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Topological nodal line semimetals*

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We review the recent, mainly theoretical, progress in the study of topological nodal line semimetals in three dimensions. In these semimetals, the conduction and the valence bands cross each other along a one-dimensional curve in the three-dimensional Brillouin zone, and any perturbation that preserves a certain symmetry group (generated by either spatial symmetries or time-reversal symmetry) cannot remove this crossing line and open a full direct gap between the two bands. The nodal line(s) is hence topologically protected by the symmetry group, and can be associated with a topological invariant. In this review, (i) we enumerate the symmetry groups that may protect a topological nodal line; (ii) we write down the explicit form of the topological invariant for each of these symmetry groups in terms of the wave functions on the Fermi surface, establishing a topological classification; (iii) for certain classes, we review the proposals for the realization of these semimetals in real materials; (iv) we discuss different scenarios that when the protecting symmetry is broken, how a topological nodal line semimetal becomes Weyl semimetals, Dirac semimetals, and other topological phases; and (v) we discuss the possible physical effects accessible to experimental probes in these materials.

Keywords: topological semimetals, nodal band structures, topological invariants

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1. Introduction

Topological semimetals (TSM) are defined as systems where the conduction and the valence bands cross each other in the Brillouin zone (BZ), and the crossing is non-accidental, i.e., cannot be removed by perturbations on the Hamiltonian without breaking any of its symmetries. If there be no symmetry, two bands, when close in energy, will hybridize with each other and maintain a gap in-between, through a mechanism known as the band repulsion; however, in the presence of certain symmetries (e.g., crystalline symmetries and time-reversal symmetry), the two crossing bands may have different quantum numbers such that they cannot be hybridized. From this, we see that all band crossings, hence all topological semimetals, can only be protected by symmetries and hence belong to symmetry protected topological phases of matter.

In three dimensions (3D), two bands can cross each other either at discrete points or along a closed curve. In the former case, there are Weyl semimetals^[1,2] and Dirac semimetals^[3] that have been intensively studied in theory as well as in experiment. In the latter case, the curve where the bands cross is called a nodal line,^[4] which may either take the form of an extended line running across the BZ, whose ends meet at the BZ boundary,^[5] or wind into a closed loop inside the BZ,^[6] or even form a chain consisting of several connected loops (nodal chain).^[7] Topological semimetals with such line band cross-

ings are called topological nodal line semimetals (TNLSM). In principle, one may have TNLSM in both quasi-2D^[8] and 3D systems,^[9,10] and in this review, we will focus on the latter.

In a topological gapped phase, such as Chern insulator, topological insulator or topological crystalline insulator, the nontrivial topology of the bands can be characterized by a topological invariant, a quantum number that depends on the Bloch wave functions of the valence bands as a whole. The form of the topological invariant is determined by only two factors: dimension and symmetry. Similarly for a topological semimetal, one can also associate with each band crossing (either a point node or a line node) a topological invariant, whose form depends on the symmetry group that protects the nodal structure. Given the dimension of the node and the symmetry group that protects it, one or several topological invariants can be found, and these quantum numbers provide a full topological classification of the nodes. Up to this day, the classification of TNLSM is yet to be complete. Theoretically, people have found mirror reflection protected nodal lines with a Z-invariant,^[6,11–14] nodal line protected by inversion, time-reversal and spin rotation symmetries with two Z₂-invariants,^[5,15–26] screw rotation protected double nodal lines (to be defined later) with a Z-invariant,^[27–29] etc.

While a topological classification tells us how many topo-

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logically different types of nodes exist in systems with a given symmetry group, only numerical calculation, mainly first-principles calculations that compute the band structure and Bloch wave functions, can compute this invariant in a given compound and determine if it is a TSM or not. The calculation has proved more challenging than that of the topological invariant in a gapped system, because of (i) the slow convergence in a gapless system and the fact that (ii) the band crossing point is not always at a high-symmetry point, but along a high-symmetry line or even on a high-symmetry plane. We will review numerical efforts that have resulted in the proposals of various materials systems as TNLSM protected by different symmetry groups.

When the protecting symmetry is broken in a TNLSM, the nodal line is either fully gapped or gapped into several nodal points. For example, without spin-orbital coupling (SOC), TaAs was,^[30,31] in first-principles calculation, shown to be a TNLSM protected by mirror reflection and spin-rotation symmetries with two nodal lines, and when SOC was turned on, each nodal line is gapped into three pairs of Weyl nodes. Another example is the double nodal line in SrIrO₃, which is gapped into a pair of non-symmorphic Dirac nodes when a certain mirror reflection symmetry is broken.^[27,32] Therefore, understanding how a nodal line evolves upon symmetry breaking can help predict new topological materials.

Unlike most topological phases, TNLSMs in general do not have protected boundary modes.^[28] Therefore, identifying them in experiments poses a challenge to the experimentalists. Angle resolved photoemission spectroscopy (ARPES) has been used to resolve the nodal band structure in the bulk,^[11,12] but the limited momentum resolution in the perpendicular direction prevents these efforts from being deterministic. Quantum oscillation measurements^[33] were performed to map the Fermi surface of ZrSiSe and ZrSiTe as well as the Berry phase along a closed loop on the Fermi surface, partially supporting the proposal of TNLSMs in these materials. Another unanswered question is the fate of these materials in the presence of electron correlation. It has been proposed that the screening effect is qualitatively different from that in normal metals,^[34] and that in the presence of strong electron interaction, the quantum phase transition from a TNLSM to a nodal point semimetal or a gapped system belongs to a new universality class.^[35]

Below is the outline of the review. In Section 2, we briefly go through the topological classification of TNLSM protected by several symmetry groups by writing down the expression of the topological invariants in terms of the Bloch wave functions. In Section 3, we review the several materials proposed by first principle calculations to be TNLSMs. In Section 4, we discuss various scenarios of how the nodal line evolves when the protecting symmetry is broken. In Section 5 we discuss the experimental consequences and the many-body effects in

TNLSMs so far proposed in the literature.

