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Multimodal Approach Reveals the Symmetry-Breaking Pathway to the Broken Helix in EuIn₂As₂

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Understanding and manipulating emergent phases, which are themes at the forefront of quantummaterials research, rely on identifying their underlying symmetries. This general principle has been particularly prominent in materials with coupled electronic and magnetic degrees of freedom, in which magnetic order influences the electronic band structure and can lead to exotic topological effects. However, identifying symmetry of a magnetically ordered phase can pose a challenge, particularly in the presence of small domains. Here we introduce a multimodal approach for determining magnetic structures, which combines symmetry-sensitive optical probes, scattering, and group-theoretical analysis. We apply it to $EuIn_2As_2$, a material that has received attention as a candidate axion insulator. While first-principles calculations predict this state on the assumption of a simple collinear antiferromagnetic structure, subsequent neutron-scattering measurements reveal a much more intricate magnetic ground state characterized by two coexisting magnetic wave vectors reached by successive thermal phase transitions. The proposed high- and low-temperature phases are a spin helix and a state with interpenetrating helical and Néel antiferromagnetic order termed a "broken helix," respectively. Employing a multimodal approach, we identify the magnetic structure associated with these two phases of $EuIn_2As_2$. We find that the highertemperature phase is characterized by a variation of the magnetic moment amplitude from layer to layer, with the moment vanishing entirely in every third Eu layer. The lower-temperature structure is similar to the broken helix, with one important difference: Because of local strain, the relative orientation of the magnetic structure and the lattice is not fixed. Consequently, the symmetry required to protect the axion phase is not generically protected in $EuIn_2As_2$, but we show that it can be restored if the magnetic structure is tuned with uniaxial strain. Finally, we present a spin Hamiltonian that identifies the spin interactions that account for the complex magnetic order in EuIn₂As₂. Our work highlights the importance of a multimodal approach in determining the symmetry of complex order parameters.

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I. INTRODUCTION

The search for materials exhibiting novel emergent properties relies on the identification of their characteristic symmetries. Examples of such properties include the anomalous and topological Hall effects arising from Berry curvature in momentum [1] and real space [2], respectively, momentum-dependent Zeeman splitting of electronic bands [3], and quantized response functions in topological systems [4,5]. These phenomena all depend on the underlying magnetic order and present the exciting prospect of tuning by manipulating the symmetry of the magnetic state.

Promising material candidates for observing these phenomena can be identified by *ab initio* calculations [6,7]. Of necessity, such calculations are based on an assumed magnetic structure, as it is difficult to reliably predict the magnetic ground state. An effective search strategy is to combine *ab initio* calculations with the magnetic order

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deduced from scattering measurements. However, in some of the most interesting material systems the interpretation of scattering data can lead to ambiguities of critical importance. For example, states that exhibit order characterized by multiple symmetry-related wave vectors (Q) pose a challenge, since the diffraction pattern does not readily distinguish phase-sensitive mixed order from an equal population of domains characterized by a single **O**. Similarly, domains complicate the determination of the orientation of magnetic moments even in a single-Q structure. The distinct scenarios consistent with a given diffraction pattern are radically different from a symmetry perspective, and have distinct consequences for response functions and topological properties, motivating specialized scattering experiments [8–10] and comparisons with complementary experimental techniques [11,12].

Here we introduce a multimodal approach for determining magnetic symmetries, which is based on combining experimental information from scattering and symmetrysensitive optics with group-theoretical analysis. The power of our methodology comes from the fact that the scattering and symmetry-sensitive optics have direct access to complementary information. Scattering experiments directly measure ordering wave vectors and can obtain further information about magnetic structure through comparison of the diffracted intensities with predictions from a model. In contrast, optical experiments directly probe point group symmetries, such as rotation and inversion, as well as timereversal symmetry. The role of group theory is to identify all magnetic structures that can arise from the parent paramagnetic group and are consistent with both sets of experimental observations.

In this manuscript, we demonstrate the power of our multimodal approach on EuIn₂As₂, whose structure is shown in Figs. 1(a) and 1(b). We choose this compound because it is a perfect example of the scenario outlined above. Ab initio calculations based on an assumed antiferromagnetic structure [(AFM) Fig. 1(c)] predicted that it hosts the elusive axion insulator state [13] exhibiting quantized responses to electromagnetic fields [14,15]. However, a more complex magnetic behavior was uncovered by neutron-scattering measurements, with two close-by transitions at $T_{N1} \approx 17.5$ K and $T_{N2} \approx 16$ K [16], in which two distinct propagation vectors appeared sequentially. Although neither of the two phases appearing at T_{N1} and T_{N2} are consistent with the previously assumed AFM structure, the magnetic structures assigned to the two phases [Figs. 1(d) and 1(e)] were shown to host the axion state [16]. However, this conclusion was based on symmetries deduced from neutron scattering alone. The complications in interpreting neutron-scattering data in a material in which domains are naturally expected to form motivated us to explore the additional information that can be accessed through the multimodal approach.



FIG. 1. (a) The side view and (b) the top view of the crystal structure of $EuIn_2As_2$. (c)–(e) A table showing the three magnetic structures discussed for $EuIn_2As_2$ in previous work: the theoretically assumed *A*-type antiferromagnet (AFM), the helix proposed in phase I, and the broken helix proposed in phase II. In all structures, the moments lie in Eu planes, and individual layers are ferromagnetically aligned. For each structure, we show the layer-resolved spin orientation, with the hexagon denoting the lattice orientation, as well as a top view (last row), with spins labeled by the layer number. (f) The magnetic unit cell corresponding to three crystallographic unit cells. We mark spin orientation in odd and even layers with white and colored arrowheads, respectively.

Our multimodal approach reveals a different picture of how the magnetic order of EuIn₂As₂ evolves with the temperature, and we identify two magnetic states that would have remained hidden to the application of any single technique. A higher-temperature phase I (T_{N1} > $T > T_{N2}$) is characterized by the onset of a single wave vector \mathbf{Q}_1 , which we identify as a "nodal amplitudemodulated state," in which the expectation value of the magnetic moment vanishes on every third Eu layer. In the lower-temperature phase II $(T < T_{N2})$, an additional wave vector \mathbf{Q}_2 appears, forming an "unpinned broken helix," in which the orientation of the magnetic moments with respect to the crystal axes varies continuously with the location on the sample. Only special orientations, with moments aligned to high-symmetry directions of the lattice, maintain symmetries that protect the topological phase.

Although the experimentally informed group-theory analysis mentioned above is well suited to identifying the magnetic phases that emerge in EuIn_2As_2 , it does not reveal their microscopic origin. To address this point, we propose a minimal spin model that captures the unpinned broken helix as the ground state. We show that this state requires the exchange interactions $J(\mathbf{Q})$ to peak sharply at two values of the wave vector \mathbf{Q} . This requires long-ranged interactions naturally arising from the coupling of itinerant electrons to the magnetic degrees of freedom. We further show that such exchange interactions alone cannot account for the higher-temperature amplitude-modulated state. Whether it can be stabilized by thermal fluctuations close to the two transitions is an interesting question posed by our work.

This paper is organized as follows. We summarize the current understanding of EuIn₂As₂ in Sec. II. In Sec. III, we introduce two optical techniques sensitive to rotational and time-reversal symmetries, and show the results of these experiments as a function of the temperature, position on the sample, and cooling protocol. We demonstrate that the symmetries revealed by these measurements are inconsistent with the previous understanding of magnetic phases in EuIn₂As₂, and therefore motivate a new analysis of the order parameters. In Sec. IV, we perform a systematic analysis of symmetry-allowed magnetic structures by combining scattering and optical data with group theory. We identify phases I and II as a nodal amplitude-modulated state and an unpinned broken helix, respectively. We complement this analysis with a phenomenological Landau free-energy model (Sec. V), demonstrating the symmetry-breaking pathway toward the ground state. Furthermore, we show experimentally and theoretically how the symmetry of the magnetic ground state and electronic topology can be tuned by uniaxial strain (Sec. VI). Finally, in Sec. VII we introduce a microscopic spin Hamiltonian pointing to the importance of the coupling between electronic and magnetic degrees of freedom. We conclude by emphasizing that our measurements and analyses show that EuIn₂As₂ hosts a remarkably rich and tunable system of coupled electrons and localized moments, opening doors for future efforts to control topological phases.

II. CURRENT UNDERSTANDING OF EuIn₂As₂

EuIn₂As₂ is a rare-earth-based magnetic material with triangular layers of Eu stacked along the crystallographic *c* direction and separated by blocks of In₂As₂. Hereafter, we set *c* parallel to \hat{z} [Figs. 1(a) and 1(b)]. Magnetism arises from the localized moments of the Eu²⁺ ions (S = 7/2, L = 0). The moments in each Eu layer are ferromagnetically aligned with respect to each other and localized in the Eu-Eu planes. This easy-plane anisotropy confirmed by magnetization and neutron-scattering experiments [16] is implicitly assumed throughout this paper; when we refer to "moment orientation," we are referring to the orientation within the Eu planes.

The low-energy electronic bands dominated by the 5s orbitals of In and 4p orbitals of As were predicted to host topological properties. Combined with the Eu-based magnetism, this makes $EuIn_2As_2$ a promising platform to explore the interplay between topology and

magnetism, as has been proposed and debated in the context of several Eu-based magnetic materials [17–24]. More specifically, EuIn₂As₂ was predicted to host the axion insulator state [13,16], a phase characterized by half-quantized magnetoelectric coupling in the bulk and half-quantized Hall surface conductivity. The axion phase requires T to be broken in order to gap the surface states, while the quantization is protected by another symmetry that reverses the sign of the magnetoelectric coupling constant, such as spatial inversion (\mathcal{P}), \mathcal{T} combined with half-translations, or the product of \mathcal{T} and twofold rotations ($\mathcal{T}C_2$).

The original prediction of an axion insulator state in EuIn₂As₂ was based on first-principles calculations that showed an insulating bulk and an A-type antiferromagnetic order illustrated in Fig. 1(c). Note that this Néel-like magnetic order does not change the periodicity of the lattice, since the paramagnetic unit cell already contains two Eu layers; as a result, the magnetic ordering vector coincides with the Bragg wave vector (0, 0, 1). Importantly, this magnetic configuration preserves spatial inversion, which protects the axion phase [13]. Experiments, however, uncovered a different, but intriguing, picture: EuIn₂As₂ shows metallic dc [25,26] and optical [27] transport, it has a Fermi surface [25,28], and the magnetic structure is considerably more complex. Both neutron-scattering [16] and resonant x-ray-scattering [29] experiments found two consecutive magnetic transitions ($T_{N1} \approx 17.5$ K, $T_{N2} \approx 16$ K) corresponding to the onsets of two distinct wave vectors. At T_{N1} , a single propagation vector $Q_1 \approx (0, 0, 1/3)$ is observed, while at T_{N2} an additional propagation vector $Q_2 = (0, 0, 1)$ emerges. It is important to note that while the neutronscattering [16] and x-ray-scattering [29] data are broadly consistent, they observed a slightly different Q_1 : In Ref. [16], Q_1 varies between (0, 0, 0.33) at T_{N1} and (0, 0, 0.303) at T_{N2} , and remains temperature independent at $T < T_{N2}$. In contrast, in Ref. [29], $Q_1 = (0, 0, 1/3)$ is commensurate and temperature independent. This difference cannot be accounted for by any experimental considerations and indicates a slight sample-to-sample variation. To remove any ambiguities stemming from such variations, we performed all optical measurements on the same crystal used for x-ray scattering.

