical details we defer the reader to Refs. [1] and [2].

As constructed, every band structure that can be obtained from localized orbitals that respect all crystal symmetries transforms according to some band representation. Furthermore, in analogy with the concept of a finite dimensional irreducible representation, all band representations can be expressed as a sum of “elementary building bricks.” Following Zak we refer to these as elementary band representations (EBRs), or, if we also enforce the role of time-reversal symmetry, physically elementary band representations (PEBRs). All band structures that can be continuously deformed to an atomic limit without breaking either crystal (or time-reversal) symmetries transform under a sum of (P)EBR. As a corollary, any disconnected group of bands which does not transform under some sum of (P)EBRs must give a topological insulator. In the following, we will show how, by applying our graph theory mapping to the set of bands in a (P)EBR, we are able to enumerate topological band structures.

### B. Connectivity and topology

While the theory of band representations tells us which little group representations appear together at all high-symmetry \( k \)-points and lines in atomic limit band structures, it does not in itself tell us how these energy bands are allowed to be connected in a real material. Applying the ideas of Ref. [1] discussed above, we see that the connectivity of elementary and physically elementary band representations lies at the heart of the search for topological insulators: if the bands transforming in a (P)EBR are disconnected in the Brillouin zone, then filling only logical insulators: if the bands transforming in a (P)EBR is a symmorphic space group with primitive tetragonal Bravais lattice. Taking the lattice vectors \( \{e_1, e_2, e_3\} \) to be aligned with the Cartesian directions, we can express the point group \( C_{4v} \) as the group generated by

\[
C_{4z} : \{e_1, e_2, e_3\} \rightarrow \{-e_2, e_1, e_3\} \\
m_x : \{e_1, e_2, e_3\} \rightarrow \{-e_1, e_2, e_4\}.
\]

Since all point group elements act trivially on the \( e_3 \) lattice vector, this space group can be viewed as stacks of \( 2D \) planes, each consisting of a square lattice with wallpaper symmetry group \( p4mm \) (wallpaper group number 11, generated by \( C_4 \) and \( m_x \)).

There are three maximal Wyckoff positions (Wyckoff positions with site-symmetry groups \( G_4 \) that are maximal finite subgroups of the space group) in this 3D space group, as indicated in Fig. [7]. First, the \( 1a \) position corresponds to vertical lines through the Bravais lattice sites, and has representative \( \{q^a\} = \{(0, 0, z)\} \). Second is the \( 1b \) position, with representative \( \{q^b\} = \{1/2, 1/2, z\} \).
This position lies on a vertical line extending upward from the center of the 2D square lattice unit cell. Both of these positions have multiplicity 1, and so have site-symmetry groups isomorphic to the full point group $C_{4v}$. As the 1a position lies on the Bravais lattice sites, its stabilizer $G_{\text{st}}$ is the full point group, generated by $\{C_{4v}|000\}$ and $\{m_x|000\}$. The stabilizer of the 2b position, on the other hand, is generated by $\{C_{4v}|000\}$ and $\{m_x|100\}$. In Table I we give the character table for $C_{4v}$. Because both the 1a and 1b Wyckoff positions have multiplicity 1, all band representations induced from these positions are trivially connected, with connectivity given by the dimension of the site-symmetry representation. To see this, note that since the stabilizer groups of both these positions are isomorphic to the point group, the connectivity graphs for these band representations contain a single node in the partition corresponding to the Γ point, and labelled by the site-symmetry irrep. If a connectivity graph has a single node in any partition, then it is impossible to divide it into disconnected subgraphs, as the node would need to be split in (at least) two. Thus, all elementary band representations induced from these Wyckoff positions give topologically trivial bands.