2. Topological classification

Topological invariant gives a quantitative description of the topology of a system. Let us first review its definition in a ‘gapped’ band structure, where ‘gapped’ means that at each momentum k in BZ, there is a finite direct gap between the conduction and the valence bands, while the indirect gap is allowed to close. Given two Hamiltonians H_1 and H_2 , if H_1 can be tuned to H_2 without (i) closing the gap or (ii) breaking a given symmetry group, then H_1 and H_2 are said to be topologically equivalent under this symmetry group. By this equivalence, one can divide the Hamiltonians into different equivalent classes, and there is a one-to-one isomorphism mapping each class to a set of integers: the explicit form of this isomorphism is the topological invariant(s) protected by the symmetry group. Well-known examples of topological invariants include: Chern numbers in 2D Chern insulators (quantum anomalous Hall states),^[36] Z_2 indices in 2D and 3D insulators protected by time-reversal and charge conservation,^[37–39] the spin Chern numbers in 2D quantum spin Hall states protected by spin rotation about the z axis,^[40,41] etc.

For TSM, the definition of topological invariant needs modification as we cannot have a well-defined conduction or valence bands throughout the BZ: at some points the two bands cross each other. Given a nodal structure, say a point node, in BZ, we first use an imaginary manifold to enclose without touching the node. On the enclosing manifold, the conduction and the valence bands do not touch each other, having a full direct gap. Therefore, topological invariants can be defined on the enclosing manifold and we identify this invariant as the topological invariant of the node inside. We use the example of Weyl point to illustrate this process,^[1] which is the same one used in numerics to calculate the monopole charge of a Weyl point.^[2,30] Given a Weyl point, we consider a sphere in k -space to enclose this point (here one needs to make sure that only one band crossing point is inside). Then since the bands are ‘gapped’ on the surface of the sphere, we can calculate its Chern number. When the Chern number is ± 1 , we know that the Weyl point has monopole charge of ± 1 ; if the Chern number is ± 2 , we know that the nodal point is actually a double-Weyl point^[42] with monopole charge of ± 2 . Another example we use is the Dirac point protected by C_{4v} (existing in, e.g., Cd₃As₂^[43]). Here since the Dirac point is only allowed to move along the k_z -axis in the BZ, we consider two points $p_{1,2}$ along the k_z -axis above and below the Dirac point, to enclose the Dirac point. At p_1 and p_2 , the conduction and the valence bands are separated, and each band is double degenerate with C_4 eigenvalues $e^{\pm i\pi/4}$ or $-e^{\pm i\pi/4}$. At $p_{1,2}$ we count the number of valence bands that have C_4 eigenvalue $e^{\pm i\pi/4}$, denoted by $N_{1,2}$, respectively. The topological invariant of the Dirac point is given by $z = N_1 - N_2$.^[46,47]

For TNLSMs, we have three types of enclosing manifolds, being zero-dimensional, one-dimensional, and two-dimensional, respectively. If the nodal line is fixed by symmetry (usually a mirror reflection symmetry) to some high-symmetry plane, we choose two points on the same plane, on different sides of the nodal line, respectively (see Fig. 2(a)). If the nodal line is not fixed to any high-symmetry plane, then we consider the two following enclosing manifolds: a loop that links with the nodal line (see Fig. 2(b)), and a sphere or torus that encloses the nodal line (see Fig. 2(c)). In the presence of multiple symmetries (i. e., several mirror planes), several nodal lines may coexist or even meet each other at several ‘nexus points’, whose topological protection, or the absence thereof, will not be discussed in this review for concision, but see Refs. [44] and [45] for related studies.

2.1. Nodal lines protected by mirror reflection symmetries

In real space, a mirror reflection symmetry can be defined as

$$M : (x, y, z) \rightarrow (x, y, -z), \quad (1)$$

whereas in momentum space

$$M : (k_x, k_y, k_z) \rightarrow (k_x, k_y, -k_z). \quad (2)$$

The symmetry can be represented by a unitary operator acting on one-electron wave functions, satisfying

$$M^2 = (-1)^{2S}, \quad (3)$$

where S is the spin of a single particle. From Eq. (3) we see an important distinction between spinless ($S = 0$) and spinful ($S = 1/2$) particles. For the former, M has eigenvalues ± 1 and for the latter, $\pm i$. Physically, this factor of i is due to the fact that in a spin–orbit coupled system, the reflection also acts on the spin degrees of freedom. Mark that spinful systems can be viewed as spinless systems when the full $SU(2)$ spin-rotation symmetry is preserved, because the spatial and spin degrees of freedom are decoupled. When a non-interacting system described by Hamiltonian $H(k_x, k_y, k_z)$ has mirror reflection symmetry M , we have

$$MH(k_x, k_y, -k_z)M^{-1} = H(k_x, k_y, k_z). \quad (4)$$

Note that $k_z \rightarrow -k_z$ on the left is due to that the mirror operation also flips the z -component of the momentum. By Eq. (4), we see that at the two high-symmetry planes in the BZ, namely $k_z = 0$ and $k_z = \pi$, the mirror operator and the Hamiltonian have the same eigenstates, such that we can use the eigenvalues of M to label the bands on these two planes. Without other symmetries, all bands are generically non-degenerate, and suppose there are two bands labeled by $+1$ and -1 respectively (assuming $S = 0$), then these two bands are disallowed

by mirror symmetry to hybridize with each other. Therefore, two bands with opposite mirror eigenvalues can cross each other at k -points satisfying

$$E_+(\mathbf{k}) = E_-(\mathbf{k}). \quad (5)$$

Since \mathbf{k} has two free components (k_z being fixed to 0 or π), in Eq. (5) we have two variables to satisfy one equation, meaning that the solution space is generically one-dimensional, i.e., a nodal line. The same discussion follows when $S = 1/2$. We emphasize that the two bands can cross only at $k_z = 0$ and $k_z = \pi$ planes: away from them, there is no quantum number to distinguish the bands or prevent hybridization.

