Hierarchy of orientational phases and axial anisotropies in the gauge theoretical description of generalized nematic liquid crystals

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The paradigm of spontaneous symmetry breaking encompasses the breaking of the rotational symmetries $O(3)$ of isotropic space to a discrete subgroup, i.e., a three-dimensional point group. The subgroups form a rich hierarchy and allow for many different phases of matter with orientational order. Such spontaneous symmetry breaking occurs in nematic liquid crystals, and a highlight of such anisotropic liquids is the uniaxial and biaxial nematics. Generalizing the familiar uniaxial and biaxial nematics to phases characterized by an arbitrary point-group symmetry, referred to as generalized nematics, leads to a large hierarchy of phases and possible orientational phase transitions. We discuss how a particular class of nematic phase transitions related to axial point groups can be efficiently captured within a recently proposed gauge theoretical formulation of generalized nematics [K. Liu, J. Nissinen, R.-J. Slager, K. Wu, and J. Zaanen, Phys. Rev. X 6, 041025 (2016)]. These transitions can be introduced in the model by considering anisotropic couplings that do not break any additional symmetries. By and large this generalizes the well-known uniaxial-biaxial nematic phase transition to any arbitrary axial point group in three dimensions. We find in particular that the generalized axial transitions are distinguished by two types of phase diagrams with intermediate vestigial orientational phases and that the window of the vestigial phase is intimately related to the amount of symmetry of the defining point group due to inherently growing fluctuations of the order parameter. This may explain the stability of the observed uniaxial-biaxial phases as compared to the yet to be observed other possible forms of generalized nematic order with higher point-group symmetries.

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I. INTRODUCTION

“Vestigial” or “mesophases” of matter are a well-established part of the canon of spontaneous symmetry breaking [1]. It might well happen that due to thermal [2] (or even quantum [3]) fluctuations a phase is stabilized at intermediate temperatures (or coupling constant at $T = 0$) characterized by a symmetry intermediate between the high-temperature isotropic phase and the fully symmetry broken phase at low temperature (small coupling constant). Iconic examples are liquid crystals [2], occurring in between the high-temperature liquids and the low-temperature crystals, characterized by only the breaking of the rotational symmetry (“nematics”), followed potentially by a partial breaking of translations (“smectic” or “columnar” phases) before full solidification sets in.

In the general sense of phases of matter that break the isotropy of Euclidean three-dimensional space, crystals are completely classified in terms of space groups. Nematics, on the other hand, are in principle classified in terms of all subgroups of O(3): the family of three-dimensional (3D) point groups. There are a total of seven infinite axial families and large this generalizes the well-known uniaxial-biaxial nematic phase transition to any arbitrary axial point group in three dimensions. We find in particular that the generalized axial transitions are distinguished by two types of phase diagrams with intermediate vestigial orientational phases and that the window of the vestigial phase is intimately related to the amount of symmetry of the defining point group due to inherently growing fluctuations of the order parameter. This may explain the stability of the observed uniaxial-biaxial phases as compared to the yet to be observed other possible forms of generalized nematic order with higher point-group symmetries.

Another well-established form is the “biaxial nematic” formed from platelets with three inequivalent director axes, characterized by the $D_{2h}$ point-group symmetry [5–13]. $D_{2h}$ is a subgroup of $D_{2hh}$, and it is well understood that the uniaxial nematic can be a vestigial mesophase that can occur in between the isotropic and biaxial phase. In order for such vestigial rotational sequences to occur, special microscopic conditions are required: dealing with molecule-like mesogenic constituents, special anisotropic interactions have to be present.

More concretely, in terms of a theory with lattice regularization, the degrees of freedom of the coarse-grained orientational constituents can be parametrized in terms of an O(3)-rotation matrix $R = (\{l_{i}, m_{i}, n_{i}\})$, i.e., an orthonormal triad $n_{i}^{\alpha} = \{l_{i}, m_{i}, n_{i}\}_{\alpha=1,2,3}$ in the body-fixed frame of the mesogen [14]. The orientational interaction between the mesogens is in general determined by their relative orientation of nearest neighbor sites $i,j$ and therefore a function of the relative direction cosines, i.e., $H_{ij} \sim -\text{Tr}[R_{i}^{T}J R_{j}] = -\sum_{\alpha\beta} J^{\alpha\beta} n_{i}^{\alpha} \cdot n_{j}^{\beta}$, where $J^{\alpha\beta}$ is a symmetric matrix; see Fig. 1. It turns out that without loss of generality this matrix can be diagonalized and the eigenvalues $J_{1}, J_{2}, J_{3}$ of $J$ characterize the interaction in terms of three perpendicular axes. Furthermore, the local axes $n_{k}^{\alpha} = \{n_{k}^{\alpha}, n_{k}^{\beta}, n_{k}^{\gamma}\}_{k=\langle i\mid j\rangle}$ are identified under the local point-group symmetries $\Lambda_{i} \in G$ in their body-fixed frame as $n_{k}^{\alpha} \simeq \Lambda_{G_{k}}^{\alpha\beta} n_{k}^{\beta}$ and the form the matrix $J$ is constrained by the point-group symmetry $G$ of the mesogens; see Sec. III. It is the case that the point groups are classified into two classes: the seven finite polyhedral groups $T, T_{h}, T_{d}, O, O_{h}, I, I_{h}$ that allow for only an isotropic $J = J_{3}$ and the seven infinite families of groups $C_{n}, C_{n}v, C_{n}h, S_{2n}, D_{n}, D_{nh}, D_{nd}$, where anisotropy in $J_{1} - J_{2} - J_{3}$ should be in general expected since it is allowed by the symmetries.
Aside from the uniaxial $D_{\infty h}$-nematic with a single director axis, the main focus regarding such anisotropies has been on a particular point-group symmetry generalizing the uniaxial ordering to three dimensions: the biaxial $D_{2h}$ "platelet" with three inequivalent director axes. The expectation is then that the biaxial phase is stabilized by sufficient anisotropy in the constituents and/or interactions [5,6,13,15].

These point groups have been the main focus of attention in mesogenic systems, and we are aware of only a few other additional point groups that have been considered in similar detail. That is, besides the $D_{2h}$ symmetry, only mesophases of $C_{2v}$ point-group-symmetric "banana-shaped" constituents have recently been investigated in some detail [12,16] in experimental systems, subsequently followed by theoretical considerations [17–19], as well as theoretical studies of other mesogenic symmetries [20–22]. However, in these systems the $C_{2v}$ constituents seem to organize into complicated mesogenic aggregates in the observed liquid crystals, thereby many of the systems form columnar and smectic phases [16].