The higher- and lower-temperature phases that we refer to as phase I and phase II, respectively, are interpreted as a 60° helix [Fig. 1(d)] and a novel magnetic state exhibiting interpenetrating type-A AFM and spin-helical orders, respectively. The latter phase is termed "broken helix" [Fig. 1(e)]. Despite this complexity, both phases still host an axion insulator state that, in this case, is protected by a magnetic symmetry TC_2 that is the product of timereversal and twofold rotation [16]. In the following, we demonstrate that the findings of our optical experiments challenge this established picture.

III. OPTICAL PROBES

We measure optical reflectance to extract information on the threefold rotational symmetry around \hat{z} (C_{3z}) and timereversal symmetry (T); the latter we probe in two complementary ways. In this section, we briefly describe the physical principles behind these measurements and the relevant experimental considerations. We then show the findings of these measurements performed on EuIn₂As₂ as a function of the temperature, position on the sample, and cooling protocol. We discuss how the unique capability of these probes reveals information incompatible with the current understanding of EuIn₂As₂.

A. Symmetry-sensitive experiment

The reflectance at normal incidence is described by a 2×2 matrix, which can be parametrized as

$$\mathbf{r} = r_0 \mathbb{1} + \delta r_s \begin{pmatrix} \cos 2\theta_0 & \sin 2\theta_0 \\ \sin 2\theta_0 & -\cos 2\theta_0 \end{pmatrix} + \delta r_a \begin{pmatrix} 0 & 1 \\ -1 & 0 \end{pmatrix}, \quad (1)$$

where 1 denotes the unit matrix, and θ_0 and $\theta_0 + \pi/2$ correspond to the principal optical axes. r_0 is the isotropic contribution allowed in all materials, whereas δr_s and δr_a known as birefringence and the Kerr effect, respectively, contain distinct symmetry information. The crucial difference between them is their behavior with respect to the exchange of indices: δr_s is symmetric and δr_a antisymmetric. The general reciprocity relations [30,31] enforce $\delta r_a = 0$ in T-invariant systems; thus, $\delta r_a \neq 0$ is an unambiguous probe of time-reversal symmetry breaking. In a reflectivity experiment the Kerr effect manifests as a change of polarization orientation upon reflection by an angle of $\delta r_a/r_0$ rad.

In contrast, δr_s in not sensitive to \mathcal{T} , but is forbidden by rotational symmetry C_{nz} if $n \ge 3$. A nonzero δr_s means that the reflectance depends on the relative orientation of the light polarization and the lattice. Like δr_a , δr_s also induces a polarization rotation upon reflection; however, the magnitude of the change depends on the incident polarization. In particular, the rotation vanishes when the incident polarization is aligned to the principal optical axes. As we show below, this fact is the basis for our detection of rotational symmetry breaking.

Further symmetry-sensitive information can be gleaned from the change of reflectance in an applied magnetic field H_z . To linear order the symmetric and antisymmetric components of the field-induced change to reflectance r_H can be parametrized as

$$\mathbf{r}_{H} = \delta \alpha_{s} H_{z} \begin{pmatrix} \cos 2\theta_{H} & \sin 2\theta_{H} \\ \sin 2\theta_{H} & -\cos 2\theta_{H} \end{pmatrix} + \delta \alpha_{a} H_{z} \begin{pmatrix} 0 & 1 \\ -1 & 0 \end{pmatrix}.$$
(2)

With the additional \mathcal{T} -odd factor H_z , the reciprocity relations now allow the antisymmetric term ($\delta \alpha_a$) in all

materials, while the symmetric term ($\delta \alpha_s$) indicates breaking of time-reversal symmetry. A nonzero ($\delta \alpha_s$) is usually referred to as the linear magneto-optic effect [32] or linear magnetobirefringence (LMB). The principal axes of the LMB response are given by θ_H and $\theta_H + \pi/2$. Although both δr_a and $\delta \alpha_s$ require time reversal to be broken, the symmetry conditions that enforce them to be nonzero are different. This follows from the first term of Eq. (2), which shows that C_{3z} must be broken in addition to T for $\delta \alpha_s$ to be nonzero.

The three distinct symmetry-sensitive quantities that we obtain through reflectance measurements are therefore δr_s , δr_a , and $\delta \alpha_s$. All of them induce changes of light polarization upon reflection and can be distinguished from each other by our optical techniques [33–35]. The fundamental observable is the rotation of the angle of linear polarization $(d\phi)$ about the optical axis at normal incidence as a function of sample orientation. Since we cannot physically rotate the sample, we access the same information by rotating the incoming light polarization (ϕ). The change of polarization upon reflection at temperature *T* and field H_z is given by

$$d\phi = A(T, H_z) \sin\left[2(\phi - \theta(T, H_z))\right] + B(T, H_z). \quad (3)$$

The amplitude of the sinusoidal variation $A(T, H_z)$ and the principal axis orientation $\theta(T, H_z)$ are determined by the symmetric part of the reflectance tensors (δr_s , $\delta \alpha_s$), while the constant offset B(T, H) originates from the antisymmetric δr_a and $\delta \alpha_a$. It is now clear how the quantities of interest can be experimentally distinguished: In an $H_z = 0$ measurement the sinusoidally varying and constant $d\phi$ originate from δr_s and δr_a , respectively, while $\delta \alpha_s$ is captured by a sinusoidal variation proportional to a magnetic field.

In practice, we perform measurements in two modes (Fig. 2; see Appendix B for details). In the temperaturemodulated mode (*T*-mod), we modulate the temperature at a frequency $f \approx 2$ kHz using a second laser beam as a heater (780 nm, 125 μ W), therefore measuring the temperature derivative of Eq. (3). This experimental procedure reveals the same information about symmetry as the unmodulated experiment at $H_z = 0$ (δr_s , δr_a), while enhancing sensitivity and rejecting contributions from inevitable setup imperfections.

In the field-modulated mode (*H*-mod), we modulate the magnetic field supplied by a copper coil ($H_z \approx 3$ mT, $f \approx 100$ Hz), and measure the field derivative of Eq. (3). This experiment is sensitive to $\delta \alpha_s$. Therefore, the symmetry constraints to observe nonzero temperature and field derivatives of A ($\partial_T A$ and $\partial_H A$, respectively) differ. As we will demonstrate below, the ability to simultaneously measure these quantities is crucial to determine the symmetry of the two ordered phases in EuIn₂As₂.



FIG. 2. Schematic of the optical setup used for measurements of the polarization rotation as a function of the incident polarization. For thermal modulation (*T*-mod), an optically chopped second beam (pump, 780 nm, 125 μ W, chopping frequency 2 kHz) is spatially overlapped with the probe beam on the sample surface and used to modulate the temperature of the sample by heating it. During field modulation (*H*-mod), an ac magnetic field ($H_z \approx 3$ mT, $f \approx 100$ Hz) is applied along the *c* axis of the sample with a coil. During thermal (field) modulation, the field (pump beam) is turned off. A helium neon laser is used as a probe beam for all the experiments (633 nm, 50 μ W).

B. Onset of the two phases

Figure 3 illustrates the onset with the decreasing temperature of the optical signatures of broken symmetry. The polar plots in Fig. 3(a) show the *T*-mod and *H*-mod rotation of polarization as red and blue symbols, respectively, measured at three temperatures. The orientations of the principal axes obtained by the two modulation modes differ by $\theta_H - \theta_0 = 45^\circ$, a point we return to later. The temperature dependence of the *T*-mod and *H*-mod amplitude and principal axis orientation are plotted in Figs. 3(b) and 3(c), respectively. Both amplitudes exhibit a sharp onset separated by 1.4 K. The onset of the *T*-mod birefringence coincides with the appearance of the Q_1 peak (phase I) and the *H*-mod signal to the appearance of antiferromagnetic Q_2 peak (phase II), as detected by x-ray-scattering measurements performed on the same crystal [29].

The two optical experiments, which probe two distinct symmetries, revealed the two transitions, and indicated that the symmetry-breaking pathway proposed on the basis of scattering experiments cannot be correct. The commensurate 60° -helix structure previously assumed to describe phase I preserves the C_{3z} symmetry of the lattice, and is thus incompatible with the birefringence onset at T_{N1} . In fact, birefringence was expected to arise only at T_{N2} , in contrast to the observation. It is important to consider whether the observed C_{3z} breaking could arise from something other than intrinsic magnetic order characterized by $Q_1 = (0, 0, 1/3)$. One concern might be the surface



FIG. 3. (a) Polar plots of thermally modulated (*T*-mod, red circles) and field-modulated (*H*-mod, blue diamonds) polarization rotation $\delta \varphi$ at different temperatures reveal distinct signatures of broken rotational symmetry in each of the two magnetic phases. Since rotation by 180° leaves light polarization invariant, $\delta \varphi$ measured as a function of the incident polarization angle from $-90^{\circ} \rightarrow 90^{\circ}$ is replicated for $90^{\circ} \rightarrow -90^{\circ}$. (b) Temperature dependence of *T*-mod and *H*-mod amplitude. The *T*-mod signal onsets at the transition temperature associated with the higher-temperature phase, whereas the *H*-mod signal onsets at the lower transition temperature. The amplitudes are normalized at 10 K. (c) Temperature dependence of the principal axes orientations corresponding to the *T*-mod and *H*-mod signals. The principal axes associated with the two signals are oriented 45° relative to each other and remain constant with the temperature.

sensitivity of optics, with typical penetration depth of approximately 50–100 nm: Could it be that our experiment is sensitive to the orientation of the spin at the top surface? This is not plausible: A uniform surface layer magnetization could be realized only if the naturally grown sample surface was atomically flat across micron length scales probed by our beam. Because of the short period of the helix, any realistic surface morphology would result in the surface layer exhibiting equal population of the three spin orientations of the helix, and the signals arising from those three would cancel. We emphasize that if $Q_1 \neq (0, 0, 1/3)$, C_{3z} symmetry is formally broken even by the helix state.

same crystal was therefore crucial to conclude that the optical measurements are incompatible with the helix state. Furthermore, spatially resolved measurements discussed in the following section reveal additional information, which is incompatible with the past understanding of both phases in $EuIn_2As_2$.