![Diagram](https://example.com/diagram.png)

**FIG. 7.** Maximal Wyckoff positions for space group $P4mm$ (99), projected along the $\hat{z} = [001]$ direction. The blue star indicates the 1a position at the Bravais lattice sites, the red diamond indicates the 1b position at the center of the unit cell, and the black circles denote the 2c Wyckoff position at the middle of the edges. Because space group $P4mm$ (99) has no symmetries constraining $z$, the coordinates of these positions extend in the $z$ direction (perpendicular to the plane of the page).

| Rep | $E$ | $C_{2v}$ | $\{m, \hat{E}m\}$ | $\{C_{4v}, m, \hat{E}C_{4v}, m\}$ | $\mathbf{E}$ | $\mathbf{EC}_{4v}$ |
|-----|-----|-----|-----------------|-----------------|------|------|
| $\Gamma_1$ | 1 | 1 | 1 | 1 | 1 | 1 |
| $\Gamma_2$ | 1 | 1 | -1 | 1 | -1 | 1 |
| $\Gamma_3$ | 1 | 1 | -1 | 1 | 1 | 1 |
| $\Gamma_4$ | 1 | 1 | 1 | -1 | -1 | 1 |
| $\Gamma_5$ | 2 | -2 | 0 | 0 | 0 | 2 |
| $\Gamma_6$ | 2 | 0 | $-\sqrt{2}$ | 0 | 0 | $-2\sqrt{2}$ |
| $\Gamma_7$ | 2 | 0 | $\sqrt{2}$ | 0 | 0 | $-2\sqrt{2}$ |

**TABLE I.** Character table for the (double) point group $C_{4v}$. Note that the conjugacy class denoted by $\{m, \hat{E}m\}$ contains $m_x$, $m_y$, and their inverse $\hat{E}m_x$ and $\hat{E}m_y$; hence they all have the same characters. Similarly, the conjugacy class $\{C_{4v}, m, \hat{E}C_{4v}, m\}$ contains $m_{x\pm\frac{\pi}{2}}$, $m_{y\pm\frac{\pi}{2}}$, and their inverses. Note that spinful orbitals transform in the same representation. For instance, the symmetry operation $\Gamma_6$ or $\Gamma_7$ representations depending on their total azimuthal quantum number $m_s - s$ orbitals (with $m_z = \pm\frac{\pi}{2}$) transform in the $\Gamma_7$ representation, as do $P^\frac{\pi}{2}$ states, and $P^\frac{\pi}{2}$ states with $m_z = \pm\frac{\pi}{2}$. The remaining $P^\frac{\pi}{2}$ states with $m_z = \pm\frac{\pi}{2}$ transform in the $\Gamma_8$ representation.

More interesting for our purposes is the maximal 2c position, with representatives $\{q_1^c, q_2^c\} = \{(0, 1/2, z), (1/2, 0, z)\}$. The site-symmetry group $G_{q_1^c}$ of the representative $q_1^c$ is isomorphic to the group $C_{2v}$, and is generated by $\{C_{2v}|010\}$ and $\{m_z|000\}$. The representations of this group are given in Table II. Let us focus on the spin-orbit coupled case, where there is a unique double-valued site-symmetry representation $\bar{\Gamma}_5$, and hence only the band representation $\bar{\Gamma}_5 \uparrow G$ can be induced from orbitals at this Wyckoff position. This representation is generated by a single $s$ orbital with spin up and spin down states at each of the 2c sites, forming stacked layers of a checkerboard lattice. Consulting Refs. [1] and [10] we see that with TR symmetry these orbitals furnish a 4-band elementary band representation. By computationally constructing the connectivity graphs for this band representation following the algorithms outlined in Sec. V and elaborated upon in Ref. [15] we find from Ref. [16] that this band representation has disconnected connectivity graphs (corresponding to TIs), which we now examine further.