As we will discuss in detail in the next section, the symmetry structure and anisotropic interactions that are behind the $D_{2h}$ uniaxial-biaxial phase descendence are actually perfectly compatible with all axial groups! As a consequence, the generalization of the special uniaxial-biaxial type of vestigial symmetry lowering is possible for this vast number of symmetries. In fact, the axial groups roughly divide into two subclasses in this particular regard. $D_{2h}$ belongs to the symmetry classes that are characterized by a horizontal mirror plane, and the $J_1$-$J_2$-$J_3$ type of anisotropy allows for just a single vestigial phase where fluctuations restore rotational symmetries in the mirror plane, which is always $D_{\infty h}$, the uniaxial nematic. However, in the other case, such a mirror plane is lacking, and we show in Sec. II that this makes possible a second generic "biaxial" phase with an extra mirror symmetry along the main axis compared to the original low-temperature biaxial phase.

Aside from pure symmetry considerations, the next question is how do the stability of the vestigial phase(s) and the fully ordered phase depend on general conditions such as the couplings and the nature of the point-group symmetry of the constituents? As we discussed elsewhere in much detail [23], the order parameter theories of "generalized nematics" characterized by symmetries beyond the simple $D_{\infty h}$. $D_{2h}$ are barely explored. The difficulty is with the complicated tensor structure of these order parameters. We introduced an extremely convenient mathematical formalism, borrowed from high-energy physics, to address these matters: $O(3)$ matrix matter coupled to discrete non-Abelian point-group $G$ gauge theory. On the technical side, the gauge-theoretic framework is a convenient device to construct the explicit order parameter tensors [23], but we also found that it is remarkably powerful to address the order-out-of disorder physics behind the occurrence of the vestigial phases [14]. We found thermal fluctuations of unprecedented strength lowering the transition temperatures to very low values in case of the most symmetric point groups ($T$, $O$, $I$), giving rise to a natural occurrence of a spontaneous vestigial chiral phase dealing with chiral point groups. How does this motive relate to the present context of "generalized" uniaxial-biaxial sequences?

It is actually the case that the $J_1$-$J_2$-$J_3$ type of anisotropy that arises in the gauge theory allows one to incorporate the generalized biaxial transitions in a natural manner, thereby making it possible to study such transitions with remarkable ease. We will discuss this in more detail in Sec. III how to use the gauge theory to compute quantitative phase diagrams. As expected, we recover the generic topology of the phase diagrams as a function of the anisotropy parameters. The advantage is that in the gauge theory one can compare apples with apples and pears with pears in the sense that the strength of the microscopic interactions including their anisotropy can be kept the same, facilitating a qualitative comparison of the phase diagrams for different point-group symmetries. The conclusion is that the stability region of the vestigial uniaxial phase grows rapidly as a function of increasing symmetry of the point group, which suppresses the fully ordered generalized biaxial phases considerably.

This mirrors the general motive that we already identified in the context of the chiral vestigial phases [14]: for the more symmetric point groups the thermal fluctuations grow in severity. This has on the one hand the effect of suppressing the ordering temperature of the fully ordered generalized biaxial phases, while at the same time the vestigial phase to a degree profits from the thermal fluctuations. As we will further discuss in the conclusion section, this raises the question whether for systems made from constituents characterized by highly symmetric point groups it will be ever possible to find the fully ordered phases before other mesophases and/or solidification sets in (these are beyond the description of our orientational lattice model). Any microscopic anisotropy might well render the vestigial uniaxial phase to be the only one that can be realized.

The remainder of this paper is organized as follows. In Sec. II we discuss the possible axial nematic phase transitions in terms of symmetries. For a realization of these phase
transitions, we review the lattice gauge theory model and define its anisotropic coupling parameters in Sec. III. Section IV is devoted for the phase diagrams and phase transitions obtained in Monte Carlo simulations. We conclude with an outlook in Sec. V.

II. THE STRUCTURE OF NEMATIC ORDER PARAMETERS AND GENERALIZED BIAXIAL TRANSITIONS

Three-dimensional generalized nematics break the rotational group O(3) down to a 3D point group. By the Landau–de Gennes symmetry paradigm, phase transitions between any two nematic phases related by the subgroup structure of O(3) are allowed, in addition to the transitions between the isotropic O(3) phase and a generalized nematic phase. In this section, we will show that the order parameter structure of axial nematics provides a natural way to realize some of the symmetry-allowed transitions. In Sec. III we then discuss how to realize these phase transitions by tuning the couplings in our gauge theoretical setup [14].

A. Point groups and nematic order parameters

Three-dimensional point groups are classified as seven finite polyhedral groups, \( \{T, T_d, T_h, O, O_h, I, I_h\} \), and seven infinite families of axial groups, \( \{C_n, C_{nv}, S_{2n}, C_{nh}, D_n, D_{nh}, D_{nd}\} \) [24,25]. The associated nematic order parameters are tensors that are invariant under the given point-group symmetry. A full classification of these order parameters and their derivation is given in our recent paper [23]. For the present purposes we therefore review the results that are of importance in the following.

Three-dimensional orientation can be parametrized in terms of a O(3) matrix

\[
R = (1 \ m \ n)^T.
\]

The rows \( n^\sigma = [l, m, n] \) of \( R \) form an orthonormal triad and satisfy the additional O(3) constraint

\[
\sigma = \det R = \epsilon_{abc}(l \otimes m \otimes n)_{abc} = l \cdot (m \times n) = \pm 1,
\]

where \( \sigma \) is the chirality or handedness of the triad \( n^\sigma \) associated with \( R \).

The order parameter tensors are constructed from tensor products of \( R \), and we use the point-group conventions of Ref. [23]. In case of the polyhedral nematics \( G = \{T, T_d, T_h, O, O_h, I, I_h\} \), the general form of the order parameter is given by \( \mathcal{O}^G = \{\mathcal{O}^G[l \ m \ n], \sigma\} \), where \( \mathcal{O}^G[l \ m \ n] \) describes the orientational order of the phase and \( \sigma \) is a chiral order parameter needed for the proper polyhedral groups \( \{T, O, I\} \). The polyhedral groups have several higher order rotation axes and transform the triads \( [l, m, n] \) irreducibly, and in these cases we need only one tensor to describe the orientational order [23].