C. Spatial distribution of optical signals

To investigate the spatial dependence of the two optical signals, we raster scanned the sample under the beam focus and repeated the polarization rotation measurements across a $240 \times 260 \ \mu\text{m}^2$ sample region, with measurements taken every 20 μ m. These measurements revealed two surprising facts: (a) The principal optical axes in EuIn₂As₂ can assume any orientation with respect to the lattice, and (b) despite that breadth of orientations, the fundamental *H*-mod signal is the same in every position on the sample.

1. T-mod signal

First we focus on the H = 0 signal. In Figs. 4(a) and 4(b), we show a map of the spatial distribution of the principal axis orientation θ_0 and the corresponding histogram. We find a single broad peak in the distribution of the principal axis orientations spanning more than 30° [Fig. 4(b)], as emphasized by the polar plot showing the polarization dependence of the *T*-mod signal taken at two sample positions [Fig. 4(c)]. Since the two high-symmetry directions in the Eu planes are separated by 30°, these measurements show that the principal optical axes can take any orientation with respect to the lattice.

The observation of a broad distribution of principal axes is surprising, since breaking of the discrete C_{3z} symmetry is expected to yield three domains related by C_{3_7} , which would manifest as three narrow peaks in the histogram separated by 60° [33,35–37]. This observation could be interpreted in two ways, which are important to distinguish. Either the mapped region is dominated by a single domain, and the broadening of the histogram is caused by contributions of the other two domains, which cannot be resolved with our diffraction-limited resolution of 1 µm (microdomain scenario), or the magnetic order microscopically assumes a continuum of orientations (orientation continuum scenario). We note that the broken-helix structure proposed for phase II [Fig. 1(e)] protects the axion state only for one orientation of the structure with respect to the lattice; the orientation continuum scenario would therefore drastically change the topological properties. Below, we show that only the orientation continuum scenario is consistent with the data, requiring a reevaluation of symmetry and topology in EuIn₂As₂.

The orientation continuum scenario is proven by analyzing the amplitude of birefringence as a function of the position. In the microdomain scenario, the amplitude is the largest when the signal is dominated by a single domain and is reduced by averaging over microdomains, yielding a



FIG. 4. Principal axis orientations associated with T-mod and H-mod signals. (a),(d),(g) Maps of the spatial distributions of the principal axis orientations for T-mod (θ_0), H-mod (θ_H), and their difference $(\theta_H - \theta_0)$, respectively. (b),(e),(h) Histograms of the distributions in (a), (d), and (g), respectively. The orientations of the T-mod and H-mod signals are broadly distributed and continuously varying, whereas the difference between the orientations $\theta_H - \theta_0$ is sharply peaked at 45°. (c),(f) Polar plots of the polarization rotation at points labeled P1 and P2 in (a) and (d), respectively, overlaid on hexagons to demonstrate the lack of registration to the crystalline axes. The principal axes of the two points are oriented approximately 30° from each other in each of the T-mod and H-mod maps. (i) Polar plot of the T-mod and H-mod polarization rotations at point P1, demonstrating the principal axes of the T-mod and H-mod signals to be oriented 45° relative to each other.

well-defined prediction for the relationship between the birefringence amplitude and orientation (see Appendix D for more details). In contrast, the amplitude is independent of the angle in the orientation continuum scenario. As evident in Fig. 5(a), the amplitude is uniform across the sample region in which the principal axis orientation spans an angle larger than 30° , i.e., the full range between the high-symmetry directions of the lattice. In contrast, in the microdomain scenario the amplitude would systematically vary across this region [simulated in Fig. 5(b)].

The confirmation of the orientation continuum scenario suggests that the local principal axes are chosen by a builtin strain, and that the distributions in Figs. 4(a) and 4(b) reflect the distribution of built-in strain axes. A somewhat similar scenario was proposed to explain the rotation of the nematic director in the superconducting nematic phase of



FIG. 5. Measured and simulated birefringence amplitude in a T-mod experiment: (a) Thermally modulated birefringence amplitude in the same region as the maps in Fig. 4. The amplitude is nearly constant over the region. (b) Birefringence amplitude simulated using the microdomain model prediction for birefringence amplitude from orientation. The microdomain model predicted amplitude is expected to vary as the measured birefringence angle is continuously tuned, in contrast with the observed experimental amplitude.

twisted bilayer graphene [38,39]. However, it is unusual to observe an orientation continuum in a magnetic system, since magnetocrystalline anisotropy (MCA) tends to pin moments to the high-symmetry direction of the lattice. Our observation therefore suggests a very weak MCA, such that an inevitable built-in strain dominates, and selects an arbitrary orientation at a generic sample position. This hypothesis is proven below by deliberate application of uniaxial strain (Sec. VI), showing that the magnetic space group can be manipulated with important consequences for topology.

2. H-mod signal

The *H*-mod signal measured over the same sample region reveals an angle distribution θ_H of principal optical axes of similar width [Figs. 4(d)–4(f)]. However, it is quite striking that the distribution of the difference of the two angles ($\theta_H - \theta_0$) is remarkably narrow and centered at 45° [Figs. 4(g)–4(i)].

This finding raises two questions, which we address in the remainder of this section: why the distribution is so narrow and why time-reversed domains of $\theta_H - \theta_0 = -45^\circ$ are not observed anywhere on the sample.

The narrow distribution of the difference $\theta_H - \theta_0 = 45^\circ$ suggests that this relationship is enforced by symmetry, as has been observed in some other magnetic systems. For example, in Ref. [33] it was shown that the magnetic point groups 2/m and 2'/m' allow only for $\theta_H - \theta_0 = \pm 45^\circ$ and $\theta_H - \theta_0 = 0^\circ, 90^\circ$, respectively. However, further consideration reveals that this cannot be the case in EuIn₂As₂: Since the principal axis orientation generically does not coincide with the high-symmetry directions of the crystal, no in-plane rotational axes or vertical mirror planes remain valid symmetries at a generic sample position, alone or in combination with time reversal. Therefore, there is no symmetry that can enforce a relationship between θ_H and



FIG. 6. (a) *H*-mod signal as a function of the incident polarization angle measured at 5 K in a sample cooled in a training field of -3 and 3 mT (blue and green symbols, respectively). (b) *H*-mod signal measured using incident polarization of 50°, as a function of a training dc field applied during cooling. Both panels clearly demonstrate that the signal can be trained by a magnetic field, proving its time-reversal-odd nature.

 θ_0 , and a microscopic mechanism must instead underlay the sharp angle distribution in Fig. 4(h). Identifying why $\theta_H - \theta_0 = 0^\circ, 90^\circ$ is not observed, although it is allowed by symmetry, is beyond the scope of our work, and we hope that our findings motivate further *ab initio* investigation to identify the origin of the observation reported in Figs. 4(g)-4(i).

The remaining puzzle is the absence of the TR domain $(\theta_H - \theta_0 = -45^\circ)$. A closely related observation is that the same $\theta_H - \theta_0 = 45^\circ$ domain was observed on each cooldown through the transition, which is not expected of a \mathcal{T} -breaking order parameter that condensed in the absence of a magnetic field. To explore this, we measured *H*-mod birefringence at 5 K following cooling in a dc magnetic field [Fig. 6(a)]. The sign of the signal can be trained by the field, confirming its \mathcal{T} -odd nature. The two time-reversed states are characterized by values of θ_H that differ by 90° and correspond to $\theta_H = \theta_0 \pm 45^\circ$.

We further monitor the maximum signal at 50° of incident polarization, as a function of the training field [Fig. 6(b)], and find that even at a vanishing applied

training field the signal reaches a value close to saturation. This indicates that cooling the sample through the magnetic transitions in Earth's field is sufficient to train the domains, consistent with the observation that the sign of the signal does not change with sample position, or between different cooldowns. We conclude that the \mathcal{T} -breaking domains in EuIn₂As₂ are easily switched, and their sign can be detected through LMB.

IV. DETERMINING THE MAGNETIC STRUCTURES

In this section, we use a multimodal approach to determine the magnetic structures in EuIn₂As₂. Both optical and scattering experiments show evidence of two magnetic transitions as a function of the temperature, yielding two magnetic phases. These experiments impose complementary constraints on the corresponding magnetic structures. Our symmetry-sensitive optical measurements show that (i) C_{3z} is broken in phases I and II; (ii) \mathcal{PT} is broken in phase II, but preserved in phase I; (iii) the in-plane magnetocrystalline anisotropy is negligible. Additionally, scattering experiments [16,29] revealed the propagation vectors: (iv) $\mathbf{Q}_1 = (0, 0, 1/3)$ in phase I and (v) $\mathbf{Q}_1 = (0, 0, 1/3)$ and $\mathbf{Q}_2 = (0, 0, 1)$ in phase II. While observation (i) is inconsistent with the 60° -helix previously associated with phase I [16,29], we show in this section that it is possible to reconcile all the experimental results for both phases.

In phase, I we find a collinear state with varying amplitude of the magnetic moments that we hereafter refer to as nodal amplitude-modulated collinear order, while in phase II we find a state similar to the previously proposed broken helix [16], but with one key difference: The moments in the experimentally observed structure are not pinned to the crystalline axes. We refer to this state as the unpinned broken helix, and show that it can be manipulated with uniaxial strain (Sec. VI), raising the possibility of on-demand control of electronic topology in $EuIn_2As_2$. This tunability is a direct consequence of weak magnetocrystalline anisotropy, which can be overpowered even by modest built-in strain.

In the remainder of this section, we highlight the main steps involved in our experimentally guided symmetry analysis, while more details are given in Appendixes E and F. In Sec. V, we complement the symmetry analysis with a phenomenological free-energy model, which captures the sequence of broken symmetries and the corresponding evolution of magnetic structure. Finally, we summarize the main results of the symmetry and free-energy analyses; a reader more interested in those results than in the reasoning that led to them may immediately proceed to Sec. V B. Illustrations of the magnetic structures associated with the two phases are shown in Figs. 7(e), 8(c), and 8(e), respectively.



FIG. 7. Top row: the layer-resolved moment orientation in [(a),(b)] the two helical structures of opposite helicities $(H_1 \text{ and } H_2)$; [(d),(e)] the two symmetry-distinct amplitude-modulated structures $(A_1 \text{ and } A_2, \text{ respectively})$. Middle row: two equivalent choices for the basis function for the $m\Delta_6$ irrep—(c) helical and (f) amplitude-modulated states. Bottom row: symmetries associated with the four different states—helical, A_1 , and A_2 . We note only the symmetries that are present regardless of the relative orientation of the lattice and the moments (for more details, see Table I in the Appendix). We mark in red and blue the symmetries consistent with the *T*-mod and *H*-mod measurements, respectively.