We can construct the disconnected realizations of this band representation by examining the compatibility relations for space group $P4mm$ (99). The maximal $k$-vectors in the Brillouin zone relevant to the 2D system are $\Gamma = (0, 0, 0), X = (0, 1/2, 0)$, and $M = (1/2, 1/2, 0)$. Note that $C_{4v}$ symmetry relates $X$ to $X' = (1/2, 0, 0)$, and hence all these high-symmetry points, the little groups and their representations are independent of $k_z$. The little co-groups of the points $\Gamma$ and $M$ are the full point group, $C_{4v}$, while the little co-group of $X$ is isomorphic to the group $C_{2v}$. To examine $k$-space compatibility, we also look at the lines connecting $\Gamma$, $M$, and $X$. Particularly relevant are the lines $\Sigma = (k_x, k_x, 0)$, $\Delta = (0, k_y, 0)$ and $T = (k_x, 1/2, 0)$, with $k_x, k_y \in [0, \frac{1}{2}]$. $\Sigma$ connects the points $\Gamma$ and $M$, and has little co-group $C_s$ generated by $m_x C_{2s}$. The line $\Delta$ connects $\Gamma$ and $X$, and has little co-group $C_s$ generated by $m_x$. Finally, the line $T$ connects $X$ and $M$, and has little co-group $C_s$ generated by $m_y$. 

**TABLE II.** Character table for the (double) group $C_{2v}$, for both single- and double-valued representations. The single-valued representations $\Gamma_1, \Gamma_4$ are all one dimensional. The unique double-valued representation $\bar{\Gamma}_5$ is the two-dimensional spin-$\frac{1}{2}$ representation. In terms of the Pauli matrices it is given concretely as $\bar{\Gamma}_5(C_{2v}) = i\sigma_z, \bar{\Gamma}_5(m) = i\sigma_y$.
Performing the induction procedure for the band representation induced from the 2c position, and omitting the details (c.f. Ref.1 and 32), the representations appearing in the Brillouin zone are presented in Table III.

Next, we analyze the compatibility between these representations. To do so, note that the little group irreps at $M$, $\Gamma$, and $X$ are all two-dimensional, as expected from Kramers’s theorem. Furthermore, by consulting Tables 1 and II, we see that all mirror elements in these irreps have character 0. This implies that along the high symmetry lines $\Delta$, $\Sigma$, and $T$, each two-dimensional little group irrep decomposes into one copy of $\Sigma_4$ and one copy of $\Sigma_3$, whose sum of mirror characters is zero. It is now clear that there are two disconnected compatibility graphs, depending on whether $\Gamma_6$ connects to $\overline{M}_6$ or $\overline{M}_7$ along the line $\Sigma$. We show a visual depiction of these two disconnected graphs in Fig. 8. In Eqs. (46) and (47), we show the two distinct Laplacian matrices for the $\Gamma - \Sigma - M$ subgraph of the connectivity graph. Both of these disconnected solutions correspond to topological phases.

In order to determine whether this is a strong, weak, or crystalline topological phase, we will impose inversion symmetry as an additional space group symmetry. This will allow us to use the Fu-Kane inversion eigenvalue formula[3] to compute topological indices from knowledge of only the valence-band little group representations. Adding inversion takes us from space group $P4mm$ (99) to space group $P4/mmm$ (123). The addition of spatial inversion forces the $z$-coordinate of our symmetry centers $\{(0, \frac{1}{2}, z), (\frac{1}{2}, 0, z)\}$ to be fixed at either 0 or 1/2. We focus on the $z = 0$ case for concreteness, since the analysis and results are identical for the $z = 1/2$ case. Note that in SG $P4/mmm$ (123), the po-