On the other hand, the axial groups \( \{C_n, C_{nv}, S_{2n}, C_{nh}, D_n, D_{nh}, D_{nd}\} \) are defined with respect to a symmetry plane involving rotations and/or reflections and a perpendicular, axial direction. Their irreducible representations are in general one- or two-dimensional. Correspondingly, the order parameter tensors of the axial point groups have the general structure

\[
\mathcal{O}^G = \{A^G, B^G, \sigma\},
\]

where \( A^G \) defines the ordering related to the orientation of the primary axial axis perpendicular to the symmetry plane and \( B^G \) describes the in-plane ordering. We refer to \( A \) as the axial order and \( B \) as the in-plane (or just biaxial) order [23]. Similarly, \( \sigma \) is the chiral ordering for the proper axial groups \( \{C_n, D_n\} \). Note that the O(3) constraints can reduce the number of independent order parameter tensors in the set \( \{A^G, B^G, \sigma\} \) [23]. Following the conventions in Ref. [23], \( n \) is chosen always to be along the primary, axial axis. It follows that the axial order parameter tensor \( A^G \) depends only on \( n \) by construction. Similarly, the in-plane order parameter \( B^G = B^G[l \ m] \) depends only on \( l \) for the symmetries \( G = \{C_n, C_{nv}, C_{nh}, D_n, D_{nh}\} \) but is a tensor polynomial \( B^G = B^G[l \ m, n] \) of all the three triads for the symmetries \( \{S_{2n}, D_{nd}\} \) with rotoreflections. We have discussed these ordering tensors in Ref. [23], but for the convenience of readers, we show the relevant selection of order parameter tensors for the axial groups in Table III.

Moreover, because of the common structure of the axial point groups, the tensors \( A^G \) and \( B^G \) are not unique to a given symmetry, though the axial point group ordering can be uniquely defined by the full set of order parameters \( \{A^G, B^G, \sigma\} \). For instance, the symmetry groups \( C_n \) and \( C_{nv} \) do not transform the primary axis \( n \), thus the axial ordering for symmetries in these types is simply a vector,

\[
A^{C_n}[n] = A^{C_{nv}}[n] = A^{C_{nh}}[n] = n,
\]

where \( C_{nv} \equiv \text{SO}(2) \) is the continuous limit of \( C_n \) and \( C_{nh} \equiv \text{O}(2) \) is the continuous limit of \( C_{nv} \). The symmetries \( \{S_{2n}, C_{nh}, D_n, D_{nh}, D_{nd}\} \), however, transform \( n \) to \( -n \) and therefore have the same axial ordering tensor

\[
A^{D_{nh}}[n] = A^{D_{nh}}[n] = A^{D_{nh}}[n] = A^{D_{nh}}[n] = A^{D_{nh}}[n] = A^{D_{nh}}[n] = n \otimes n = - \frac{1}{2} \mathbb{I},
\]

which is just the well-known director order parameter of \( D_{nh} \)-uniaxial nematics. Note that \( D_{nh} \) can be considered as the continuous limit of the finite groups \( D_{nh} \), and \( D_{nd} \), whereas \( C_{nh} \) arises from the limit of \( C_{nh} \). Similarly, axial nematics with the same \( n \)-fold in-plane symmetries have the same ordering tensor \( B^G \):

\[
B^{C_n}[l \ m] = B^{C_{nh}}[l \ m],
\]

\[
B^{C_{nv}}[l \ m] = B^{D_n}[l \ m] = B^{D_{nh}}[l \ m].
\]

Note that, though the axial and the biaxial ordering tensors are distinct and transform irreducibly, they are not completely independent due to the O(3) constraints of orthonormality and Eq. (2).

B. Generalized biaxial phases and transitions

The distinction between the primary axis \( n \) and the in-plane axes \( l \) and \( m \) for axial nematics allows the disordering of the axial and in-plane order separately.

A familiar example is the biaxial-uniaxial-isotropic liquid transitions of \( D_{2h} \)-biaxial liquid crystals [5,6,26–29]. The order parameter tensors of the \( D_{2h} \) nematic are defined by two linearly independent rank-2 tensors, \( \mathcal{O}^{D_{2h}} = \{A^{D_{2h}}[n], B^{D_{2h}}[l \ m]\} \), where \( A^{D_{2h}}[n] \) has been given in Eq. (4), and
TABLE I. Generalized biaxial phase transitions. The first column specifies the generalized nematic symmetries, and the second column the minimal set of order parameter tensors for their characterization. Relations of the order parameters given by Eqs. (3)–(5) are indicated. For the specific forms of these order parameters see Ref. [23]. The third and fourth columns show the order parameter tensors involved in the generalized biaxial-uniaxial transitions in Eq. (8) and the biaxial-biaxial* transitions in Eq. (11), respectively. The symbol “→” indicates the replacement of an order parameter that becomes nonvanishing for the higher symmetry biaxial* (or uniaxial*) phases.

| Symmetry | Order parameters | Uniaxial-biaxial transitions | Biaxial-biaxial* (uniaxial-uniaxial*) transitions |
|----------|------------------|-------------------------------|--------------------------------------------------|
| $C_n$    | $A^{C_n} = A^{C_{n,∞}} [n], B^{C_n} = B^{C_{nh}} [l,m,n]$ | $B^{C_{n,∞}} [l,m,n], \sigma$ | $A^{C_{n,∞}} [n] \rightarrow A^{D_{nb}} [n], \sigma$ |
| $C_{nv}$ | $A^{C_{nv}} = A^{C_{n,∞}} [n], B^{C_{nv}} = B^{D_{nb}} [l,m]$ | $B^{D_{nb}} [l,m]$ | $A^{C_{n,∞}} [n] \rightarrow A^{D_{nb}} [n]$ |
| $S_{2n}$ | $A^{S_{2n}} = A^{D_{nh}} [n], B^{S_{2n}} [l,m,n]$ | $B^{S_{2n}} [l,m,n]$ | $B^{D_{nh}} [l,m,n] \rightarrow B^{C_{nh}} [l,m]$ |
| $C_{nh}$ | $A^{C_{nh}} = A^{D_{nh}} [n], B^{C_{nh}} [l,m]$ | $B^{C_{nh}} [l,m]$ | None |
| $D_n$    | $A^{D_n} = A^{D_{nh}} [n], B^{D_n} = B^{D_{nh}} [l,m,n], \sigma$ | $B^{D_{nh}} [l,m,n], \sigma$ | None |
| $D_{nh}$ | $A^{D_{nh}} = A^{D_{nh}} [n], B^{D_{nh}} [l,m,n]$ | $B^{D_{nh}} [l,m,n]$ | None |
| $C_{2n}$ | $A^{C_{2n}} [n]$ | None | $A^{C_{n,∞}} [n] \rightarrow A^{D_{nb}} [n]$ |
| $D_{2n}$ | $A^{D_{2n}} [n]$ | None | None |

$B^{D_{nh}} [l,m] = l \otimes 1 - m \otimes m$. (6)

In terms of the symmetries, the biaxial nematic order allows for the phase transitions

$$D_{2n} \rightarrow D_{2nh} \rightarrow O(3),$$

with the uniaxial phase occurring before the isotropic liquid. That is, upon increasing temperature, the biaxial order is destroyed, first leading to the restoration of the in-plane O(2) symmetry of uniaxial nematics before the transition to the fully disordered isotropic phase takes place.