A. Multimodal approach

Scattering experiments reveal the successive onset of two propagation vectors \mathbf{Q}_1 and \mathbf{Q}_2 resulting in a nonzero expected value of the Eu magnetic moment that can be expressed as the sum over Fourier components:

$$\mathbf{M}_{\alpha}(\mathbf{r}_{i}) = \sum_{\mathbf{Q}} e^{i\mathbf{Q}\cdot\mathbf{r}_{i}} \mathbf{M}_{\alpha}(\mathbf{Q}), \qquad (4)$$

where \mathbf{r}_i denotes the position of an Eu site, and $\alpha = 1, 2$ is the Eu sublattice index. $\mathbf{Q} = \pm \mathbf{Q}_1$ in phase I and $\mathbf{Q} = \pm \mathbf{Q}_1, \pm \mathbf{Q}_2$ in phase II. However, the scattering experiments are not sufficient to uniquely determine $\mathbf{M}_{\alpha}(\mathbf{Q})$; removing this ambiguity is the objective of our analysis. We note that since both propagation vectors are perpendicular to the Eu layers ($\mathbf{Q}_1 || \mathbf{Q}_2 || \hat{\mathbf{z}}$), it is sufficient to consider the average magnetic moment in each Eu layer $\mathbf{M}_{\alpha}(z_i) = \langle \mathbf{M}_{\alpha}(\mathbf{r}_i) \rangle_{z=z_i}$ rather than the magnetic moment at each site.

The first step is to identify all magnetic structures $\mathbf{M}_{\alpha}(z_i)$ consistent with the scattering experiments, and with the fact



FIG. 8. Top row: the layer-resolved spin orientation in (a) the Néel state; (c) in the mixed structure obtained by combining A_2 order with the Néel order; the blue arrows indicate the orientation of Ψ_L ; (e) the mixed structure obeying the equal-moment condition [Eq. (9)]. Bottom row: (b) the basis functions of the $m\Gamma_5^+$ irrep corresponding to the two perpendicular orientations of the Néel order. (d),(f) The top view of two orthogonal orientations of the orientation of the Néel order shown in (c) and (e), respectively. θ denotes the orientation of the Néel order. The structure shown in panel (f) with $\theta = \pi/2$ corresponds to the broken helix considered in previous work [16].

that the magnetic moments lie in Eu planes. The next step is to identify the magnetic space groups associated with each of those magnetic configurations. The symmetry-sensitive information obtained from the optical experiments can then be used to rule out a vast majority of structures obtained in the previous step. In phase I of EuIn₂As₂, this symmetrybased approach has proven sufficient to uniquely identify the magnetic structure, while in phase II it needs to be complemented with further considerations, as explained in Sec. V.

B. Phase I: Amplitude-modulated collinear order

Each magnetic structure lowers the system symmetry from the parent paramagnetic group $P6_3/mmc1'$ (No. 194.264) to a magnetic space group (MSG) that is a subgroup of the paramagnetic group, and has the symmetry properties of one or more of its irreducible representations (irreps). We find that all magnetic configurations $\mathbf{M}_{\alpha}(z_i)$ that are consistent with the observed magnetic Bragg peaks and in-plane magnetic moments transform according to the $m\Delta_6$ irrep (see Appendixes E and F for a detailed analysis). One of them is the 60° helix proposed in earlier works [16,29], which is inconsistent with the observed birefringence. We now show that there is only one magnetic structure for phase I consistent with all the experimental results (i)–(iii) outlined above.

It is useful to rewrite the order parameter of phase I $\mathbf{M}_{\alpha}(\mathbf{Q}_1)$ as a vector in the four-dimensional representation space of $m\Delta_6$, whose basis functions are any four linearly independent arrangements of the magnetic moments that transform as the $m\Delta_6$ irrep. One possible choice of the basis functions is the four 60°-helix states of opposite helicities and orthogonal orientations corresponding to the states H_1 , H_2 and $\phi = 0$, $\phi = \pi/2$ in Figs. 7(a)–7(c). For our purposes, however, it is more convenient to consider the amplitude-modulated collinear magnetic states of different phases shown in Figs. 7(d)–7(f) and labeled by A_1 , A_2 and $\phi = 0$, $\phi = \pi/2$. In these structures, all magnetic moments point in the same direction, while their amplitude varies sinusoidally from layer to layer.

There are two classes of amplitude-modulated structures, hereafter called nodeless (denoted by A_1) and nodal (denoted by A_2). In the nodeless structure, $\mathbf{M}_{\alpha}(z_i)$ is nonzero in all Eu layers, while in the nodal structure it vanishes in every third Eu layer [Figs. 7(d) and 7(e)]. In principle, the magnetic moments associated with each of the orders $A_{1,2}$ could assume any orientation. Therefore, the four structures obtained by $A_{1,2}$ with moments pointing along two orthogonal directions form a valid basis of the $m\Delta_6$ representation space [Fig. 7(f)]. Written in this basis, the order parameter for phase I takes the form

$$\Psi_I = (a_1 \cos \phi_1, a_1 \sin \phi_1, a_2 \cos \phi_2, a_2 \sin \phi_2), \quad (5)$$

where $\phi_{1,2}$ and $a_{1,2}$ set the orientation and amplitude, respectively, of the moments associated with $A_{1,2}$. Any magnetic structure consistent with the scattering pattern observed in phase I can be obtained by specific choices of $a_{1,2}$ and $\phi_{1,2}$, and the relationship between Ψ_I and $\mathbf{M}_{\alpha}(\mathbf{Q}_1)$ is derived in Appendix F. For instance, while the amplitudemodulated phases A_1 and A_2 in Figs. 7(d) and 7(e) are described by $a_1 \neq 0$, $a_2 = 0$, and $a_2 \neq 0$, $a_1 = 0$, respectively, the 60°-helix phases H_1 and H_2 are parametrized by $a_1 = a_2$ and $\phi_2 = \phi_1 + \pi/2$, with $\phi_1 = \pi$ for the H_1 phase and $\phi_2 = \phi_1 - \pi/2$ with $\phi_1 = 0$ for the H_2 phase.

While Ψ_I is compatible with the scattering experiments for any choice of $a_{1,2}$ and $\phi_{1,2}$, optical measurements impose further constraints. Birefringence is nonzero only in structures that break the threefold rotation symmetry along z (C_{3z}), which can be recast as a nonzero value of the composite three-state Potts nematic order parameter [35],

$$\boldsymbol{\eta} = \frac{1}{2} \begin{pmatrix} a_1^2 \cos(2\phi_1) + a_2^2 \cos(2\phi_2) \\ a_1^2 \sin(2\phi_1) + a_2^2 \sin(2\phi_2) \end{pmatrix}.$$
 (6)

The condition $\eta \neq 0$ for phase I excludes the 60°-helical states and allows any of the amplitude-modulated states. This is because, as explained above, the 60°-helical states have $a_1 = a_2$ and $\phi_2 = \phi_1 \pm \pi/2$, which makes $\eta = 0$. The absence of an *H*-mod signal in phase I enables us to distinguish between the two amplitude-modulated states, as this signal is forbidden by the product of inversion and time-reversal symmetry (\mathcal{PT}). Therefore, the state A_1 is ruled out, as the Eu sites remain centers of inversion, as they were in the parent space group, and \mathcal{PT} is broken. In contrast, in A_2 inversion symmetry is broken, while \mathcal{PT} is preserved. In fact, this nodal amplitude-modulated structure is the only one belonging to the $m\Delta_6$ irrep that allows for nonzero η and vanishing *H*-mod signal. Therefore, the only magnetic structure that is consistent with all experimental findings in phase I is A_2 characterized by $a_1 = 0$ and $a_2 \neq 0$ in Eq. (5).

One remaining degree of freedom is the direction of the magnetic moments within the *a*-*b* plane set by ϕ_2 . Setting $a_1 = 0$ in Eq. (6), we note that the direction of the magnetic moment coincides with the orientation of the nematic director. Therefore, the principal axis extracted from our optical birefringence measurements corresponds to the direction of the magnetic moments. Although the broad distribution of principal axis orientations shown in Fig. 4 was measured in phase II, we demonstrated that the orientation does not change as a function of the temperature [Fig. 3(c)]. Therefore, the distribution of orientations is equally broad in phase I, indicating that ϕ_2 is not constrained. This observation is surprising from a symmetry perspective, since there are always symmetry-allowed terms in the free energy, which can be thought of as magnetocrystalline anisotropy that favors ϕ_2 pointing along the high-symmetry directions. The experimental observation therefore indicates that the magnetocrystalline anisotropy is weak compared to built-in strain.

Our work is the first to suggest an amplitude-modulated phase in EuIn₂As₂, and our conclusion is supported by the results of Mössbauer spectroscopy (reported in the Supplemental Material Fig. 10 of Ref. [16]). These measurements also confirm that the valence Eu^{2+} in $EuIn_2As_2$ is temperature independent, fixing spins on individual sites to S = 7/2 in all phases. The amplitude modulation therefore arises from the sinusoidal variation of the thermal average of the magnetic moments. Mössbauer measurements are not sensitive to the phase of amplitude modulation and, therefore, cannot distinguish between A_1 and A_2 . We note that a nodal amplitude-modulated phase had previously been observed in the ground state of Na-doped SrFe₂As₂ [40], where it originates from itinerant magnetism, in contrast to the local moment magnetism of EuIn₂As₂. The amplitude-modulated state is also reminiscent of the sinusoidal collinear state observed in some insulating multiferroic manganites with local Mn moments (see Ref. [41] and references within), although no nodes are reported there. Intriguingly, in those materials the sinusoidal phase is stabilized in a temperature window of up to 25 K below the onset of long-range magnetic order, but it is always followed by a second phase transition into a magnetic structure with constant moment length (helical or antiferromagnetic). We will now explore whether the transition into phase II could play a similar role in $EuIn_2As_2$.

C. Phase II: Mixed order

We now turn to the analysis of phase II, whose onset is characterized by the appearance of an additional ordering vector $\mathbf{Q}_2 = (0, 0, 1)$ corresponding to the emergence of Néel order [illustrated in Fig. 8(a)]. In-plane Néel order transforms as the $m\Gamma_5^+$ irrep of the paramagnetic group of EuIn₂As₂, and its order parameter takes the form of a twocomponent vector

$$\Psi_L = l(\cos\theta, \sin\theta) \tag{7}$$

in the space of $m\Gamma_5^+$. Here, *l* denotes the amplitude of the moments in the Néel state, and θ determines their orientation with respect to the in-plane crystal axes [Figs. 8(a) and 8(b)].

Phase II is, therefore, parametrized by both Ψ_I defined in Eq. (5) and Ψ_L , and all candidate magnetic states for this phase can be described by different choices of $a_{1,2}$, $\phi_{1,2}$, l, and θ . One example with $\Psi_I = (0, 0, a_2, 0)$ and $\Psi_L = (0, l)$ is shown in Figs. 8(c) and 8(d). The blue arrows indicate the direction of the Néel component. A complete list of states compatible with phase II can be found in Table II in the Appendix, where the relationship between $\mathbf{M}_{\alpha}(\mathbf{Q}_2)$ and Ψ_L is also derived.