$C_s$ has two double valued representations which we denote by $\rho_\pm$, both one dimensional, and distinguished by whether the mirror is represented by $i$ or $-i$. Because the group $P4mm$ is symmorphic, the representation $\rho_+$ of $C_s$ uniquely determines the little group representations $\Sigma_4$ of $G_\Sigma$ and $\overline{T}_4$ of $G_T$. Similarly, the representation $\rho_-$ uniquely determines the little group representations $\Sigma_3$ of $G_\Sigma$ and $T_3$ of $G_T$. Physically, this means that the representations of the little groups $G_\Sigma$ and $G_T$ do not change as we move along the lines $\Sigma$ and $T$.
situation \{0, \frac{1}{2}, 0, \frac{1}{2}, 0\} is conventionally denoted 2e, and we adopt that notation for the remainder of this section. In addition to the elements of $C_2v$, enumerated above, the site-symmetry group of $q_f^x = (0, 1/2, 0)$ now also contains \{1|010\}. Thus, the site-symmetry group is isomorphic to $D_{2h}$, whose representations we enumerate below. We note, however, that each representation of $D_{2h}$ is generated by taking a representation of $C_2v$ and appending inversion represented as either plus or minus the identity matrix. Since we started with s-orbitals in SG $P4mm$ (99), we know that \{0|100\} should be represented by the identity matrix in the relevant representation. Thus, when we add inversion to the disconnected band representation in SG $P4mm$ (99), we arrive at the band representation induced from the $\Gamma_5$ representation of the site-symmetry group $D_{2h}$ in SG $P4/mmm$ (123). Consulting Ref. \[96\] we see that this band representation is physically elementary, and allows for disconnected connectivity graphs; this means we can add inversion symmetry without changing the band connectivity. To analyze the topological character of the phases where this band rep is disconnected, we construct explicitly the $k$-dependent band representation matrices. Let $s_i$ be a set of Pauli matrices operating in spin space, and let $\sigma_i$ be a set of Pauli matrices operating in sublattice space \{s_1, s_2\}. We start by constructing the matrix representative of inversion \{I|000\}. First, we note that inversion acts trivially in spin space. Next, using the fact that \begin{align*}
\{I|000\} q_f^x &= q_f^x - (0, 1, 0) \\
\{I|000\} q_f^y &= q_f^y - (1, 0, 0)
\end{align*}
we deduce using the methods of Refs. \[1\] and \[2\] that we can represent inversion as \begin{equation}
p^k(I) = s_0 \otimes \begin{pmatrix} e^{i{k_e}e_z} & 0 \\
0 & e^{i{k_e}e_z} \end{pmatrix}
\end{equation}
Let us pause to make the following observation: there are 8 inversion-symmetric momenta in the BZ: $\Gamma$, $X$, $X'$, and $M$, along with the points $Z$, $R$, $R'$, and $A$, directly above them in the $z$ direction. Assume now that our band representation $\Gamma_5 \uparrow G$ is realized as disconnected, with two valence and two conduction bands. Because of $C_4v$ symmetry, the inversion eigenvalues of the valence bands at $X$ and $X'$ are the same; so are the inversion eigenvalues at $R$ and $R'$. Examining $p^k(I)$, we see that at $\Gamma$ and $Z$ all four inversion eigenvalues are positive, while at $M$ and $A$ all inversion eigenvalues are negative. At $X$ and $R$ two eigenvalues are positive, and two are negative. Using the well-known relationship between inversion eigenvalues and $Z_2$ topological indices,\[22\] we conclude immediately that all disconnected realizations of this band representation give a weak topological insulator, with indices \{0; 111\}. In Appendix \[13\] we present an explicit tight-binding model realizing this topological phase. We emphasize again that the only phases permitted for this band representation are either a connected topological semimetal, or a disconnected topological insulator. This is analogous to the situation present in graphs,\[24, 55\] and indeed is true of any elementary band representation which has a disconnected connectivity graph.

D. Filling constraints: insulators and semimetals

We have in this section been primarily focused with using the method of connectivity graphs to find topological insulating band structures. However, the presence of disconnected groups of bands in a band structure is not in itself sufficient to guarantee that a material is insulating; it must also be true that connected bands are completely filled, i.e. that (accounting for possible charge-transfer from other bands) there is one electron per band per unit cell. When the electron filling is less than the number of bands in an isolated group, a metallic phase naturally results.