Given the general order parameter structure of axial nematics discussed in Sec. II A, this transition sequence can be directly generalized to other axial symmetries. We will refer to the associated phase transitions as generalized biaxial transitions. By first destroying the in-plane order $B$, the following generalized biaxial-uniaxial transition can be induced:

$$C_n, C_{nv} \rightarrow C_{2n}.$$

Note that in these transitions we consider situations where the in-plane order has been completely disordered, leading to full O(2) symmetry. Thus the chiral order $\sigma$ for proper groups $C_n$ and $D_n$ has been simultaneously lost. Nevertheless, we can in principle also have the restorations of only the in-plane SO(2) symmetry with the transitions

$$C_n \rightarrow C_{∞}, \quad D_n \rightarrow D_{∞}. \quad (9)$$

where the chirality $\sigma$ does not disorder [14]. However, since $\sigma$ is a composite order parameter of $[l,m,n]$ featuring also some in-plane ordering, these transitions require more fine tuning in comparison to those in Eq. (8).

In the opposite limit, if the in-plane order with order parameter $B$ is sufficiently strong in comparison to the axial ordering $A[n]$, we can disorder the primary axis $n$ without destroying the in-plane order upon increasing the temperature. Note that due to the O(3) constraints on the triads, the axial ordering is never fully independent in the presence of the perpendicular in-plane ordering that fixes $n$ up to sign.

Therefore, upon disordering the axial order, the symmetry of the phase is augmented by

$$\sigma_b = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & -1 \end{pmatrix}, \quad (10)$$

which is a simply a reflection with respect to the $[l,m]$ plane that acts trivially on the in-plane ordering. Other symmetry operations transforming $n$ to $-n$, such as the inversion or a twofold rotations about an axis in the $[l,m]$-plane, however, will simultaneously transform the in-plane order. If such symmetries belong to the original symmetry group $G$, they will lead to enhanced in-plane symmetries in combination with $\sigma_b$. Therefore the new symmetries due to the disordering of the axial order $A^[G][n]$ are generated by the elements $G^* = G, \sigma_b$, leading schematically to either the direct product structure $G^* = G^* \times 1, \sigma_b$ or the semidirect structure $G^* = G^* \times 1, \sigma_b$, where $G^*$ can be an n-fold or 2n-fold rotational group. These are transitions between phases with different "biaxial" orders $B^G$ and $B^{G^*}$ will be for convenience referred to as biaxial-biaxial* transitions, where the subscript in $G^*$ denotes the presence of the additional reflections in comparison with the low-temperature symmetries $G$. The behavior of the associated ordered in the generalized uniaxial-biaxial transitions Eq. (8) and biaxial-biaxial* transition are summarized in Table I.

More specifically, in the "biaxial-biaxial*" phase transition the disordering of the primary axis with order parameter $A^[G][n]$ will lead to the phase transition of the generalized nematics with symmetries $[C_n, C_{nv}, S_{2n}, D_n, D_{nh}, D_{2n}]$

$$C_n \rightarrow C_{nh}, \quad S_{2n} \rightarrow C_{2nh}, \quad C_{nv}, D_n \rightarrow D_{nh}, \quad D_{nh} \rightarrow D_{2nh}, \quad (11)$$

as follows from the subgroup structure of O(3). Since $\sigma_b$ is already contained in the groups $C_{nh}$ and $D_{nh}$, the biaxial* phase is not present for these nematics.

Indeed, we see that these transitions have more interesting features than the generalized uniaxial-biaxial transitions in
The matter fields \(\{R_i\}\) live on the sites of a cubic lattice and are O(3) matrices, as in Eq. (1). The gauge fields \(\{U_{ij}\}\) are elements of the point group \(G\) and live on the links \((ij)\). In the Hamiltonian, \(H_{\text{Higgs}}\) is a Higgs term [32] describing interactions between the matter fields \(R_i\) and gauge fields \(U_{ij}\), parametrized by the coupling matrix \(J\) determining how the local axes \(\{n^i\}\) are coupled; see Fig. 1. The Hamiltonian in Eq. (12) is invariant under local gauge transformations

\[
R_i \rightarrow \Lambda_i R_i, \quad U_{ij} \rightarrow \Lambda_i U_{ij} \Lambda^T_j, \quad \forall \Lambda_i \in G. \tag{15}
\]

which leads to the identifications

\[
R_i \simeq \Lambda_i R_i, \quad n^i \simeq \Lambda^αβ n^i_β, \quad \Lambda_i \in G. \tag{16}
\]

Thus \(H_{\text{Higgs}}\) effectively models the orientational interaction between two \(G\)-symmetric “mesogens” [14]. In addition, \(H_{\text{Higgs}}\) has the global O(3)-rotation symmetry

\[
R_i \rightarrow R_i \Omega^T, \quad \Omega \in \text{O}(3). \tag{17}
\]

Since gauge symmetries cannot be broken [33], the fully ordered Higgs phase of \(H_{\text{Higgs}}\) will develop long-range order characterized by \(G\)-invariant tensor order parameters and thus realizes spontaneous symmetry breaking of Eq. (17) from an isotropic O(3) liquid phase to a generalized nematic phase [14, 23].

The term \(H_{\text{gauge}}\) in the Hamiltonian describes a point-group-symmetric gauge theory [34]. The term \(U_{ij} = \prod_{(ij) \in \square} U_{ij}\) denotes the oriented product of gauge fields around a plaquette \(\square\) and represents the local gauge field configuration on the lattice. Plaquettes with nontrivial flux \(U_{ij} \neq 1\) represent nonvanishing gauge field strength. Due to the gauge symmetries, gauge fluxes in the same conjugacy class are physically equivalent; therefore, the coupling \(K_{\text{C}}\) is a function on the conjugacy classes \(\text{C}\) of the group \(G\). These gauge fluxes are elements of the point group \(G\) and correspond to the Volterra defects in nematics [14, 35], and thus \(K_{\text{C}}\) equivalently assigns a finite core energy to the topological defects in the nematic [31]. However, for the purpose of realizing the generalized biaxial transitions in Eqs. (8) and (11), the Hamiltonian \(H_{\text{Higgs}}\) is sufficient, and for simplicity we will take \(K_{\text{C}} = 0\) in the following.