To analyze the optical responses in phase II, we first note that any nonzero Ψ_L also introduces a Potts-nematic order parameter:

$$\boldsymbol{\xi} = \frac{l^2}{2} \begin{pmatrix} \cos 2\theta \\ \sin 2\theta \end{pmatrix}.$$
 (8)

Therefore, in the lowest-temperature phase, both $\boldsymbol{\xi}$ and $\boldsymbol{\eta}$ [Eq. (6)] contribute to the observed birefringence via the combination $\boldsymbol{\eta} + \boldsymbol{\xi}$ (see Appendix G for details). Furthermore, the Néel component breaks \mathcal{PT} , accounting for the onset of the *H*-mod signal in phase II.

From a symmetry perspective only, we cannot completely constrain the magnitudes of $a_{1,2}$ and l, or their relative orientations. Many different structures with Ψ_I and Ψ_L both nonzero break C_{3z} symmetry and \mathcal{PT} symmetry, and are therefore consistent with the optical measurements. Indeed, Table II shows various phases characterized by different combinations of the order parameters that result in different magnetic space groups. However, by taking into account the local moment nature of the EuIn₂As₂ magnetism, we can narrow down the candidates for the magnetic ground state. The magnetism originates from the Eu²⁺ localized magnetic moments (S = 7/2), with no valence mixing with Eu⁺³ [16]. At the lowest temperatures, where fluctuations associated with the magnetic phase transitions are completely suppressed, the moments in each layer should have the same magnitude. Note that this statement is not in contradiction with our proposed amplitudemodulated phase I, where the magnetic moments are small and fluctuate strongly because of their proximity to the magnetic phase transitions. Indeed, as we show below, the amplitude-modulated phase can smoothly evolve from a configuration near T_{N2} in which the moments in each layer are not the same to a configuration at lower temperatures in which the moments all have the same magnitude, consistent with the aforementioned Mössbauer spectroscopy data.

Enforcing the moments to have the same magnitude corresponds to the additional conditions on the components of the Ψ_I and Ψ_L order parameters:

$$a_1 = \sqrt{a_2^2 + 2l^2} - \sqrt{2}l, \quad \phi_1 = \phi_2 \pm \frac{\pi}{2}, \quad \theta = \phi_2 \mp \frac{\pi}{2}.$$
 (9)

We note that the broken-helix state proposed in Refs. [16,29] is a special combination of Ψ_I and Ψ_L which fulfills the equal-moment condition [Figs. 8(e) and 8(f)]. Although the equal-moment condition cannot be obeyed throughout phase II (in contrast to assumptions in the previous work), the fact that there are no phase transitions between T_{N2} and the lowest measured temperature (2 K) indicates that the symmetry at $T \leq T_{N2}$ is the same as in the equal-moment ground state. The free-energy model presented in the following section will provide a perspective on the evolution of the magnetic structure from the amplitude-modulated state of phase I to the equal-moment ground state.

In addition to the moment-length variation, the key difference between the broken helix considered previously and our findings is that the magnetic order in phase II is not constrained by the underlying lattice, and we therefore refer to this structure as the unpinned broken helix. As we discussed in the context of phase I, this observation indicates that the magnetocrystalline terms in the free energy are weak compared even to the inevitable built-in strain.

D. Comparison of the multimodal approach with other experimental techniques

We have demonstrated in the previous sections that the multimodal approach can be used to identify the magnetic structures in phase I and phase II of EuIn₂As₂. To assess the usefulness and uniqueness of this approach in determining complex magnetic structures in general, it is important to compare it to other, well-established, techniques. Conventional scattering experiments are necessary to find the propagation vectors but are not sufficient to fully determine the structure. Here we consider whether in principle, or in practice, a more advanced version of those experiments would be sufficient.

Two advanced scattering approaches could be taken: neutron diffraction with spin polarization, and resonant x-ray diffraction with polarization analysis. In a single magnetic domain both approaches could distinguish the amplitude-modulated and helical structures; this is done by comparing the measured intensities with calculations based on assumed structures. However, in a realistic sample this analysis is complicated by the presence of multiple domains within the measured volume. This problem is particularly severe for neutron-scattering experiments, which typically probe volumes of mm³, but even a focused x-ray beam is typically 1–2 orders of magnitude larger than a diffraction-limited laser spot in the visible range. Further, the usual polarized x-ray technique requires rotating the sample about the scattering vector **Q**, making it challenging to ensure that all the measurements are taken on the same sample location. A less commonly used technique called the full linear polarization analysis can avoid this problem since the azimuth of the sample is fixed; providing the x-ray beam could be focused on a single magnetic domain, this technique would offer the best chance to determine the magnetic structures in the two phases of EuIn₂As₂ without the optical measurements.

In addition to the challenges related to the domains, it is important to note that the advanced scattering experiments described above require long measurements using specialized setups at large-scale facilities. In contrast, the optical measurements can be performed in a laboratory setting. What is more, small modifications of the experimental conditions of the optical experiments yield sensitivity to distinct symmetries. For instance, a second-harmonicgeneration (SHG) experiment [42] sensitive to inversion symmetry (\mathcal{P}) could be performed in the same setup on the same sample positions as the birefringence measurements. Indeed, an SHG experiment with sensitivity to rotational anisotropy, combined with scattering and a group-theory analysis, could offer an alternative way to characterize the two magnetic phases in EuIn₂As₂. In general, the versatility of the optical probes adds to the power of the multimodal approach.

We conclude that combining conventional diffraction experiments with symmetry-sensitive optics is a powerful and efficient step to identifying complex magnetic states. In $EuIn_2As_2$, this was sufficient to fully determine the magnetic structure. Even when it is not sufficient, this approach would motivate and inform more involved scattering experiments, making the use of valuable facility time more efficient.

V. LANDAU FREE-ENERGY EXPANSION

In the previous section, we identified the symmetries of the two magnetic phases in $EuIn_2As_2$. Here, we introduce a phenomenological description of the evolution of the magnetic structure from the nodal amplitude-modulated state to the unpinned broken helix. The key finding is that it is not necessary to include two separate instabilities to describe the two phase transitions. Instead, the order Ψ_L naturally arises from coupling to the order Ψ_I of phase I.

A. Phenomenological description of the two transitions

Our starting point is the Landau free energy

$$F(\Psi_I, \Psi_L) = F_A(\Psi_I) + F_L(\Psi_L) + F_{AL}(\Psi_I, \Psi_L).$$
(10)

It has three classes of terms: F_A involves only the order parameter Ψ_I of phase I [Eq. (5)], which transforms as the $m\Delta_6$ irrep, F_L depends only on the Néel component Ψ_L of the order parameter of phase II, which transforms as the $m\Gamma_5^+$ irrep, and F_{AL} accounts for the coupling between them. The most general expressions for these three terms are given in Appendix H. For the discussion in this section, it suffices to know that only combinations of the components of Ψ_I and Ψ_L that are invariant under all symmetries of the parent paramagnetic group, including time-reversal symmetry, are allowed. In particular, F_A has a quadratic term,

$$F_A(\Psi_I) = \alpha_I (T - T_{N1}) |\Psi_I|^2 + (...), \qquad (11)$$

whose coefficient changes sign to induce the transition into phase I at T_{N1} . The remaining parameters of F_A are temperature independent and can be chosen to favor the experimentally observed nodal amplitude-modulated state.

Similarly, F_L has a quadratic term

$$F_L(\Psi_L) = \alpha_L |\Psi_L|^2 + (\dots), \qquad (12)$$

and the second transition at T_{N2} could in principle be induced by a change of its sign. However, obtaining two transitions so close in temperature would require finetuning of model parameters. Instead, Ψ_L can be induced by the coupling term F_{AL} . The form of F_{AL} determines whether these two transitions take place simultaneously or at different temperatures. Because Ψ_I and Ψ_L transform as different irreps, there can be no bilinear coupling between them. This, combined with the fact that the free energy is time-reversal even, implies that all couplings Ψ_I and Ψ_L happen at quartic order. As we show in Appendix H, there are two types of quartic couplings: a linear-cubic coupling between Ψ_L and Ψ_I and a biquadratic coupling. The former implies that the onset of Ψ_I necessarily triggers Ψ_L , i.e., that the two transitions are simultaneous. Since two transition temperatures are observed in EuIn₂As₂, linear-cubic coupling has to vanish in phase I. Indeed, as we show in Appendix H, the linearcubic term generically vanishes if the condition $a_1 = 0$ is imposed. Therefore, the observation of two transitions combined with the analysis of the Landau free energy provides independent evidence that A_2 is the state realized in phase I.

The subsequent transition to phase II can arise from the biquadratic coupling between Ψ_I and Ψ_L , which is allowed even if $a_1 = 0$. As the temperature is lowered and $|\Psi_I|^2$ increases, the biquadratic coupling renormalizes the quadratic term of the Néel order parameter according to

$$F(\boldsymbol{\Psi}_{I}, \boldsymbol{\Psi}_{L}) = (\alpha_{L} - \beta |\boldsymbol{\Psi}_{I}|^{2}) |\boldsymbol{\Psi}_{L}|^{2} + (\dots), \quad (13)$$

such that it eventually changes sign if $\beta > 0$, triggering the second phase transition at T_{N2} (see Appendix Sec. H 3 for a detailed discussion).

We have therefore shown that a symmetry-based Landau approach can reproduce the observed sequence of phase transitions. However, this approach does not uniquely predict the evolution of the magnetic structure. In particular, it does not capture the tendency toward equal-moment magnitude, which is expected based on the low-temperature Mössbauer results and microscopic considerations. Although the Landau free-energy expansion is not justified far from the phase transitions, it can capture the tendency toward the equal-moment condition if constraints on F beyond those that follow from symmetry alone are introduced. In Appendix H4, we demonstrate that fixing the ratios between some of the coefficients of F_A and F_{AL} results in a term whose minimization gives precisely the equal-moment condition. We introduce a temperature dependence of this term to model the increase in stiffness with respect to variation in moment amplitude with reduced temperature, and show that it is sufficient to induce the transition into phase II, and that the magnitudes of the moments in each layer tend to acquire the same value as one moves farther below T_{N2} .

B. The symmetry-breaking pathway

In the previous two sections, we identified the order parameters in the two phases of $EuIn_2As_2$, derived the most general form of the free energy, and constrained its parameters based on experimental findings and physical arguments. We now analyze the magnetic structures obtained by minimization of the free energy as a function of the temperature, and compute quantities that can be compared with experimental results.