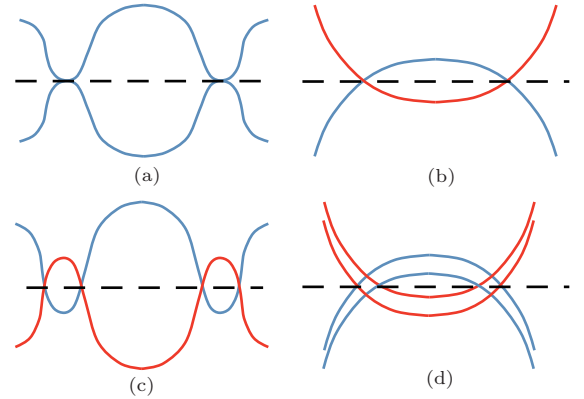


Fig. 1. (color online) Dispersion along a cut across the nodal lines protected by mirror reflection symmetries carrying different topological invariants. Different colors represent opposite mirror eigenvalues. (a) An accidental nodal line which has $\zeta_0 = 0$. (b) A protected nodal line carrying $\zeta_0 = 1$. (c) Two nodal lines (one inner and one outer) carrying opposite invariants. (d) Two nodal lines carrying the same invariant. [28]

Since the nodal line is fixed to the high-symmetry planes by mirror symmetry, we use the zero-dimensional enclosing manifold. On the two sides of the nodal line we pick two points p_1 and p_2 . At $p_{1,2}$, the conduction and the valence bands are separated in energy, and one can count the number of bands below the Fermi energy that has mirror eigenvalue of $+1$, denoted by $N_{1,2}$. The topological invariant is then given by

$$\zeta_0 = N_1 - N_2. \quad (6)$$

$\zeta_0 = 0$ corresponds to the case shown in Fig. 1(a), and it means that the crossing is only accidental and can be removed without breaking the mirror symmetry; $\zeta = 1$ corresponds to the case shown in Fig. 1(b), and it means that the crossing is between two bands that have opposite mirror eigenvalues. $\zeta = 2$ corresponds to the case shown in Fig. 1(d), and it means that there are two nodal lines resulting from two pairs of bands with opposite eigenvalues. Here we remark that the 0D invariant ζ_0 is a Z -invariant, not Z_2 -invariant. Suppose by tuning the parameters we can put two nodal lines at the same k -point in BZ, then if the invariant is Z_2 , the two nodal lines will ‘cancel’ each other and create a full gap; but if the invariant is Z ,

it depends on whether the two lines have the same or opposite invariants. If the invariants are the same, then the two lines will not gap each other (see Fig. 1(d)); and if the invariants are opposite, putting these nodal lines together will create a full gap (see Fig. 1(c)).

2.2. Nodal lines protected by inversion, time-reversal, and SU(2) spin-rotation symmetries

Here we first assume that all the three symmetries are present in our system. Since SU(2) is a symmetry, we can redefine time-reversal operator, combining it with a π spin rotation about the y -axis,

$$T \rightarrow T e^{i s_y \pi}, \quad (7)$$

after which we have $T^2 = +1$ instead of -1 for fermions. Since both inversion, P and T , reverse the momentum $\mathbf{k} \rightarrow -\mathbf{k}$, $P * T$ is an anti-unitary symmetry that preserves the momentum. Since $[P, T] = 0$, we have

$$(P * T)^2 = P^2 T^2 = 1. \quad (8)$$

Equation (8) dictates that it can be represented as

$$P * T = K, \quad (9)$$

where K is the complex conjugation, in a proper orbital basis. In this basis, $P * T$ -symmetry ensures that

$$H(\mathbf{k}) = H^*(\mathbf{k}), \quad (10)$$

or that $H(\mathbf{k})$ is real at each \mathbf{k} .

Away from the crossing lines, the Hamiltonian can be ‘flattened’ as

$$Q(\mathbf{k}) = I - \sum_{n \in \text{occ.}} |u_n(\mathbf{k})\rangle \langle u_n(\mathbf{k})|. \quad (11)$$

From Eq. (11), we see that the eigenfunctions of $Q(\mathbf{k})$ are the same as those of $H(\mathbf{k})$, but eigenvalues of $Q(\mathbf{k})$ are either zero or one, depending on whether n is an occupied or unoccupied band. Then, we note that on any compact manifold that does not contain any crossing point, the Hamiltonian $H(\mathbf{k})$ can always be deformed into $Q(\mathbf{k})$ without breaking any symmetry or closing the gap, i.e., $H(\mathbf{k})$ and $Q(\mathbf{k})$ have the same topology.

Before going further, a brief review of the concept of homotopy groups in algebraic topology is due. Consider continuous mappings from an n -sphere S^n to a space M (in this case the space of all occupied bands). If two such mappings, ϕ_1 and ϕ_2 , can be continuously deformed into each other, then we say that ϕ_1 and ϕ_2 are equivalent to each other. Mappings that are equivalent to each other form an equivalent class, and each class corresponds to an element in the homotopy group, denoted by $\pi_n(M)$. For example, if the homotopy group has

only one element, $\pi_n(M) = \{e\}$, then we know that all mappings from S^n to M are equivalent to each other, and therefore equivalent to a trivial mapping where all points in S^n map to the same point in M . If the homotopy group is Z_2 , $\pi_n(M) = Z_2$, then we know that all mappings are either equivalent to a trivial mapping or to a nontrivial mapping. This mathematical definition can be paraphrased in our physical context: $S^{n \geq 1}$ is our enclosing manifold; M is the Hilbert space spanned by all occupied bands; the Hamiltonian $H(\mathbf{k})$ is the mapping; and the condition that two mappings can continuously deform into each other corresponds to the case where $H_1(\mathbf{k})$ can be adiabatically transformed to $H_2(\mathbf{k})$ without gap closing. Therefore, the homotopy group of $Q(\mathbf{k})$ exactly gives the topological classification of the nodal line enclosed by the n -sphere.