### B. Anisotropic couplings and generalized biaxial transitions

In order to analyze the Higgs interaction in terms of the nearest-neighbor local triads \(n^i = [1, m^i, n^i]\) and \(n^i\) identified under (16), we can define a local triad vector \(n^i_α = U_{ij}^{ij} n^i_α\) at a site \(j\), which has been brought (“parallel transported”) into the same local gauge as \(n^i_α\) at the site \(i\); see Fig. 1. In the gauge theory Eq. (15), each triad \(n^i\) represents a local frame of the mesogens and the gauge fields \(U_{ij}\) (elements of the point group) on the links encode the relative orientations of the local frames that are ambiguous up to the point-group symmetry of the mesogens. Therefore, in order to analyze the physical orientational interaction between the triads \(n^i\) and \(n^i\), we need to consider \(n^i \cdot n^i\) that correctly measures the relative orientation. This is mathematically known as the “parallel transport” of the triad in the gauge potential [34, 37] and is hardwired in the gauge theory. The Higgs interaction
This shows explicitly that the symmetric matrix $J_{ab}^{\rho\rho}$ parametrizes the interaction between the local triads; see Fig. 1. Naturally the interaction specified by the bilinear form $J_{ab}$ under a given 3D point-group gauge symmetry $\mathcal{G}$ only introduce a common sign in the $(\pm)$, since in the case of $C_1$ and $C_t \simeq S_2 = \{1, -1\}$, there are no rotational gauge symmetries $U_{ij}$ to begin with. It is easy to see that the monoclinic symmetries only introduce a common $\pm$ sign in the $\langle \textbf{l} \cdot \textbf{m} \rangle$ plane with the nondiagonal couplings. Therefore without loss of generality we can diagonalize the couplings,

$$J = \left( \begin{array}{ccc} J_1 & J_2 & J_3 \\ J_2 & J_1 & J \end{array} \right),$$

with $J_1, J_2, J_3 \geq 0$ for nematic alignment. For the monoclinic symmetries, this requires $J_{13} \leq \sqrt{J_1 J_2}$, and we do not consider negative or “antinematic” couplings [38,39]. We further note that the couplings also respect the symmetries of the auxiliary cubic lattice and favor alignment of the triads, leading to homogenous nematic states without any modulation or sublattice structure in the order parameters. Concerning the strength of alignment of the three perpendicular axes, the line of thought can actually be reversed in the sense that we can take couplings $J_1, J_2, J_3$ to be a measure of the effective three-dimensionality of the “mesogens” $R_i$. One realizes that they provide tuning parameters for the phase transitions involving the axial and in-plane ordering.

For the purpose of realizing the transitions in Eq. (8) and Eq. (11), we can consider the following form of $J$ for simplicity:

$$\beta J = \beta \left( \begin{array}{ccc} J_1 & J_1 & J_3 \\ J_1 & J_1 & J_3 \\ J_1 & J_1 & J_1 \end{array} \right),$$

where $J_1$ specifies the coupling of the in-plane degrees of freedom and $J_3$ the coupling between the primary axes. Therefore this form of $J$ is allowed for all axial groups and quantifies the anisotropy between the in-plane order and axial order, as was considered in Sec. II A in terms of the symmetries.

The fact that the phase transitions are tuned with respect to the temperature $\beta = 1/T$ reduces the the independent dimensionless couplings to two in terms of the reduced temperatures $\beta J_1$ and $\beta J_3$. Alternatively, we can consider the temperature $T$ as the tuning parameter in a thermotropic system and the anisotropy $\beta J_1$ as a fixed microscopic parameter. The ratio $\frac{J}{T}$ is in fact an analog to the so-called biaxiality parameter of $D_{2h}$ nematics [13,15,40,41]. Accordingly, when $\frac{J}{T}$ is sufficiently small, upon increasing temperature we expect that the in-plane order will be lost while the axial order still persist, leading to the generalized biaxial-uniaxial transition given in Eq. (8). In the opposite limit, where $\frac{J}{T}$ is sufficiently large, it is possible to disorder the axial order while the in-plane order is still maintained, leading to the generalized biaxial-biaxial transitions characterized by Eq. (11). Between these two limiting cases we expect direct transitions from the biaxial nematics to the O(3) isotropic liquid. Note, however, that in general the “biaxial” in-plane order is much more fragile than the uniaxial order of the primary, axial axis. Furthermore, the biaxial in-plane order reinforces the uniaxial order since it fixes the perpendicular axial order up to a sign. Conversely, the presence of the axial order reinforces the biaxial order much less, since ordering along $\textbf{n}$ still leaves in-plane SO(2) fluctuations before the full ordering sets in. As has been discovered in Ref. [14], the highly symmetric order parameter fields experience giant fluctuations and generalized biaxial nematics with a more symmetric in-plane structure require much larger $\frac{J}{T}$ to stabilize the in-plane order.
Nevertheless, although $\frac{J_3}{J_1}$ parameterizes the anisotropy of the in-plane and axial order of general biaxial nematics, they are defined in the gauge theory, so their values do not directly indicate the relative strength of the in-plane order and axial order. Therefore $\frac{J_3}{J_1} > 1$ does not necessary mean the in-plane order is favored and vice versa. Moreover, due to the O(3) constraints, naturally only two of the orthonormal triads are fully independent. In the gauge theoretical effective Hamiltonian terms respecting all the symmetries and the O(3) constraints, i.e., all gauge invariant combinations, appear order by order. That is, gauge invariant interactions such as $(\mathbf{l} \times \mathbf{m}_i) \cdot (\mathbf{l} \times \mathbf{m}_j) = \sigma_i \sigma_j \mathbf{n}_i \cdot \mathbf{n}_j$ or $(\mathbf{l} \cdot \mathbf{1})^2 + (\mathbf{l} \cdot \mathbf{m}_j)^2 + (\mathbf{m}_i \cdot \mathbf{m}_j)^2 \sim (\mathbf{n}_i \cdot \mathbf{n}_j)^2$ are present with coefficients parametrized by powers of $J_1$. Therefore, even though $J_3 = 0$, effective axial interactions $J_{3, \text{eff}}(J_1)\sigma_i \sigma_j \mathbf{n}_i \cdot \mathbf{n}_j$ or $J_{3, \text{eff}}(J_1)(\mathbf{n}_i \cdot \mathbf{n}_j)^2$ (pseudovector or uniaxial terms) are generated at all orders for all axial groups if allowed by the symmetries. In particular this affects higher order axial symmetries that have high rank order parameter tensors with large fluctuations. Among other things, due to the induced axial terms that are more relevant than the higher order in-plane interactions, the uniaxial (or uniaxial$^*$) phase is always stabilized before the biaxial (or biaxial$^*$) phase for in-plane symmetries with higher symmetries. The qualitative effect of these induced terms on the phase diagram is depicted in Fig. 3. We will see concrete examples how these induced interactions affect the numerical phase diagrams in Sec. IV.