In Fig. 9(a), we show the evolution of the magnetic structure with the temperature. The transition between the paramagnet and phase I at T_{N1} is characterized by breaking of the rotational C_{3z} symmetry. The magnetic structure in phase I is identified as the nodal amplitude-modulated state, in which the average magnetic moment vanishes in every third Eu layer [A_2 , Fig. 7(e)]. This moment-length variation is enabled by strong fluctuations in the vicinity of the transition, but it becomes increasingly unfavorable as the temperature is lowered. The nonlinear coupling between the nodal amplitude-modulated state and the Néel order [Ψ_L ; Fig. 8(a)] triggers the second transition, and both the Néel order and the nodeless A_1 component [Fig. 7(d)]



FIG. 9. Results of the free-energy minimization of the constrained free energy \tilde{F} [Eq. (H13)] as a function of the temperature: (a) magnetic structure; (b) scattering intensity at \mathbf{Q}_1 (red) and \mathbf{Q}_2 (blue) computed as $|\Psi_I|^2$ and $|\Psi_L|^2$, respectively; (c) the magnitude of the nematic order parameter computed as $|\eta + \xi|$; (d) the mean (red) and standard deviation (green) of the magnetic moment length extracted from the magnetic structures obtained by energy minimization. The values for the parameters of the Landau free energy are given in Appendix H. For illustrative purposes, the coefficients of the magnetocrystalline anisotropy terms in the Landau functional are set to zero (see Appendix H), so the magnetic moments in the resulting structure do not point along high-symmetry directions, consistent with the experiment. Of course, in the experiments, the magnetic moments are ultimately pinned by the local strain. All the axes are in arbitrary units.

develop at T_{N2} . At this point, the moments are neither of the same magnitude nor point along the same direction, as can be verified in Fig. 9(a). As the temperature is lowered, the minimization of the constrained free energy $[\tilde{F}, \text{Eq. (H13)}]$ yields structures with reduced moment-length variation [Fig. 9(a)], as expected from the construction of the model.

To compare the results of the energy minimization with the experimental findings, we compute $|\Psi_I|^2$ and $|\Psi_L|^2$ [Fig. 9(b)], the nematic order parameter $[|\eta + \xi|$; Fig. 9(c)], and the mean magnetic moment length, as well as its standard deviation [Fig. 9(d)]. The results of minimization of the constrained Landau free energy as a function of the temperature capture the main features of the experimental findings remarkably well: (a) $|\Psi_L|^2$ onsets at a slightly lower temperature than $|\Psi_I|^2$ and shows a steeper temperature dependence, consistent with scattering experiments [16,29]; (b) nematic order onsets at T_{N1} , consistent with the observed birefringence reported in this work; (c) the expected value of the magnetic moment varies substantially close to the transition, but the variation is suppressed by the onset of phase II, consistent with Mössbauer spectroscopy [16]. We conclude that our free-energy model captures the essential aspects of all the optical and spectroscopic measurements on EuIn₂As₂ to date.

VI. STRAIN TUNING THE MAGNETIC SYMMETRY

The sharp onset of birefringence at T_{N1} proves its magnetic origin. On the other hand, the experimental observation of the broad distribution of the principal axis orientation (Fig. 4) implies that the orientation of magnetic moments at each sample location is determined by built-in strain, which locally breaks the C_{3z} symmetry (Fig. 4). This in turn suggests that uniaxial strain can be used to control the principal axis orientation. As we demonstrate below, this is indeed possible, with implications for the tuning of magnetic symmetry.

To test the hypothesis of axis tunability, we apply compressive and tensile uniaxial strain to a bar-shaped sample using a commercial strain device (see Appendix C for experimental details). It is immediately clear from the *T*-mod birefringence measured at several strain values that the principal axes orientation can indeed be rotated by strain [Fig. 10(a)]. Systematic strain-dependent measurements [Fig. 10(b)] reveal a smooth strain dependence of the orientation, spanning the range of approximately 0°–90° corresponding to the exchange of fast and slow optical axes.

To show that the smooth strain dependence is a consequence of weak magnetocrystalline anisotropy, we include strain e in our free-energy model through coupling to the nematic order parameters η and ξ [Eqs. (6) and (8)]:

$$F_{\epsilon} = -\epsilon \boldsymbol{e} \cdot (\boldsymbol{\eta} + \boldsymbol{\xi}), \qquad (14)$$

where e is the total strain, and e is the coupling constant assumed to be the same for both η and ξ , since they both originate from the Eu²⁺ moments (Appendix G). The total strain is the sum of the built-in strain e_0 and applied strain e_a . To obtain the curve in Fig. 10(b), we vary the applied strain e_a and minimize the total free energy (other parameters are the same as in Fig. 9, and $T - T_{N1} = -3.5$, e = 0.1). We find excellent agreement with the measured strain dependence of principal axis orientation for $|e_0| =$ 0.056% [Fig. 10(b)]. The principal axis rotates smoothly, in agreement with the expectations of a three-state Potts order parameter coupled to a conjugate field in the limit of weak magnetocrystalline anisotropy [43].

To explore the consequences of this tunability, we calculate the magnetic structure that minimizes the free energy as a function of the orientation of total strain, while



FIG. 10. (a) The polarization rotation as a function of the incident polarization measured at four different values of applied strain. (b) Symbols: principal axis orientation as a function of the applied strain (positive values denote tensile strain). Agreement between the data taken while increasing and decreasing strain indicates that the tuning is reproducible, with no plastic deformations. The line is calculated by minimizing the free energy (see text for more details; parameters are the same as in Fig. 9, and $T - T_{N1} = -3.5$, $\epsilon = 0.1$), assuming a built-in strain, whose magnitude and orientation $(0.056\%, \text{ oriented at } -65^\circ)$ are extracted from the fit. The applied strain is oriented at -82° , indicating a slight misalignment (8°) between the applied strain axis and the principal optical axes of the laboratory. (c) The magnetic structures found by free-energy minimization for different orientations of total strain. The rotation of the magnetic structure has stark consequences for symmetry, with the vast majority of orientations belonging to MSG 4.7; only when L, denoted by the blue arrows, is oriented along the Eu-Eu bond direction, or perpendicular to it, are different MSGs found: 20.31 and 20.33, respectively.

keeping its magnitude fixed [Fig. 10(c)]. As expected, we find that the relative orientation of the magnetic structure and the lattice can be freely tuned. Furthermore, we analyze the symmetry of all of the structures, and we find that for

most orientations the magnetic point group is $P2_1$ (No. 4.7), which contains C_{2z} combined with a translation by three Eu layers and identity as the only symmetries. However, when the Néel vector is aligned to one of the high-symmetry directions of the lattice, additional symmetries are restored. In particular, when the Néel vector is perpendicular to the Eu–Eu bond direction, the magnetic space group identified for phase II in Ref. [16] is restored, protecting the axion insulator phase. We therefore find that it is possible to use uniaxial strain to switch on and off the symmetries required for the axion insulator state in EuIn₂As₂.

VII. SPIN HAMILTONIAN

So far, we have shown that our measurements and symmetry analysis lead to a unique identification of the two magnetic phases, as well as describing the two phase transitions. A question that remains is the microscopic origin of this behavior, which we address by developing an effective spin Hamiltonian.

As we show in the following sections, there are two prerequisites for forming the broken-helix order: (i) a long-range Heisenberg interaction \mathcal{J}_{ij} and (ii) fourth-order exchange \mathcal{J}_{ijkl} . Our proposed spin Hamiltonian

$$H = \sum_{ij} \mathcal{J}_{ij} \mathbf{S}_i \cdot \mathbf{S}_j + \sum_{ijkl} \mathcal{J}_{ijkl} (\mathbf{S}_i \cdot \mathbf{S}_j) (\mathbf{S}_k \cdot \mathbf{S}_l)$$
(15)

includes these terms and neglects terms that depend on the spin orientation with respect to the lattice. To capture the effects of the easy-plane anisotropy, and the fact that the moment length is the same for every Eu^{2+} ion in the ground state, we parametrize each spin by an angle $S_i = (\cos \theta_i, \sin \theta_i)$.

Our two theoretical approaches are complementary: The Landau free-energy expansion captures the sequence of broken symmetries and is valid in the vicinity of the transitions. On the other hand, the spin Hamiltonian describes the microscopic origin of the magnetic ground state but provides no information about the pathway between the paramagnetic state and the ground state. In Sec. VII C, we will complement the spin Hamiltonian with atomistic magnetic simulations at nonzero temperatures, hence connecting the two approaches.

A. Heisenberg term

The Heisenberg term in Eq. (15) can induce magnetic structures of arbitrary wave vector Q if coupling beyond nearest neighbors is included, with the propagation vector determined by the Fourier transform \mathcal{J}_Q of the spatially dependent exchange interaction \mathcal{J}_{ij} . The two periodicities in the ground state of EuIn₂As₂ imply that \mathcal{J}_Q is peaked at both Q_1 and Q_2 , while the sequence of transitions indicates that Q_1 is energetically more favorable. We capture this physical intuition with a Ruderman-Kittel-Kasuya-Yosida- (RKKY)



FIG. 11. (a) Main plot: Heisenberg energy of a single $Q = (0, 0, q_z)$ magnetic structure as a function of q_z . The distance dependence of the Heisenberg interaction is given by Eq. (16), with parameters $(k_1 = 0.4\pi, k_2 = \pi, J_1 = -0.1,$ $J_2 = 0.125$) chosen to reproduce the minima at $q_z = 1/3$ and $q_z = 1$, and favor the $q_z = 1/3$ state. We consider interactions up to tenth neighbor (the numerical values of the exchange parameters are given in Table III of Appendix I 2). Top panel: magnetic structures corresponding to the helix and antiferromagnetic states. (b) Main panel: the energy of two forms of the fourthorder exchange—biquadratic ($\mathcal{J}_{1212} = 0.2$, green curve) and a combination of the biquadratic and a four-spin three-site exchange ($\mathcal{J}_{1212} = \mathcal{J}_{1223} = 0.2$, purple curve), as a function of the magnetic structure interpolated between the helical (l = 0) and the antiferromagnetic (l = 1) states, while maintaining the spin normalization condition [Eq. (18)], and setting the length of each spin to 1. Biquadratic exchange exhibits degenerate minima at the helical (l = 0) and mixed (0 < l < 1) states, and therefore is not sufficient to stabilize the mixed state. However, biquadratic exchange combined with the four-spin three-site exchange exhibits one minimum in the mixed state and can stabilize it. The top row shows the magnetic structures corresponding to l = 0 (helix), l = 0.5 (mixed order), and l = 1 (AFM).

inspired expression for the effective exchange coupling (for more details, see the Appendix I 1):

$$\mathcal{J}_{ij} = J_1 \frac{\cos\left[k_1(z_{ij} - 1)\right]}{k_1 z_{ij}} + J_2 \frac{\cos\left[k_2(z_{ij} - 1)\right]}{k_2 z_{ij}}, \quad (16)$$

where $z_{ij} = |z_i - z_j|$.

In Fig. 11(a), we show the energy of a single $Q = (0, 0, q_z)$ state as a function of q_z calculated from Eq. (16) assuming a ten-neighbor Heisenberg Hamiltonian. The parameters $k_{1,2}$ and $J_{1,2}$ (given in the caption) are chosen so that the energy has two local minima at $q_z = 1/3$ and $q_z = 1$, and the minimum at $q_z = 1/3$ is slightly lower.