If $Q(\mathbf{k})$ is real, it is an element of the real Grassmanian manifold, or

$$Q(\mathbf{k}) \in \frac{O(m+n)}{O(m) \oplus O(n)}. \quad (12)$$

It is known that the homotopy groups of this manifold are

$$\pi_1 \left(\frac{O(m+n)}{O(m) \oplus O(n)} \right) = \pi_2 \left(\frac{O(m+n)}{O(m) \oplus O(n)} \right) = Z_2, \quad (13)$$

for $m, n > 2$. Equation (6) means that if we enclose the nodal line with either a ring or a sphere, the topological classification of the wave functions on the ring/sphere is Z_2 . This means that for a nodal ring, we have two independent Z_2 indices (denoted by ζ_1 and ζ_2 , defined on a ring that links with the line (Fig. 2(b)) and on a sphere that encloses the whole line (Fig. 2(c)).

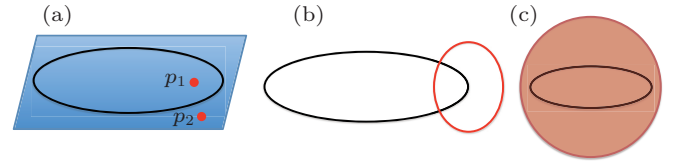


Fig. 2. (color online) Manifolds of different dimensions (S^0 , S^1 , and S^2) that enclose a nodal line. (a) Two points (S^0) inside and outside the nodal line pinned to (protected by) a mirror plane; (b) a loop (S^1) that threads the nodal line; and (c) a sphere (S^2) surrounding the entire nodal line. [28]

If ζ_1 is zero, we infer that the line crossing is purely accidental and can be removed by an arbitrarily small perturbation that preserves all symmetries. Therefore, all topological nodal rings protected by this symmetry group must all have $\zeta_1 = 1$. The explicit expression of ζ_1 is simply the Berry phase on the ring that links with the nodal line,

$$(-1)^{\zeta_1} = \oint d\mathbf{k} \mathbf{A}(\mathbf{k}) \cdot d\mathbf{k}, \quad (14)$$

where

$$\mathbf{A}(\mathbf{k}) = -i \sum_{n \in \text{occ.}} \langle u_n(\mathbf{k}) | \partial_{\mathbf{k}} | u_n(\mathbf{k}) \rangle \quad (15)$$

is the Abelian Berry connection. It can be proved that when $H(\mathbf{k})$ is real, the Berry phase associated with any loop must be quantized to either 0 or π . If it is zero, then we can smoothly

shrink this loop to a single point; but if it is π , the loop cannot shrink to a point, as an infinitesimal loop necessarily has zero Berry phase. In the latter case, there must be a point inside the loop where the Berry phase cannot be defined, where the conduction and the valence bands cross.

We have understood that $\zeta_1 = 0$ and $\zeta_1 = 1$ mean that the nodal ring is accidental and protected, respectively. What is the physical meaning of the second index? When the second index ζ_2 is zero, although the nodal line is stable against perturbations, the nodal line may still shrink to a point and be gapped by a continuous tuning of the Hamiltonian. This point can be illustrated by the following example. Consider an effective Hamiltonian near Γ

$$H(\mathbf{k}) = (m - k^2)\sigma_z + k_z\sigma_x, \quad (16)$$

where $P = \sigma_z$ and $T = K\sigma_z$ and of course $P * T = K$ as promised. If $m > 0$, the two bands cross each other on the $k_z = 0$ -plane, making a nodal circle of radius $r = \sqrt{m}$. It is obvious that as m decreases, the nodal circle shrinks, and at $m = 0$, it shrinks into a single point at $\mathbf{k} = 0$, and when we further decrease m to $m < 0$, the nodal circle vanishes (see Fig. 3(a)). This is a typical example when the second index of a nodal ring is zero. $\zeta_2 = 1$ means that on the surface of the sphere that surrounds the nodal line, the periodic part of the Bloch wave functions cannot be adiabatically tuned to the same function, and also means that one cannot shrink the sphere to a single point without meeting a singularity in the process. In this case, the nodal line cannot shrink to a point and be gapped out, which can be proved by contradiction: suppose the nodal line can vanish by tuning the Hamiltonian, one can first gap out the nodal line by shrinking it, so that there is no singularity inside the sphere, then one can shrink the sphere to a single point, contradicting the assumption that $\zeta_2 = 1$. We use the following example to illustrate a nodal line with $\zeta_2 = 1$.

$$H(\mathbf{k}) = k_x s_x + k_y \tau_y s_y + k_z s_z + m \tau_x s_x, \quad (17)$$

where τ_i and s_i are Pauli matrices acting on two isospin degrees of freedom. The spectrum is given by

$$E(\mathbf{k}) = \pm \sqrt{k_z^2 + (\sqrt{k_x^2 + k_y^2} \pm m)^2}. \quad (18)$$

The band crossing can be found by solving $E(\mathbf{k}) = 0$, yielding $k_z = 0$ and $\sqrt{k_x^2 + k_y^2} = |m|$, i.e., a nodal line on the xy -plane of radius $|m|$. As m changes from positive to negative, the radius decreases and shrinks to zero at $m = 0$ but increases again when m becomes negative (see Fig. 3(b)). Therefore, this nodal line cannot shrink to a point and then be gapped out by tuning the parameters.

One can draw an analogy between the ζ_2 of the nodal line and the monopole charge of a Weyl point, as both are defined on a sphere that encloses the nodal structure, with a key difference that here the monopole charge is a Z_2 -charge in the

former and Z -charge in the latter. This difference implies that when two nodal rings with $\zeta_2 = 1$ meet each other, they will necessarily annihilate each other, while two Weyl points with $C = 1$ will not annihilate each other, but form a double Weyl point. The Nelson–Nanomiya theorem also applies to this Z_2 -charge, which states that in a lattice model, the nodal lines with $\zeta_2 = 1$ must appear in pairs. This is another distinction between nodal lines with $\zeta_2 = 0$ and $\zeta_2 = 1$: while the ring can be annihilated or created in singles in the former case, in the latter case they can only be annihilated or created in pairs.