C. Topology of the phase diagrams

Based on the discussions in Secs. II B and III B, we can now identify the topology of phase diagrams of biaxial nematics at different temperatures and anisotropies of $J$ as defined in Eq. (21). These are shown in Figs. 2 and 3. In Fig. 2 we show the conventional phase diagram in terms of the temperature

![FIG. 2. The schematic temperature-anisotropy phase diagram of axial nematics with conventional twofold biaxial symmetries. Small and large $\frac{\beta}{J}$ correspond to weak and strong in-plane order, respectively. $(\frac{\beta}{J})_U$ and $(\frac{\beta}{J})_V$ are the critical anisotropies where the generalized biaxial-uniaxial transitions in Eq. (8) and the biaxial-biaxial$^*$ transitions in Eq. (11) terminate, respectively. Solid lines in the phase diagram are present for all axial symmetries $\{C_n,C_{nv},S_{2n},C_{nh},D_n,D_{nh},D_{ad}\}$ with finite $n$, while the dashed line transition is present only for the symmetries $\{C_n,C_{nv},S_{2n},D_n,D_{nh}\}$ and the “biaxiality” parameter $\frac{J_3}{J_1}$. In Fig. 3 we vary the reduced axial and in-plane couplings $(\beta J_1, \beta J_3)$ independently since these relate more directly to the independent coupling strengths of the separate nematic orders in contrast to the relative anisotropy.

Let us start with the features of the phase diagram shown in Figs. 2 and 3 (upper panel). As we discussed, the strength of the biaxial order should reinforce the uniaxial order more than the uniaxial order reinforces the biaxial ordering, affecting the transition temperatures. Moreover, as has been discussed in Sec. III B, biaxial nematics with a more symmetric in-plane structure require larger $\beta J_1$ to stabilize the in-plane order. The critical anisotropy $(\frac{\beta}{J})_c$ for the uniaxial-biaxial transitions

![FIG. 3. The schematic $(\beta J_1, \beta J_3)$ phase diagrams of axial nematics. (a) The phase diagram in Fig. 2 in terms of $(\beta J_1, \beta J_3)$. As in Fig. 2, for low-order groups with two- and threefold symmetries the effective couplings stabilizing the biaxial and uniaxial order are of the same order and lead to a transition directly to the biaxial phase. (b) For higher in-plane symmetries, the biaxial phase is suppressed in comparison to the uniaxial phase. When allowed by the symmetries, axial terms with a vector or second rank uniaxial order parameter appear always in the Hamiltonian even at $J_3 = 0$ due to the O(3) constraints. These always stabilize the uniaxial order while the higher order biaxial order is still fluctuating. Solid lines in the phase diagram are present for all axial symmetries $\{C_n,C_{nv},S_{2n},C_{nh},D_n,D_{nh},D_{ad}\}$ with finite $n$, while the dashed biaxial$^*$-transition is present only for the symmetries $\{C_n,C_{nv},S_{2n},D_n,D_{nh}\}$ and the dotted uniaxial$^*$-transition for $\{C_n,C_{nv}\}$.](https://example.com/fig3)
TABLE III. A selection of 3D nematic order parameters. The first column specifies the symmetries, the second column the type \( A, B \) of the order parameter, and the third column gives the explicit form of the order parameter tensors [23]. Besides the tensors shown here, chiral nematics \( D_n \) have in addition a chiral order parameter \( \sigma \) defined by Eq. (2).

| Symmetry groups | Type | Ordering tensors | Tensor rank |
|-----------------|------|-----------------|-------------|
| \( D_2, D_{2h} \) | \( B[1,m] \) | \( 1 \otimes 1 - m \otimes m \) | 2 |
| \( D_3, D_{3h} \) | \( B[1,m] \) | \( (1^3 - 1 \otimes m^2 - m \otimes 1 \otimes m = m^2 \otimes 1) \) | 3 |
| \( D_4, D_{4h} \) | \( B[1,m] \) | \( 1^2 \otimes m^2 + m^2 \otimes 1^2 - \frac{4}{15} \delta_{ab} \delta_{cd} \otimes \mu = a, b, c, d \) \( e_\mu \) | 4 |
| \( D_5, D_{5h}, D_{5ch} \) | \( A[n] \) | \( n \otimes n \) | 2 |

will therefore move to the right for biaxial nematics having a larger in-plane \( n \)-fold rotational symmetry or more in-plane reflections. One the other hand, since a weaker in-plane order in turn means effectively stronger axial order, the critical anisotropy \( (\frac{J}{J})_c \) for the biaxial-biaxial\(^*\) transitions will correspondingly also move to the right. Therefore this phase region shrinks, while the uniaxial phase should become more prominent.

In the \((\beta J_1, \beta J_3)\)-phase diagram of Fig. 3, the corresponding points move to the opposite directions, similarly enlarging the uniaxial region and shrinking the biaxial\(^*\) phase. At the same time, as the symmetry increases, the biaxial order fluctuates more strongly leading to the transition to the biaxial phase at considerably lower temperatures. In addition to these general trends, for higher order symmetries, the presence of the induced axial couplings rounds the phase transitions to the uniaxial phase from the isotropic liquid and leads to a finite region where only the uniaxial phase is stabilized without a direct transition to the biaxial phases. In this region, at small enough \( \beta J_3 \), it is possible to stabilize only the more disordered uniaxial\(^*\) phase with higher \( n \rightarrow -n \) symmetry, if the original uniaxial order is vectorial. At larger \( \beta J_1 \), the uniaxial vector order is again lost in the biaxial\(^*\)-biaxial transition. As summarized in Table I, the uniaxial\(^*\) phase occurs only for the groups \( C_n, C_{3n} \). In the case of \( D_n \), the uniaxial\(^*\)-transition is not possible, but the biaxial\(^*\)-biaxial transition persists due to the nonzero chiral order parameter \( \sigma \) in the \( D_n \) biaxial phase, whereas the biaxial\(^*\) phase has the symmetry \( D_{nh} \).

Last, although the gauge theoretical formulation is not realized microscopically in any condensed matter system, it encodes the mesogenic symmetries very efficiently, and we expect the qualitative features and the topology of the phase diagrams to be applicable to many generalized nematic systems. This is clear from the biaxial-uniaxial phase diagrams (symmetries \( D_2 \) and \( D_{2h} \)) where all expected features of the mean-field phase diagram are recovered [13,15]. Moreover, in agreement with Ref. [15], we also see evidence of a tricritical point along the biaxial-uniaxial line, as will be discussed in more detail in Sec. IV.