An energy landscape with two well-defined minima as shown in Fig. 11(a) requires narrow energy wells and consequently long-ranged real-space interactions. Two possible mechanisms for the long-range coupling are the RKKY interaction mediated by conduction electrons and the dipolar interaction, which has been suggested to induce A-type antiferromagnetism in several Eu compounds [44]. A possible scenario is that the dipolar interaction promotes the AFM order, while the RKKY interaction promotes Q_1 . Regardless, the exchange parameters used here [Fig. 11(a)] capture the tendency toward ordering at two Q values, and RKKY interaction is required for at least Q_1 .

B. Fourth-order terms

Crucially, no parameter choice within a purely Heisenberg model will promote the coexistence of two periodicities; the system simply chooses the lower of the two energy minima in Fig. 11(a), i.e., an antiferromagnet or a helix. To account for the coexistence of the two periodicities, it is necessary to move beyond the Heisenberg model. It has been shown in several itinerant systems that various forms of the four-spin exchange $[\mathcal{J}_{iikl}; \text{Eq. (15)}]$ can promote multi-q states composed of symmetryequivalent single-q states [45-47]. As we show below, we find that fourth-order terms can stabilize the coexistence of states even if they belong to different irreps, as is the case with the Q_1 and Q_2 orders in EuIn₂As₂. We note that firstprinciples calculations [48], effective Hubbard models [47], and perturbative expansions of Kondo models [49] have shown that coupling described by \mathcal{J}_{ijkl} naturally arises in itinerant systems and can be crucial for promoting exotic magnetic order.

First, we find that the biquadratic term $(S_i \cdot S_{i+1})^2$ is not sufficient to promote the mixed order. However, when combined with the symmetry-allowed three-site four-spin term, it leads to an interaction which can stabilize the mixed ground state:

$$H_4 = \mathcal{J}_{1212} (\mathbf{S}_i \cdot \mathbf{S}_{i+1})^2 + \mathcal{J}_{1223} (\mathbf{S}_i \cdot \mathbf{S}_{i+1}) (\mathbf{S}_{i+1} \cdot \mathbf{S}_{i+2}).$$
(17)

We demonstrate this by showing that the energy of the fourth-order term is lower in the mixed state than in either of the single-Q states characterized by equal moments (helix H, AFM Ψ_L). To that end, we construct a continuum of magnetic structures with varying ratios between the amplitudes of the two order parameters ($|H|/|\Psi_L|$). In these structures, equal-moment length in each layer (M) is maintained by applying the constraint expressed in Eq. (9), leading to

$$a_1 = \frac{M-l}{\sqrt{2}}, \qquad a_2 = \frac{\sqrt{-3l^2 + 2lM + M^2}}{\sqrt{2}}, \qquad (18)$$

with the orientations $\phi_1 = \phi_2 \pm \pi/2$ and $\theta = \phi_2 \mp \pi/2$. For l = 0, we find $a_1 = a_2 = M/\sqrt{2}$ corresponding to a helix, while for l = M the structure is antiferromagnetic $(a_1 = a_2 = 0)$. For each value of 0 < l < M, we calculate a_1 and a_2 , determine the corresponding layer-dependent spin orientation, and compute the energy of the different Hamiltonian terms.

In Fig. 11(b), we plot the energy of the fourth-order term [Eq. (17), $\mathcal{J}_{1212} = \mathcal{J}_{1223} = 0.2$, purple curve) as a function of l, for M = 1. The fourth-order exchange clearly exhibits a minimum for 0 < l < 1, and therefore favors the mixed state over either of the single-Q states. In contrast, the biquadratic term alone ($\mathcal{J}_{1212} = 0.2$, $\mathcal{J}_{1223} = 0$, green curve) exhibits two degenerate minima at l = 0 and 0 < l < 1, and therefore is not sufficient to stabilize the mixed state.

C. Evolution of the magnetic structure

The above analysis suggests that combining the longrange Heisenberg exchange with the four-spin terms can induce the mixed structure. To verify this, we find the magnetic ground state by solving the Landau-Lifshitz-Gilbert (LLG) equation, as implemented in the SPIRIT framework for atomistic spin simulations [50]. The broken helix is indeed the ground state, and its Fourier transform [Fig. 12(a)] shows good agreement with the scattering data [16,29]. We therefore identify a Hamiltonian whose ground state is the observed mixed order.

We extend the calculation to nonzero temperature implemented within the SPIRIT framework by adding a stochastic thermal field, whose distribution is determined by the fluctuation-dissipation theorem [50]. In Fig. 12(b), we show the amplitudes of the two scattering peaks as a function of the temperature and find two second-order transitions, with Q_2 vanishing at a lower temperature upon warming, in agreement with the experiments. However, the simulations reveal a helical, rather than amplitudemodulated (AM) state in phase I, in contrast to our experimental findings. It is perhaps not surprising that the effective spin Hamiltonian cannot capture the AM phase, since neither of its terms is likely to favor it: The Heisenberg energies of the helix and AM phases are degenerate, and the four-spin exchange terms are negligible just below the second-order transition at T_{N1} when the AM phase is stabilized due to the small value of the magnetic moment. Magnetocrystalline anisotropy, which has been suggested to promote the AM state in a few gadolinium compounds [51,52], is not included in the calculation because our data suggest it is not experimentally relevant.

Although the free energy originating from the spin Hamiltonian alone does not capture the nodal amplitudemodulated phase, the strong coupling of magnetic and electronic degrees of freedom can account for additional contributions to entropy. To illustrate why this is the case, we first look at individual snapshots of the magnetic structure in phase I obtained by the atomistic simulations, which reveal peaks at both Q_1 and Q_2 . In contrast, the average over an ensemble of 10^5 such snapshots shows only the helical Q_1 peaks, as is characteristic of phase I. Therefore, the transition into phase II is anticipated by



FIG. 12. (a),(b) Results of atomistic LLG simulations [50] based on the Hamiltonian shown in Eq. (15), with the same parameters used in Fig. 11. Each structure is obtained by averaging over 10⁵ microscopic configurations, with other details given in Appendix I2. (a) The Fourier transform of the spin structure found at T = 0, revealing two peaks at Q_1 and Q_2 . (b) The temperature dependence of scattering intensity at Q_1 and Q_2 , revealing two second-order transitions, consistent with the experiment. (c) The Fourier transform of snapshots of the magnetic structure in phase I, as well as the Fourier transform of the structure found by averaging 10⁵ such snapshots. This demonstrates antiferromagnetic fluctuations in phase I, despite vanishing static AFM order. (d) The temperature dependence of the temperature derivative of reflectivity measured at 633 nm, showing a sharp change of electronic structure at T_{N2} . This indicates that the AFM fluctuations shown in panel (d) are accompanied by fluctuations of the electronic structure.

strong AFM fluctuations. Since magnetic fluctuations are slow on electronic timescales, AFM fluctuations are accompanied by electronic ones.

Crucially, due to the strong coupling of itinerant and localized degrees of freedom in EuIn₂As₂, the fluctuations of the electronic structure that accompany AFM fluctuations are significant. Infrared studies [27] have shown that electronic structure dramatically reconstructs with the onset of the AFM order in phase II. This is consistent with our own measurement of the temperature derivative of reflectivity on the same sample used for symmetry-sensitive measurements [Fig. 12(d)], which exhibits a sharp kink at T_{N2} . Entropic effects in EuIn₂As₂ therefore cannot be correctly captured by localized spin models alone. Our findings suggest that the entropy associated with the joined system of localized spins and itinerant electrons stabilizes

the nodal amplitude-modulated state, providing a strong motivation for explicit calculations considering the magnetic and electronic degrees of freedom in $EuIn_2As_2$ on equal footing. They also yield constraints for such models: A correct calculation needs to reproduce both the nodal amplitude-modulated state in phase I and the broken-helix ground state.

The key message of our spin model is that the itinerant electrons are crucial for stabilizing the magnetism of EuIn2As2: The long-range Heisenberg interaction and the fourth-order term coupling three Eu layers are the simplest terms that can stabilize the magnetic ground state, but neither can arise just from direct exchange of the localized 4f orbitals. Instead, they are likely to be mediated by conduction electrons, with possible contributions from coupling to the lattice, as proposed in Ref. [29]. The understanding of the origin of the magnetic interactions offers a natural way to resolve a slight difference in Q_1 reported in the two scattering experiments: Only Soh et al. [29] found $Q_1 = (0, 0, 1/3)$ within experimental resolution, while Riberolles *et al.* [16] found $Q_1 \sim (0, 0, 0.303)$. Since the magnetism is mediated by conduction electrons, such differences can naturally be explained by variations in stoichiometry. In addition to explaining the existing data, this suggests that electron density could be a powerful tuning knob for magnetism in EuIn₂As₂.

VIII. DISCUSSION AND CONCLUSIONS

To summarize, we demonstrated a multimodal approach that combines scattering and symmetry-sensitive optical measurements with group-theory analysis to identify the magnetic structures in two phases of $EuIn_2As_2$. In addition to answering the open questions regarding the magnetism of $EuIn_2As_2$, we set the scene for future research in materials with strongly intertwined magnetic and electronic degrees of freedom. Finally, we identified the coupling of local magnetic and itinerant electronic degrees of freedom as the origin of the rich magnetic behavior observed in $EuIn_2As_2$.

Each of the magnetic phases that we identified has unique aspects. Amplitude-modulated structures in systems of localized moments are rare. They have been deduced on the basis of thermodynamic measurements in several Gd^{3+} compounds [51–54], as well as in multiferroic manganites [41]. In both cases, magnetocrystalline anisotropy is thought to be the driver of amplitude modulation. The fact that magnetocrystalline anisotropy is experimentally irrelevant in EuIn₂As₂ raises the intriguing possibility that its amplitude-modulated state is stabilized by the entropic effects mentioned in Sec. VII C.

A further consequence of the decoupling of magnetic states from the $EuIn_2As_2$ lattice is that the magnetic symmetry can be controlled by uniaxial strain. Although the axion insulator state in $EuIn_2As_2$ is incompatible with its metallicity, strain tuning could allow the exploration of

approximate symmetries in other candidate topological systems where coupling to the lattice is weak. Such measurements would address the following question: If a symmetry protects a topological state for a particular orientation of the magnetic structure and the lattice, how do the response functions change when the orientation is infinitesimally changed? In other words, how robust are topological responses to continuous tuning parameters? Our work identifies Eu^{2+} -based compounds, many of which are predicted to host topological states, as promising platforms for such studies.