Crucially, we can use our graph theory mapping to assess when a set of $k \cdot p$ Hamiltonians can be patched together to form a band structure which allows for a topological semimetallic phase. By this we mean a group of bands that, at the appropriate filling, can host a zero-volume point or line-like Fermi surface (possibly in the presence of additional compensated Fermi pockets). While we defer the systematic treatment of semimetallic connectivity to a future work, we can already see some interesting examples in the connectivity graphs we have examined previously. First, let us return to the example of Sec. \[IVD\] where we examined the connectivity of bands originating from $P_{32}$ orbitals in the cubic space groups $I432$ (211) and $I4_{1}32$ (214). Returning to Figs. \[6a\] and \[6b\] we recall that both band structures have only connected connectivity graphs, with a single representation at the $\Gamma$ point, and with two representations at the $P$ point. We see that at half-filling, the band structure corresponding to the connectivity graph Fig. \[6a\] in SG $I432$ can yield a semimetal with a pointlike Fermi surface at $\Gamma$; this is in fact a “Spin-3/2 Weyl” semimetal first predicted and discussed in Ref. \[38\]. On the contrary, by counting the number of edges in Fig. \[6b\] we see that a topological semimetallic structure in SG $I4_{1}32$ (214) necessarily hosts Fermi surface features away from the $\Gamma$ point. In fact, if we consider additionally the partition $N$ corresponding to the $N$ point (with coordinates $\frac{1}{2}g_3$), we find\[25\] that there are two nodes each corresponding to the two-dimensional representation $\overline{\Gamma}_5$ of the little group $G_N$. Comparing the connection $\Gamma \leftrightarrow \Sigma \leftrightarrow N$ with the connection $P \leftrightarrow D \leftrightarrow N$ using the BANDREP application, we can decisively rule out the existence of a semimetal with features only at $\Gamma$ in this connectivity graph: there must be an additional Fermi pocket centered near $P$ or $N$.

A second example of topological semimetallic behavior is given by connectivity graphs with symmetry-enforced crossings along high-symmetry lines and planes, as in Fig. \[6b\]. Along mirror planes in mirror symmetric systems, these types of crossings are generically allowed, but not required. However, along glide planes and screw axes...
in non-symmorphic systems, these crossings are generically required, leading to the “movable, but not removable” band crossings first pointed out by Michel and Zak. At half-filling, band structures in these glide- and screw-symmetric systems will generically yield Weyl and nodal-line semimetals.

VII. DISCUSSION

We have shown how graph theory can be used to solve the problem of constructing and classifying global band structures consistent with crystal and time-reversal symmetries. While symmetry strongly constrains the allowed connections of energy bands, there are often many consistent global band structures with different numbers of disconnected bands; our graph theory mapping provides the tools necessary to algorithmically enumerate all valid band structures. In combination with the theory of elementary band representations of Refs. 1 and 2 we demonstrated that the mapping of global band structures to connectivity graphs gives a powerful method for enumerating topological band structures. In the accompanying Ref. 45 we show how to practically implement this programme to algorithmically compute all allowed topologically disconnected elementary band representations in all 230 space groups both with and without time reversal symmetry; the results of these computations are now publicly available on the Bilbao Crystallographic Server.

While we have discussed the theory of connectivity graphs largely within the context of the electronic structure of time-reversal invariant crystals, the applications of our method are far grander in scope. The methods here can in principle be adapted to the electronic structure of magnetic materials, the dispersion relation of vibrational modes, and spin-wave spectra. Furthermore, we expect connectivity graphs to play a central role in the classification of topological semimetals, and the theory of non-interacting topological phase transitions.

Appendix A: Discussion of the Graph Laplacian

In this Appendix, we explore the motivation for the name “Graph Laplacian.” Let us consider the graph \(G\) obtained by discretizing \(D\)-dimensional Euclidean space with a regular hypercubic lattice with lattice constant \(a = 1\). This is an infinite graph with nodes \(N(G) = \{x^n = (x^n_1, \ldots, x^n_D)\}\) corresponding to points in the lattice, each with degree (coordinating number) \(2D\). Edges connect nearest-neighbor lattice sites in the usual way. Now, if we have some function \(f : \mathbb{R}^D \rightarrow \mathbb{R}\) on space, we can discretize it by restricting to the points \(N(G)\). The discretization of the Laplacian \(\nabla^2 f\) of \(f\) at a lattice site \(x_a\) can be written