IV. QUANTITATIVE PHASE DIAGRAMS OF THE GAUGE THEORETICAL DESCRIPTION

Having introduced the general concepts and framework, we still need to explicitly verify the generalized biaxial phase transitions given by Eqs. (8) and (11) departing from the gauge theoretical description. For this purpose we have simulated the temperature-anisotropy phase diagram and the \( J_1-J_3 \) phase diagrams for various symmetries, using the standard Metropolis Monte Carlo algorithm. These simulations were performed on lattices having dimensions \( L^3 = 8^3, 10^3, 12^3, 16^3 \). The associated order parameters and their characterizations relevant for the phase transitions are collected in Table III and Table I, respectively. As we detail below, the obtained results completely agree with the general scenario of generalized biaxial phase transitions as discussed in the previous sections.

A. Determination of the phases

To determine the symmetry of a nematic phase with tensor order parameter \( O^G \), one in principle needs to consider all the entries of \( O^G \). However, for interactions favoring homogeneous distribution of the order parameter fields, such as the interaction in the gauge model Eq. (12), the symmetry of the phase can be revealed by the strength of the order parameter defined as

\[
q = \sqrt{\left(\langle O^G_{abc} \rangle \right)^2},
\]

where \( O^G = \frac{1}{L^3} \sum_i O^G \), averages the order parameter tensor over the system, \( a, b, c, \ldots \) denote the tensor components, and contractions for repeated tensor indices are assumed. In combination with symmetry arguments, the scalar order parameter is enough to fix the symmetry of the phase, and the nematic ordering strength will develop a finite value in the ordered phase and vanish in the disordered phase (for more details, see, e.g., Refs. [14,23]).

For axial nematics, we accordingly need to define the ordering strength for the axial order \( A^G \) and the in-plane order \( B^G \), respectively,

\[
q_A = \sqrt{\left(\langle A^G_{ab} \rangle \right)^2},
\]

\[
q_B = \sqrt{\left(\langle B^G_{ab} \rangle \right)^2}.
\]

A transition is then identified by monitoring the appearance of a peak in the associated susceptibility

\[
\chi(q_{A,B}) = \frac{L^3}{T} \left( \langle q_{A,B}^2 \rangle - \langle q_{A,B} \rangle^2 \right),
\]

where \( \langle \ldots \rangle \) denotes the thermal average.
in the usual way,

$$C_v = \frac{1}{T^2 L^3} \left( \langle E^2 \rangle - \langle E \rangle^2 \right),$$  \hspace{1cm} (26)$$

$$\chi(\sigma) = \frac{L^3}{T} \left( \langle \sigma^2 \rangle - \langle \sigma \rangle^2 \right),$$  \hspace{1cm} (27)$$

where $E$ is the internal energy of the system, and $\sigma = \frac{1}{L^2} \sum_i \sigma_i$ is the global chiral order parameter.

**B. Phase diagrams involving temperature versus anisotropy**

A salient feature of our results is that we can retrieve the well-known temperature-anisotropy phase diagram of $D_{2h}$ and $D_2$ nematics within our gauge theoretical setting (12); see Fig. 4. In the region of small $\frac{J}{J_3}$, where the stiffness of the in-plane order is weaker than that of the axial order, we see that upon increasing the temperature, the in-plane order is destroyed first, leaving room for a vestigial $D_{\infty h}$-uniaxial phase. This vestigial uniaxial phase vanishes at a critical anisotropy $(\frac{J}{J_3})^\ast$, after which the in-plane and axial order become of comparable strength, and the transitions merge into a single transition between the biaxial phase and the $O(3)$ liquid phase. On the other hand, for large $\frac{J}{J_3}$, when the in-plane order is sufficiently strong, there will be a vestigial $D_{2h}$-biaxial phase in the $D_2$ case [Fig. 4(b)]. This realizes the biaxial-biaxial$^*$ transition in Eq. (11). We note, however, that, as discussed in Sec. III B, although the in-plane coupling can effectively induce an axial coupling, the axial order is not fully destroyed during this transition. The resulting behavior of the associated order parameters across these transitions is given in Table I.

Moreover, we find that the direct transition between the $D_{2h}$-biaxial nematic phase and the $O(3)$ isotropic liquid phase in Fig. 4(b) is first-order-like. Both $\chi(q_B)$, $\chi(\sigma)$ and $C_v$ exhibit a sudden peak at the transition, and the magnitude of their peak grows dramatically with the lattice size. This discontinuity continues to the biaxial-uniaxial transition line. Therefore, we identify a triple point where the three transition lines in Fig. 4(b) meet and the three phases can coexist. Moreover, in the middle of the biaxial-uniaxial transition line we find evidence for a tricritical point where the first order phase transition terminates and the transition becomes continuous.

These observations exactly agree with the mean field phase diagram and the experimental results of biaxial nematics in Refs. [10,15].

Besides these two familiar examples, we have also verified the generalized uniaxial-biaxial transitions in Eq. (8) as well as the uniaxial-uniaxial$^*$ and biaxial-biaxial$^*$ transition in Eq. (11) for nematics having symmetry \{$S_2$, $C_2$, $C_{2v}$, $C_{2h}$, $D_{2d}$, $S_4$, $D_3$, $D_{3h}$, $C_{4v}$, $D_4$, $D_{4h}$, $D_6$, $D_{6h}$\}. These comprise all seven types of axial groups and include symmetries with low and high symmetric in-plane structure. For nematics with a low symmetry, including the cases \{$S_2$, $C_2$, $C_{2v}$, $C_{2h}$, $D_{2d}$, $S_4$, $D_3$, $D_{3h}$\}, the phase diagrams have been checked to have the same topology as those of the $D_2$ or $D_{2h}$ case and are thus not presented here. For nematics with a high symmetry, comprising the cases \{$D_4$, $D_{4h}$, $D_6$, $D_{6h}$\}, the generalized biaxial transitions will however be affected dramatically by the induced axial coupling, as discussed in Sec. III C, and the phase diagrams are different. In the next section, we will discuss these phase diagrams for each of these symmetries.