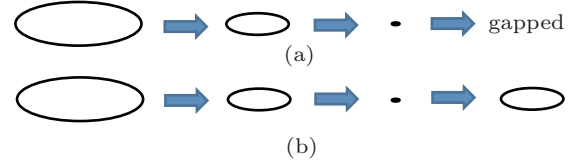


Fig. 3. (color online) (a) By tuning m from positive to negative in Eq. (16), the nodal line with $\zeta_2 = 0$ is fully gapped; (b) by tuning m from positive to negative in Eq. (17), the nodal line with $\zeta_2 = 1$ first shrinks to a point, then grows into a line again.^[28]

The explicit form of ζ_2 , in terms of the wave functions, is not very concise, and readers are encouraged to read the Appendices of Ref. [28] for more information. Here we give an intuitive understanding of this invariant. Since the Hamiltonian $H(\mathbf{k})$ is real, we can find for each \mathbf{k} -point a set of real eigenfunctions of $H(\mathbf{k})$, denoted by $|u_n(\mathbf{k})\rangle \in \text{Real}$. Then, a natural question is whether there is a choice of $|u_n(\mathbf{k})\rangle$ that are smooth on the entire sphere that encloses the nodal line. $\zeta_2 = 1$ means that this ‘smooth, real gauge’ does not exist.

Before closing this subsection, a few technical comments are due. (i) The first invariant ζ_1 was first identified as a Z -invariant in the literature,^[4] because the authors considered a two-band model, and $\pi_1(O(2)/O(1) \oplus O(1))$ is indeed Z . Physically, this means that if there are only two bands, two nodal lines of the same charge meeting each other will not gap out, but once more bands are introduced, they can meet with the two crossing bands and gap out the nodal line. (ii) Another special case is $\pi_2(O(4)/O(2) \oplus O(2)) = Z$, that is, when there are in total four bands and two conduction (valence) bands, the second index becomes a Z -index. This means that a nodal line may carry either positive or negative charges, and that two nodal lines with the same monopole charge will not annihilate each other, but when other bands are introduced into the model, the two nodal lines will be gapped when they meet each other.

2.3. Double-nodal lines protected by twofold screw rotation, inversion, and time-reversal

Double-nodal lines appear when both the conduction and the valence bands are doubly degenerate and they cross each other along a nodal line, where the bands are fourfold degenerate. From its definition, we see that we need a symmetry such

that the all bands are doubly degenerate, and then we need another symmetry to protect the band crossing.

Even for spin-orbit coupled systems, $P * T$ makes sure that all bands are doubly degenerate. In the presence of SOC, we cannot redefine T such that $T^2 = +1$, and therefore, we have

$$(P * T)^2 = -1. \quad (19)$$

Since $P * T$ preserves the momentum and is an anti-unitary operator, we can prove that all bands are doubly degenerate. To be specific, for any Bloch state $|\psi_n(\mathbf{k})\rangle$, $P * T|\psi_n(\mathbf{k})\rangle$ must be an eigenstate at \mathbf{k} that is orthogonal to $|\psi_n(\mathbf{k})\rangle$.

Intuitively, it is understood that it is harder for two degenerate bands to cross each other than for non-degenerate bands. Consider an effective theory near the band crossing, then one needs at least a four-band model for the former and a two-band model for the latter. A four-band model is expanded in fifteen Dirac matrices (the identity matrix having nothing to do with band crossings), while a two-band model in three Pauli matrices. Then a double-line nodal requires the coefficients of fifteen Dirac matrices to vanish and a single nodal line only requires the coefficients of three Pauli matrices to vanish. Corroborating with this intuition is the fact that mirror reflection symmetry $M : (x, y, z) \rightarrow (x, y, -z)$ is insufficient to protect the crossing.

For spinful systems, equation (1) gives that the mirror eigenvalues are either $+i$ or $-i$. Since $P * T$ commutes with M and $P * T$ is anti-unitary, $|\psi_n(\mathbf{k})\rangle$ and $P * T|\psi_n(\mathbf{k})\rangle$ have opposite mirror eigenvalues, i.e., the two degenerate bands have opposite mirror eigenvalues. Therefore, when two degenerate bands, say band one and band two, cross, the $+i$ -subband in band one will repel the $+i$ -subband in band two and similar repulsion exists between the $-i$ -subbands in the two bands, creating a full gap.

From this discussion, we may conjecture that in order for the two bands to have a protected crossing, we need the two subbands in a degenerate band to have the same quantum number of some symmetry. So far, one type of these symmetries have been found:^[28] the twofold screw rotation, which acts in real space as

$$R : (x, y, z) \rightarrow (-x, -y, z - c/2), \quad (20)$$

where c is the lattice constant in the z -direction. In the presence of inversion, the twofold screw rotation is equivalent to the following mirror plane $M' = P * R$:

$$M' : (x, y, z) \rightarrow (x, y, c/2 - z). \quad (21)$$

It is easy to see that the only distinction between M' and M defined in Eq. (1) is that the mirror plane of M' is located at $z = c/4$, while for M it is at $z = 0$. This offset of mirror plane

leads to the following commutation relation between P and M' :

$$P * M' = T_{00\bar{1}} M' * P, \quad (22)$$

where $T_{00\bar{1}}$ is the translation along the $-z$ -direction by one unit cell. At the BZ boundary $k_z = \pi$, $T_{00\bar{1}} = \exp(ik_z c) = -1$. Therefore, at $k_z = \pi$ -plane, we have

$$\{P * T, M'\} = 0. \quad (23)$$

Suppose on this plane, one subband of the conduction (or valence) bands have $M' = +i$, then the M' eigenvalue of the other subband is found to be the same, namely,

$$\begin{aligned} M'(P * T|i\rangle) &= -P * T(M'|i\rangle) = -P * T(+i|i\rangle) \\ &= +i(P * T|i\rangle), \end{aligned} \quad (24)$$

where we have used the fact that $P * T$ anti-commute with both M' and $+i$. The same steps show that if one subband has $M' = -i$, the other subband also has $M' = -i$. When the degenerate conduction and the valence bands have opposite M' -eigenvalues on the $k_z = \pi$ plane, the effective four-band Hamiltonian reads

$$H = \begin{pmatrix} \varepsilon_v(k_x, k_y, \pi) & 0 & 0 & 0 \\ 0 & \varepsilon_v(k_x, k_y, \pi) & 0 & 0 \\ 0 & 0 & \varepsilon_c(k_x, k_y, \pi) & 0 \\ 0 & 0 & 0 & \varepsilon_c(k_x, k_y, \pi) \end{pmatrix}. \quad (25)$$