\[
-\nabla^2 f(x_a) \approx -\sum_{\mu=1}^{D} \left[ f(x_a + e^\mu) + f(x_a - e^\mu) - 2f(x_a) \right],
\]

where \(e^\mu = (0, \ldots, 0, x_\mu = 1, \ldots, 0)\). We note that each term in square brackets is the discretization of the second derivative of \(f\) in the \(\mu\) direction. However, also note that

\[
\sum_{\mu=1}^{D} 2f(x_a) = 2D f(x_a) = (D_G)_{ab} f(x_b).
\]

where \((D_G)_{ab} = 2D\delta_{ab}\). Furthermore, note that the sum

\[
\sum_{\mu=1}^{D} (f(x_a + e^\mu) + f(x_a - e^\mu))
\]

is the sum of \(f\) evaluated at all nodes adjacent to \(x_a\). Using the fact that the adjacency matrix \(A_G\) as defined in Eq. (30) has matrix elements \((A_G)_{ab}\) equal to 1 for each node \(b\) connected to \(a\), and zero otherwise, we can rewrite this as

\[
\sum_{\mu=1}^{D} (f(x_a + e^\mu) + f(x_a - e^\mu)) = (A_G)_{ab} f(x_b).
\]

Putting it all together the discretized Laplacian of \(f\)

\[
-\nabla^2 f(x_a) \approx (L_G)_{ab} f(x_b)
\]

coincides with the graph Laplacian.

Spectral graph theory starts with the study of the eigenvalues of the graph Laplacian. We can motivate the relation between these eigenvalues and the connectivity of the graph using the correspondence Eq. (A5). In particular, let us consider the diffusion equation

\[
\partial_t f = \nabla^2 f
\]

on \(\mathbb{R}^d\). We know that if \(\phi_E(x)\) is an eigenfunction of \(\nabla^2\) with eigenvalue \(-E\), then

\[
f_E(x, t) = e^{-Et} \phi_E
\]

solves the diffusion equation. Because the Laplacian is negative definite, all solutions decay to zero at infinite time unless \(E = 0\). If we consider diffusion on a subspace of our discretized \(\mathbb{R}^D\) given by the union of \(N\) disconnected sublattices with Neumann boundary conditions, we know that there will be \(N\) eigenfunctions of \(\nabla^2\) with eigenvalue \(E = 0\): these are given by the functions which are 1 on exactly one of the discs, and zero on the others. These give steady states of the diffusion equation, which are constants on each of the \(N\) sublattices.

From this fact we may suspect that zero eigenvectors of the graph Laplacian \(L_G\) correspond to connected components of the graph \(G\), and we would be correct; we proved this in the main text in Eqs. (37) (38).
FIG. 9. Tight-binding band structure for s orbitals at the 2f position in SG P4/mmm (123) in two dimensions. (a) shows the band structure with no spin-orbit coupling, which yields a semimetal with a gapless point at M. (b) shows the spectrum with nonzero SOC. The band structure is fully gapped.

FIG. 10. SG P4/mmm (123) tight-binding model on a finite slab of fifty unit cells. (a) shows the full slab band structure, with two pairs of counterpropagating edge modes clearly visible in the bulk gap. Because the total center of charge of orbitals at the 2f position are not on Bravais lattice sites, our boundary conditions break inversion symmetry. (b) shows the surface density of states as a function of energy and surface momentum, showing one pair of counterpropagating edge modes. The other pair of edge modes are on the other surface of the slab (not shown).