**C. $J_1$-$J_3$ phase diagrams**

As already discussed in the introduction, within the gauge theoretical description we can in fact compare the physics of nematics with different symmetries in a common reference. In Fig. 5 we show the $J_1$-$J_3$ phase diagram for $D_{2h}$, $D_{3h}$, and $D_{4h}$ nematics. Let us first focus on the $D_{2h}$ case in Fig. 5(a). As in the temperature-anisotropy phase diagram in Fig. 4(a), in the region with small $J_1$ and large $J_3$ there is a vestigial uniaxial phase sandwiched between the fully ordered biaxial phase and the disordered liquid phase. The critical anisotropy where the vestigial uniaxial phase starts appearing is consistent with that of Fig. 4(a), up to our numerical accuracy. Moving to the $D_{3h}$ case, the increased in-plane symmetry requires a larger in-plane coupling (lower temperature and larger $\frac{J_1}{J_3}$ anisotropy) to stabilize the biaxial order, due to the more severe
FIG. 5. The $J_1$-$J_3$ phase diagram of (a) $D_2h$, (b) $D_3h$, and (c) $D_4h$ nematics. The red stars in (a) $D_2h$ and (b) $D_3h$ highlight a triple point, in which the three transition lines meet. Similar to the temperature-anisotropy phase diagram in Fig. 4, there is a vestigial uniaxial phase appearing from the region with small $J_1$ and large $J_3$ (small $\frac{J_1}{J_3}$), realizing the generalized biaxial-uniaxial transition in Eq. (8). As the symmetry increases, this vestigial uniaxial phase becomes more prominent and the fully ordered biaxial phase is remarkably squeezed. When the symmetry is sufficiently high, the vestigial uniaxial phase appears adjacent to the isotropic liquid due to the symmetry allowed axial terms. Moreover, our simulations indicate that depending on strength of the in-plane coupling, the biaxial-uniaxial transition may be either first order or second order. Therefore a tricritical point may exist in the biaxial-uniaxial transition line.

FIG. 6. The $J_1$-$J_3$ phase diagram of (a) $D_2$, (b) $D_3$, and (c) $D_4$ nematics. This is similar to Fig. 5, but there is in addition a vestigial biaxial phase at small $J_1$ region, realizing the generalized biaxial-biaxial$^*$ transition in Eq. (11). Both this vestigial biaxial phase and the fully ordered biaxial phase are squeezed considerably as the symmetry increases. The associated triple points at where transition lines meet are highlighted by large stars. As in Fig. 5, there may be a tricritical point in the biaxial-uniaxial transition line.

fluctuations. The biaxial phase is therefore squeezed by the liquid phase and the vestigial uniaxial phase.

The squeezing of the biaxial phase is even more prominent for the $D_{4h}$ nematics, where the in-plane symmetry is increased further. In particular, since very large in-plane coupling is
required to stabilize the highly symmetric $D_{2h}$ order, before the biaxial phase is realized, the induced axial coupling is always sufficiently strong for the uniaxial order. This leads to a vestigial uniaxial phase realized for all non-negative values of the “bare” axial coupling $J_3$, while the direct biaxial-liquid transition is absent. The same is true for the more symmetric $D_{4h}$ nematics, with a even larger region of the vestigial uniaxial phase.

However, one should not interpret this as a no-go theorem for a direct biaxial-liquid transition in the case of highly symmetric biaxial nematics. Instead, this simply means that in order to realize this transition, one needs to consider a model with “antinematic” coupling for the axial order to offset the induced axial coupling.

The above discussions can similarly be verified for $D_2$, $D_3$, and $D_4$ nematics as well, as shown in Fig. 6. Nonetheless, since the biaxial-biaxial* transition is possible for these cases, in the small $J_3$ region, there is in addition a vestigial biaxial* phase. This phase is also squeezed as symmetries increase, as in the case of the fully ordered biaxial phase. Moreover, in the cases of $D_2$ and $D_3$, there are direct transitions from the fully ordered biaxial phase or vestigial biaxial phase to the liquid phase. For the highly symmetric $D_4$ case, however, these transitions are replaced by a biaxial-uniaxial or a biaxial*-uniaxial transition, since a vestigial uniaxial phase exists for all non-negative values of $J_3$ as in the $D_{2h}$ case due to the induced axial couplings.

V. CONCLUSIONS AND OUTLOOK

There is a rich landscape of unexplored generalized nematics, entailing not only a diversity of orientational phases in terms of their symmetry but also an abundance in possible vestigial phases. In this paper, we have discussed the anisotropy-induced vestigial uniaxial and biaxial phases for nematics characterized by axial point-group symmetries and studied their phase transitions. Our results generalize the well-studied biaxial-uniaxial transition of $D_{2h}$ nematics to a much broader class, which can be directly accessed within our earlier proposed gauge theoretical formulation of generalized nematics [14] and follow from a priori symmetry arguments. This framework allows us in particular to compare nematics and vestigial phases with different symmetries in one common reference. Utilizing this formalism, we found that, in comparison to the familiar $D_{2h}$ biaxial nematic phase, nematic phases with high axial symmetries require much lower temperature to stabilize their order. This motivates the fact that biaxial phases with high symmetry are difficult to realize in reality and have not yet been experimentally encountered: before reaching the low temperature demanded by the biaxial order, crystallization may already start playing a role. Consequently, columnar, smectic and/or crystalline phases may occur instead of a generalized nematic phase. We stress that such states are not captured by our model that by construction encompasses only the orientational ordering. These challenges notwithstanding, the advances in the fabrication and manipulation of colloidal systems of nanoparticles appear in fact promising with regard to stabilizing generalized nematic phases in the laboratory in the near future [42–46].

Besides these generalized biaxial transitions, there may be more vestigial phases and transitions in the gauge model [Eq. (12)]. Those phases are associated with the defects in the model, which have been ignored in this work by setting $H_{\text{gauge}} = 0$ in Eq. (12), describing the confined and Higgs phases of the model. From the point of view of topological melting, phase transitions may be understood as a proliferation of topological defects [31,47,48]. To illustrate this further we can take the $D_{2h}$-biaxial nematic as an example. According to homotopy theory, topological defects of $D_{2h}$ nematics are classified by the five conjugacy classes of the quaternion group $Q_8$ [49–52]. Among these defects, there are only three elementary ones, which are the $\pi$ disclinations in the three orthogonal planes of the 3D space. In the transition of the nematic phase to the O(3) liquid phase, all these defects proliferate. In the biaxial-uniaxial transition, however, one of them stays gapped. This implies that a phase transition can be affected by the tuning of the energy cost of topological defects. The gauge model [Eq. (12)] provides a natural way to do this. Concretely, when the $H_{\text{gauge}}$ term is set to be zero, topological defects in the model only cost elastic energy by the $H_{\text{Higgs}}$ term. By tuning on the $H_{\text{gauge}}$ term, however, we can introduce a finite core energy to a particular class of topological defects and therefore modify the nature of the phase transition. While such defect terms $H_{\text{gauge}} \neq 0$ have been identified to be important in the melting of many quantum nematics [53–58], they have not yet been discovered to play a role in the realm of classical nematics and melting [59]. The rich physics associated with these ideas leaves many interesting avenues of for future research in the generalized nematic systems.

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