To understand why it takes such a simple form, we first notice that M' symmetry requires the off-diagonal block to vanish, and $P * T = K(is_y)$ requires that the two diagonal blocks to be proportional to identity. Only one Dirac matrix out of the fifteen remains, and its coefficient is $(\varepsilon_v - \varepsilon_c)/2$. The band crossing appears at

$$\varepsilon_v(k_x, k_y, \pi) = \varepsilon_c(k_x, k_y, \pi), \quad (26)$$

which is one equation with two variables: the solution space is generically one-dimensional, i.e., a nodal line. Since both crossing bands are doubly degenerate, we call this crossing a double-nodal line.

The topological invariant for the double-nodal lines is very similar to the case of single nodal lines protected by mirror reflection. Choose two points on the two sides of the double-nodal line, p_1 and p_2 , and count the number of the occupied bands at $p_{1,2}$ that have $M' = +i$, and denote them by N_1 and N_2 . The Z -invariant is given by

$$Z = N_1 - N_2. \quad (27)$$

We have several technical comments before closing this subsection. To simplify our discussion, in Eq. (20) we choose the simplest form of the twofold rotation, which passes through the inversion center (origin). In fact, the screw axis can also be shifted to $x = a/4$ or $y = b/4$. In that case, the reflection symmetry $M' = P * R$ is not a mirror reflection but a glide reflection symmetry, because it also involves half-lattice

translation along the x - or y -direction. This will slightly complicate the proof that the two subbands have the same M' eigenvalue, but the statement itself remains unchanged,^[28] that the symmetry protects the crossing of two degenerate bands. However, it should be noted^[10] that the actual shape and number of the nodal lines depend on the choice of symmetry. In either case (mirror or glide plane away from the inversion center), the bands are required to appear in groups of twos (fours if counting spin) by symmetry, i.e., the $(2n-1)$ -th band and the $2n$ -th band always appear together. In the case of a mirror plane, these two bands cross each other along an even number of non-contractible lines in the BZ and in the case of a glide plane, the two cross each other along an odd number of non-contractible lines.^[10] The $2n$ -th band and the $(2n+1)$ -th band belong to different groups and do not have to cross, but when they do, they cross along an even number of nodal rings in the case of a mirror plane, and an odd number of nodal rings in the case of a glide plane.

3. Materials realization

Since TNLSM was proposed in 2011,^[4] there have been many material proposals to realize it experimentally. According to the classification in the former section, we summarize, to the best of our knowledge, the existing proposals in Table 1. The original proposal of Burkov *et al.*^[4] is based on a fine tuned superlattice of normal insulator and topological insulator with broken time reversal symmetry, which is actually a model rather than a realistic material. The first type-A NLSM (see caption of Table 1 for definitions of types) was proposed in the half-metallic double WSM HgCr_2Se_4 .^[6] When the magnetization is long the $[001]$ axis, there is a nodal line inside the $k_z=0$ mirror plane in addition to the double Weyl nodes on the $[001]$ axis.^[42] Such spinful NLSM has also been proposed in noncentrosymmetric TiTaSe_2 ^[11] and PbTaSe_2 ,^[12] where the bands are non-degenerate due to inversion symmetry breaking though time reversal symmetry is conserved. For spinless case, TaAs ,^[30,31] ZrTe ,^[14] and CaAgX ($X = \text{P}, \text{As}$)^[13]

have nodal line as protected by the mirror symmetry. In the first two materials, the nodal line decays into Weyl points when SOC is included while the last one becomes a fully gaped TI. In 2014, a kind of all carbon graphene network, so called Mackay–Terrones crystal (MTC),^[15] was proposed to be NLSM of type B and the Bernal graphite^[48,49] studied a decade ago was also revealed to be of this type. 3D honeycomb lattices, including the hyperhoneycomb and the stripyhoneycomb lattices, are also reported to host both nodal lines and nodal points.^[50] The other proposals for type B, including Ca_3P_2 ,^[16,17] $\text{Cu}_3(\text{Pd,Zn})\text{N}$,^[18,19] LaN ,^[20] Be and other alkaline-earth metal,^[21,22] CaTe ,^[23] BaSn_2 ,^[24] black phosphorus (BP) under pressure,^[25] CaP_3 ^[26] and other carbon based materials like interpenetrated graphene network (IGN)^[5] and body-centered orthorhombic C_{16} (BCO-C16).^[51] Among them, CaP_3 has the lowest crystal symmetry and the nodal line appears at arbitrary points in the momentum space, while the others have their nodal line(s) constrained in the mirror plane(s). SOC can be neglected for compounds composed of light elements. Presently, only Be metal with very tiny SOC has been confirmed to host nodal line by the ARPES experiment.^[22] For IGN in Ref. [5], the nodal line stretches over the whole BZ and connects to its image in the next BZ instead of forming a closed ring. For type C, SrIrO_3 ^[28] and BaMX_3 ($M = \text{V}, \text{Nb}$ and Ta , $X = \text{S}, \text{Se}$)^[29] have been proposed. In SrIrO_3 , the symmetry that protects the double nodal line is a glide plane and the crossing is between a $2n$ -th band and a $(2n+1)$ -th band, so that there is only one nodal ring around the U -point.^[52] In fact, SrIrO_3 should also be considered as a filling enforced semimetal,^[53] whose band crossing happens to be a nodal ring: the space group and time-reversal requires that all band insulators have filling number $8n$, while SrIrO_3 's filling is $8n+4$. BaMX_3 has been shown to host nodal surface when SOC is not taken into account. Different from the inner connecting of the three mutually perpendicular nodal rings in MTC and $\text{Cu}_3(\text{Pd}, \text{Zn})\text{N}$, the inter connecting of nodal rings can form nodal chain state, which has been proposed for IrF_4 family compounds.^[7]

Table 1. The proposed materials to host nodal lines classified. The DSM, WSM, and TI mean the nodal lines evolve into the corresponding topological state when spin–orbit coupling (SOC) is further included. N/A means unknown. Type-A means that the nodal lines are protected by mirror reflection symmetry; type-B means that the nodal lines are protected by space inversion and time-reversal symmetry; type-C means the existence of double-nodal lines.