Appendix B: Tight-Binding model for the Disconnected EBR in SG P4/mmm

It remains for us to show that such a disconnected realization exists. This follows immediately from imposing inversion symmetry on the disconnected band structure we found for SG P4mm (99). Inversion forces bands to be doubly degenerate everywhere, and this leaves our disconnected band structure disconnected. To see this more
clearly, however, we construct an explicit tight-binding model. Noting that in SG $P4/mmm$ (123) along every vertical line in the BZ there is only one allowed two-dimensional little group representation, we know that the only possible crossings in the $z$ direction can be four-fold crossings of these representations. However, level repulsion ensures that we can eliminate all such crossings. Hence we can dispense with the $z$-direction in our model. We then construct a 2$D$ tight binding model consistent with the space group symmetries – as with graphene (discussed in Ref. [54]) as well as in the Supplementary Information of Ref. [55], the effect of inversion in this 2$D$ model is to enforce $s_z$ spin conservation. In addition to inversion, we find for the remaining space group generators

$$
\rho^k(C_{4z}) = e^{i k \cdot e_1/2} e^{-i \pi/4 s_z} \otimes \left( \frac{\cos k \cdot e_1}{2} \sigma_x + i \sin \frac{\cos k \cdot e_1}{2} \sigma_y \right)
$$
(B1)

$$
\rho^k(m_x) = -i e^{i k \cdot e_z s_z/2} s_x \otimes \left( \frac{\cos k \cdot e_1}{2} \sigma_0 - i \sin \frac{\cos k \cdot e_1}{2} \sigma_z \right)
$$
(B2)

$$
\rho^k(T) = i s_y \otimes \sigma_0 K,
$$
(B3)

where $K$ is the complex conjugation operator. We seek a Hamiltonian consistent with these symmetries, that is,

$$
\rho^k(I) H(k)(\rho^k(I))^{-1} = H(-k) \quad (B4)
$$

$$
\rho^k(T) H(k)(\rho^k(T))^{-1} = H(-k) \quad (B5)
$$

$$
\rho^k(m_x) H(k)(\rho^k(m_x))^{-1} = H(m_x k) \quad (B6)
$$

$$
\rho^k(C_{4z}) H(k)(\rho^k(C_{4z}))^{-1} = H(C_{4z} k) \quad (B7)
$$

For the sake of illustration, we will look for the shortest-ranged fully gapped Hamiltonian satisfying these properties. The shortest range Hamiltonian which can support a fully gapped insulator reads

$$
H(k) = t_1 \left[ (1 + \cos k \cdot e_1 + \cos k \cdot e_2 + \cos k \cdot (e_2 - e_1)) \sigma_x + (\sin k \cdot e_1 - \sin k \cdot e_2 - \sin k \cdot (e_2 - e_1)) \sigma_y \right] + t_2 \left[ (\cos k \cdot e_1 - \cos k \cdot e_2) \sigma_z + \lambda \left[ (1 - \cos k \cdot e_1 - \cos k \cdot e_2 + \cos k \cdot (e_2 - e_1)) s_x \otimes \sigma_z \right] + (\sin k \cdot e_1 - \sin k \cdot e_2 + \sin k \cdot (e_2 - e_1)) s_z \otimes \sigma_x \right]
$$
(B8)

where $t_1$ is a nearest-neighbor inter-sublattice hopping, $t_2$ is a next-nearest-neighbor intra-sublattice hopping, and $\lambda$ is the nearest-neighbor spin-orbit coupling (SOC). We note that a closely related model was recently considered in a different context in Ref. [59]. When $\lambda = 0$, the system is gapless at the $M$ point, as shown in Figure 9a. $s_z$-conserving “Haldane type” SOC[1169] opens a gap at the $M$ point, giving us a fully disconnected realization of this elementary band representation, as shown in Fig. 9b. Analysis of the inversion eigenvalues reveals that this is a 2$D$ strong topological insulator (layers of which form the 3$D$ weak topological insulator as above). To see this directly, we diagonalize our tight-binding model in a slab geometry, periodic in $y$ and with 50 layers in the $x$-direction. Figure 10a shows the slab band structure as a function of the surface momentum. Two pairs of counterpropagating midgap states are clearly seen, one pair coming from each boundary of the slab. To show that these states are indeed localized to the edge, we compute the surface density of states on the right boundary of the slab, shown in Fig. 10b. One pair of counterpropagating edge states is clearly visible. This confirms our assertion that this disconnected elementary band representation realizes a weak 3$D$ topological insulator (i.e. a strong 2$D$ topological insulator).

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