Class	NO SOC	+SOC
Type A	TaTa, ZrTe	Weyl semimetal
	CaAgX ($X = \text{P}, \text{As}$)	Topological insulators HgCr ₂ Se ₄ , TiTaSe ₂ , PbTaSe ₂
Type B	CaP ₃	Topological insulators
	MTC, BaSn ₂ , BP, IGN, BCO-C ₁₆	Topological insulators
	Be and other alkaline-earth metal, Ca ₃ P ₂	N/A
	Cu ₃ (Pd, Zn)N, LaN, CaTe	Dirac semimetals
Type C		SrIrO ₃ , BaMX ₃ ($M=\text{V}, \text{Nb}, \text{Ta}$, $X=\text{S}, \text{Se}$)

4. Symmetry breaking scenarios

We have established several classes of nodal lines protected by different symmetry groups. When these symmetries are partially or fully broken, the nodal lines generically vanish with them. However, unlike a point node that can simply gap out, there are at least two fates for a nodal line: it may break into several discrete nodal points, or it may also be fully gapped. In this section, we show (i) how a nodal line protected by mirror reflection and SU(2) spin rotation breaks into several Weyl points or Dirac points if SU(2) is broken and (ii) how the double-nodal line proposed in SrIrO₃ breaks into two nonsymmorphic Dirac points when one glide reflection symmetry is broken.

4.1. Nodal line broken into point nodes by SOC

As we have discussed at the end of Section 2, in a spinless (or SU(2)-symmetric) system, nodal line appears when there is a band inversion between two bands with opposite mirror eigenvalues. This has been seen in first principles calculations in many classes of materials. In reality, the spin-orbital coupling, while small, cannot be completely ignored in electronic systems. When SOC is considered, the band repulsion between opposite spins becomes nonzero at finite momenta, making the nodal line unstable. Depending on the remaining symmetries (other than SU(2)) of our system, the nodal line can either break into several pairs of Weyl points, one or several Dirac points, two separate nodal lines, or become fully gapped. In TaAs and several materials of the same family, it is found that a nodal line in the non-SOC band structure breaks into three pairs of Weyl points; in LaN, Cu₃(Pd,Zn)N and CaTe, three intersecting nodal lines break into two Dirac points. In noncentrosymmetric materials, a nodal line in the non-SOC band structure usually signifies Weyl points when SOC is turned on.

How a nodal line breaks into Weyl points can be understood in simple effective models, but the quantitative results (say how many pairs of Weyl points and their location) will differ from those from the first-principles calculations. Consider an effective Hamiltonian for TNLSM protected by mirror reflection and SU(2)

$$H_0 = (m - k^2)\sigma_z s_0 + k_z \sigma_y s_0, \quad (28)$$

where $\sigma_{0,i}$ acts on the orbital index (for example, one orbital may be s-orbital and the other p_z), and $s_{0,i}$ acts on the spin. Note that here only s_0 appears due to SU(2) symmetry. The symmetries are represented by $M = i\sigma_z s_z$ and $T = K(is_y)$. When SU(2) is broken, we can add spin-orbital terms

$$H = H_0 - m' \sigma_y s_x + k_x \sigma_x s_x, \quad (29)$$

whose dispersion takes the form of

$$E(\mathbf{k}) = \sqrt{(m - k^2)^2 + (k_z \pm m')^2 + k_x^2}. \quad (30)$$

For $m > m'^2$, this dispersion has two positive Weyl points at

$$\begin{aligned} W_{1+} &= (0, \sqrt{m - m'^2}, m'), \\ W_{2+} &= (0, -\sqrt{m - m'^2}, -m') \end{aligned} \quad (31)$$

and two negative Weyl points at

$$\begin{aligned} W_{1-} &= (0, -\sqrt{m - m'^2}, m'), \\ W_{2-} &= (0, \sqrt{m - m'^2}, -m'). \end{aligned} \quad (32)$$

At the point $m = m'^2$, W_{i+} and W_{i-} annihilate each other, and the system becomes fully gapped. In the second example, we have mirror reflection symmetry about the xz - and yz -planes, and we assume there is fourfold rotation symmetry about the z -axis and inversion symmetry

$$H_0 = (m - k^2)\sigma_z s_0 + k_x k_y \sigma_x s_0, \quad (33)$$

and the symmetries are represented by $C_4 = \sigma_z \exp(-is_z \pi/4)$, $M_{xz} = i\sigma_z s_y$, $M_{yz} = i\sigma_z s_x$, and $T = K(is_y)$. Solving for the energy dispersion of Eq. (33) we see that there are two nodal lines, which are the intersection between the $k_x k_z$ -plane and the sphere of $k = \sqrt{m}$, and the $k_y k_z$ -plane and the sphere. Then we add the SOC terms

$$H = H_0 + k_y \sigma_x s_y - k_x \sigma_x s_x, \quad (34)$$

and the dispersion becomes

$$E(\mathbf{k}) = \sqrt{(m - k^2)^2 + (\sqrt{k_x^2 + k_y^2} \pm k_x k_y)^2}. \quad (35)$$

This dispersion only has two point nodes at $\mathbf{k} = (0, 0, \pm\sqrt{m})$, and at each point, all four bands meet at the same point, i.e., the two points are Dirac points.

4.2. Double nodal line broken into nonsymmorphic Dirac points

SrIrO₃ is the first proposed material that has a double-nodal line, protected in this particular case by a twofold screw rotation about the b -axis $S_y : (x, y, z) \rightarrow (-x + a/2, y + b/2, -z + c/2)$ as well as inversion and time-reversal symmetries. The space group of the bulk material is generated by inversion $P : (x, y, z) \rightarrow (-x, -y, -z)$, S_y and $S_x : (x, y, z) \rightarrow (x + a/2, -y + b/2, -z)$. Now suppose we break S_y preserving P and S_x , the double-nodal line is no longer protected. However, one can prove that along UR, the subbands in a degenerate band have the same eigenvalues of S_x . To see this, we note that

$$S_x^2 = -T_{100} = -e^{-ik_x}, \quad (36)$$

where the minus sign comes from the full rotation of the $1/2$ -spin. From Eq. (36), we find the eigenvalues of S_x to be

$\pm i e^{-ik_x/2}$. Then consider the commutation relation between P and S_x ,

$$S_x * P = T_{110} P * S_x = e^{-ik_x - ik_y} P * S_x. \quad (37)$$

Suppose $|\phi(\mathbf{k})\rangle$ is one subband of a degenerate band along UR of S_x eigenvalue $+i e^{-ik_x/2}$, then for the other subband

$$\begin{aligned} S_x(P * T|\phi(\mathbf{k})) &= -e^{-ik_x} P * T * S_x|\phi(\mathbf{k})\rangle \\ &= -e^{-ik_x} P * T(+i e^{-ik_x/2})|\phi(\mathbf{k})\rangle \\ &= i e^{-ik_x/2} P * T|\phi(\mathbf{k})\rangle. \end{aligned} \quad (38)$$

Equation (38) shows that, as promised, the two subbands have the same eigenvalue of S_x . Therefore, if two degenerate bands have opposite S_x eigenvalues, they may cross each other at a Dirac point. The Dirac points protected by the twofold screw axis and inversion are distinct from normal Dirac semimetals in that they have topologically protected surface states (double-helicoid surface states in this case).

5. Physical consequences

For most topological materials, the observation of topological surface states has been considered a definitive confirmation of the nontrivial topology in the band structure. The underpinning of this logic is the bulk-edge correspondence principle, which asserts that a nontrivial bulk topology in d -dimensional bulk must correspond to a gapless mode ($d-1$)-dimensional edge, which cannot be realized in a real ($d-1$)-dimensional system without symmetry breaking. Here we emphasize the prerequisites of its application that (i) the symmetry group protecting the topology in the bulk must be unbroken on the edge and (ii) the interaction is weak or the edge can still be gapped into an anomalous topology order.

So far, the protection of nodal lines requires one or several of the following spatial symmetries: mirror reflection, space inversion, and twofold screw rotation. A simple inspection of them shows that there is no surface where any of these symmetries is preserved. Therefore, the bulk-edge correspondence may not be applied here to indicate the existence of topological surface modes in TNLSMs. In numerical simulations. However, there are indeed states localized on the surface in the surface BZ, appearing inside the projections of the nodal lines. Unlike the surface states of, say, topological insulators, these surface states are very flat in dispersion, and are hence dubbed ‘drumhead’ states. The drumhead states can be considered as a higher-dimensional analogy of the flat band on the zigzag edge of graphene, and are hence dubbed ‘drumhead’ states, which may host interesting correlation effects due to the high ratio of interaction and kinetic energy.^[54] As we have stated, the flat bands are not topologically protected: a change of the model parameters on the surface will destroy the ‘flatness’ of the surface modes, and can even push these surface states into

the bulk continuum spectrum. In the same way the flat band on the zigzag edge gains dispersion as soon as we turn on the intra-sublattice hopping on the edge. The lack of a topological signature on the surface poses a challenge to design a ‘smoking gun’ experimental observation of TNLSM. In Refs. [11] and [12], using ARPES, the group measures the dispersion of both the bulk and the surface states, where the results support the existence of a nodal line protected by mirror reflection symmetry in PbTaSe₂ and TiTaSe₂.

Due to the lack of surface signatures in TNLSMs, people turn to the bulk probes. In quantum oscillation, the special behavior of these materials has been predicted and measured. In Ref. [55], the Landau levels of an effective $\mathbf{k} \cdot \mathbf{p}$ model near a double-nodal line is calculated as a function of the strength and the angle of the magnetic field. It is predicted that there are zero modes in the spectrum that lead to a peak in density of states at the Fermi level. In Ref. [56], the Landau levels are calculated in a lattice model describing a 3D honeycomb lattice that has nodal rings in the BZ. A key distinction between this work and Ref. [55] is that here the magnetic field is in the toroidal direction, where the field lines are parallel to the nodal ring. It is found that the Hall conductance, $\sigma_{\rho\rho}$ (ρ means the radial direction in cylindrical coordinates) is quantized so that a 3D quantum Hall effect can be realized. In Ref. [33], the de Haas–van Alphen oscillation is measured in TNLSMs ZrSiSe and ZrSiTe. The authors use the angle-dependent oscillation frequencies to map out the Fermi surfaces in these two materials and by fitting the Lifshitz–Kosevich formula, Berry phase is found to be $\sim 0.31\pi$ for out-of-plane and $\sim \pi$ for in-plane field (the latter value matches the theoretical prediction).

The ideal Fermi surface of a TNLSM is the nodal ring itself at half-filling, which may be achieved only if there is particle-hole symmetry or chiral symmetry that pins the energy of all the points on the nodal line at the Fermi energy. The dispersion near the Fermi surface is also particular in a TNLSM: while the band splitting along the nodal line is zero, the dispersion perpendicular to the nodal line is linear in momentum. These special properties near the Fermi energy lead to new many-body effects for TNLSMs. In Ref. [34], it is predicted that the Coulomb interaction is only partially screened and still long-ranged due to the vanishing density of states at the Fermi energy. In Ref. [35], the authors using renormalization group method analyze the quantum phase transition between a topological nodal line superconductor and a fully gapped superconductor, where the symmetry protecting the nodal line is broken. It is predicted that this transition belongs to a new universality class, where the dynamic exponent, order parameter exponent, and susceptibility exponent are all 1.

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