Our understanding of itinerant electrons as mediators of magnetic interactions radically changes the picture of EuIn₂As₂: While the initial interest in this material was spurred by the notion that a simple magnetic structure will generate exotic electronic features [13], we showed that electronic states instead generate an exotic magnetic ground state. Of course, this does not mean that the electronic states themselves do not possess interesting properties. In addition to the topological phase restored for specific relative orientations of the magnetic structure and the lattice, AFM order was shown to induce altermagnetic momentum-dependent spin polarization of the electronic bands in EuIn₂As₂ [23]; the influence of such a band splitting on electron-mediated magnetic interactions is an important open question. Our work therefore challenges *ab initio* calculations to study the interplay of magnetic and itinerant degrees of freedom in EuIn₂As₂ in a realistic way, in order to capture the experimental findings on both subsystems.

The multimodal approach proved invaluable to reach the conclusions described above, offering information beyond that available to any of the techniques alone. While scattering is unique in its ability to determine ordering wave vectors, spatially resolved probes are crucial to distinguish systems that possess a given symmetry from those in which apparent symmetry is restored by averaging over domains. This was critical for recognizing that the magnetic structure in phase I is an amplitude-modulated state, rather than a helix.

Following up on our work on $EuIn_2As_2$, we concluded that the multimodal approach would be valuable in all materials in which a helical structure is deduced based on scattering alone. This is true both for the helical structures arising from frustrated exchange interactions, such as $EuIn_2As_2$, and for those arising through the Dzyaloshinskii-Moriya interaction. This combination of techniques is ideally suited to address the challenge of differentiating between multi-Q structures and an equal population of domains of a single-Q structure. It also holds great promise for guiding the emerging research field of socalled altermagnets [3], that is, materials supporting momentum-dependent spin splitting of electronic bands despite a vanishing magnetization. Distinguishing altermagnets from "conventional" antiferromagnets poses a major experimental challenge, which can be addressed by advanced scattering techniques [55,56], although such an analysis may be complicated by the presence of multiple domains. The fact that altermagnets and antiferromagnets behave differently under time-reversal symmetry makes them ideal candidates for combining scattering with optical probes of symmetry: Altermagnets can exhibit effects such as *H*-mod birefringence and the magneto-optical Kerr effect, which are prohibited by symmetry in conventional antiferromagnets. More broadly, the multimodal approach to identifying broken-symmetry phases can be employed regardless of the nature of the order: Here it was demonstrated on a material with complex magnetism, but it is also applicable to systems exhibiting charge density or orbital ordering.

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APPENDIX A: CRYSTAL GROWTH AND CHARACTERIZATION

The optical data in Figs. 2–5 were taken on the same crystal used for the resonant x-ray scattering in Ref. [29], synthesized at the University of Oxford by a method described in Ref. [29]. Since scattering and optical experiments were performed on physically the same crystal, there are no ambiguities about sample-sample variation that could complicate our multimodal approach. Samples made at UC Berkeley gave qualitatively the same optical response, with T_{N1} and T_{N2} revealed by the *T*-mod and *H*-mod measurements, respectively. The strain experiment (Fig. 10) was performed on one of those samples, with the corresponding crystal growth method described below.

Single crystals of EuIn₂As₂ were prepared by a flux method similar to Ref. [16]. Europium (99.9999% from Ames Lab) was cut up and combined in an alumina crucible under a dry nitrogen glove box atmosphere with arsenic pieces (99.99% from Alfa Aesar) and indium shot (99.9999% from Alfa Aesar) in a 1:3:12 molar ratio with total mass of 4 g. The crucible was sealed in an evacuated quartz ampule without being exposed to air. The ampule was heated in a box furnace from room temperature to 300 °C over two hours, held for one hour, then heated to 580 °C over three hours, held for two hours, then ramped to 900 °C over 20 hours before soaking at 900 °C for four hours. Finally, the ampule was slowly cooled to 770 °C at 1.8 °C/h, and spun in a centrifuge to remove excess indium flux. The method was similar to the one used in Ref. [29], with a few differences: The highest temperature in that work was 950 °C, the centrifuging temperature at 750 °C, and the Eu purity 99.99%. Both methods used heat soaks before reaching the peak temperature to avoid any high vapor pressure of arsenic and increase the homogeneity of the flux solution.

APPENDIX B: OPTICAL EXPERIMENTS

A schematic of the optical setup is shown in Fig. 13. The incident polarization angle φ of the probe laser (633 nm, 50 μ W) was set by a polarizer followed by a half-wave plate (HWP1). The measured polarization rotation from the sample was independent of the pump (780 nm, 125 μ W, chopping frequency 2 kHz) polarization. After reflecting off the sample, the pump beam was rejected by a color filter (*F*), and the probe beam polarization was further rotated by an angle $\varphi + 45^{\circ}$ by a second half-wave plate (HWP2). In the case where the polarization state was not altered by the setup or the sample, the polarization of the reflected probe beam upon exiting the second half-wave plate was an equal



FIG. 13. Schematic of the optical setup for the modulated birefringence measurements.

superposition of vertical (V) and horizontal (H) linearly polarized light. The beam was then sent through a Wollaston prism, which spatially separated the vertical and horizontal components of the light, and the two orthogonal components were each focused onto separate, unbiased photodiodes of an balanced optical bridge detector. When the V and H components had equal intensity, which occurred when the polarization state of the light was unchanged by the sample, the net photocurrent was zero; in contrast, any measured signal indicated a change of polarization, such that the technique had a high sensitivity to detecting such changes.

While changes in the final polarization state could also be introduced by birefringence or ellipticity of the setupresulting in artifacts-we largely mitigated this issue by performing thermally modulated (T-mod) and fieldmodulated (H-mod) experiments, as these setup effects were independent of both temperature and field. For thermal modulation, a pump laser was focused onto the same spot on the sample as the probe beam and optically chopped at kHz frequencies to locally modulate the temperature of the sample. A Menlo C-Fiber Femtosecond Erbium Laser (wavelength 780 nm, repetition rate 100 MHz, pulse width 100 fs) was used as a pump for convenience, but only the average power (125 μ W) was relevant for this experiment. Precisely determining the temperature increase due to the pump laser would require solving the diffusion equation with thermal diffusivity as the input. Since the absolute magnitude of the temperature variation was not relevant for the symmetry-sensitive conclusions, we did not perform this calculation. Instead, we could estimate the temperature increase from the sharpness of the observed temperaturedependent features (Fig. 3). We estimated that the temperature increase induced by the laser was no larger than 0.2 K.

For field modulation, the sample was placed in a coil that was driven with an alternating current to create an oscillating magnetic field. The experiment was sensitive only to effects which are proportional to the modulation parameter, which the artifacts arising from the setup were not. Despite this, small cross-coupling terms could still occur if there was more than one optical constant proportional to the modulation parameter, as discussed in detail in the Supplemental Material of Ref. [33].

In phase I, the sample became birefringent, rotating the polarization of the probe beam, which unbalanced the contributions of the orthogonally polarized light components entering the balanced photodiode and caused a signal to be measured. The intensity of the light admitted to each of the photodiodes could be calculated using the Jones calculus formalism, in which the polarization state of the light is represented by a vector in the (V, H) basis, and each optical component was represented by a 2×2 matrix. The Jones matrix for a half-wave plate with its fast axis rotated by θ with respect to the horizontal axis, thereby rotating the polarization by $\varphi = 2\theta$, is described by

$$J_{\rm HWP}(\theta) = \begin{pmatrix} \cos(2\theta) & \sin(2\theta) \\ \sin(2\theta) & -\cos(2\theta) \end{pmatrix}.$$
 (B1)

The Jones matrix representing the sample is

$$J_{\text{sam}}(r,b,k,\varphi_0) = R(-\varphi_0) \begin{pmatrix} r+b & k \\ -k & r-b \end{pmatrix} R(\varphi_0), \quad (B2)$$

where *r* represents the sample reflectivity, *b* the birefringence (difference in reflectivity between the *V* and *H* polarizations), and *k* for the polar Kerr effect (difference in reflectivity between the left- and right-circularly-polarized light caused by an out-of-plane magnetization). $R(\varphi_0)$ is a rotation matrix, which encodes the different orientations of the sample and its domains with respect to the lab coordinate system:

$$R(\varphi_0) = \begin{pmatrix} \cos(\varphi_0) & -\sin(\varphi_0) \\ \sin(\varphi_0) & \cos(\varphi_0) \end{pmatrix}.$$
 (B3)

Modeling the experiment in this way, the final polarization state of the reflected light is given by

$$\begin{pmatrix} E_V \\ E_H \end{pmatrix} = J_{\text{HWP2}}(\varphi/2 + 22.5^\circ) \times R(-\varphi_0) \times J_{\text{sample}}$$
$$\times R(\varphi_0) \times J_{\text{HWP1}}(\varphi/2) \times \begin{pmatrix} 1 \\ 0 \end{pmatrix}.$$
(B4)

The balanced photodiode detector measured the intensity difference between the two orthogonal polarization components $I = |E_V|^2 - |E_H|^2$. In the experiment, we rotated the two half-wave plates, which rotated the polarization of the incident probe beam relative to the sample to simulate rotation of the sample.

The data were fit to the following function:

$$I(\varphi) = A\sin[2(\varphi - \varphi_0)] + B\sin[4(\varphi - \varphi_1)] + C.$$
 (B5)

The term $A \sin[2(\varphi - \varphi_0)]$ describes the birefringence: A is the amplitude of the birefringence and φ_0 is the orientation of the sample's principal axes relative to the lab coordinate frame. The term $B \sin[4(\varphi - \varphi_1)]$ is small and arises from second-order couplings of the sample birefringence as well as weak residual birefringence and ellipticity of the setup that can cross-couple with modulated birefringence from the sample.

The offset term C typically describes the derivative of the sample magnetization with respect to the modulation parameter; this corresponds to the magneto-optical Kerr effect for the thermally modulated experiment and the magnetic susceptibility for the field-modulated experiment. However, cross-couplings between the sample and setup can result in a small offset artifact that is unrelated to these derivatives and which can be resolved when they are equal



FIG. 14. Thermally modulated (a) offset, (b) dR/dT, and (c) birefringence amplitude.

to zero. Figure 14 shows the thermally modulated offset, reflectivity, and birefringence amplitude. In the thermally modulated measurements, C is small (2 orders of magnitude smaller than the birefringence amplitude A and thermally modulated, temperature-dependent reflectivity dR/dT). Contributions to this term arise from crosscoupling between the sample and setup birefringence, ellipticity, and dR/dT, resulting in a small offset (typically at least an order of magnitude smaller than the birefringence amplitude) when no magneto-optical Kerr effect is detected in the material, as is the case for EuIn₂As₂. See Ref. [33] for an in-depth analysis of the ways in which these cross-coupling terms arise, the material and optical properties they couple, the artifacts they can create, their influence on the measured signal, and methods for mitigating them to isolate the true signal.

APPENDIX C: STRAIN EXPERIMENT AND CALIBRATION

1. Experiment

For the measurement of *T*-mode birefringence under applied strain, a $EuIn_2As_2$ sample (600 × 500 µm) was loaded into a Razorbill Instruments CS130 strain cell. The sample was secured between two titanium plates at each end with Stycast 2850FT epoxy, which were in turn bolted to the piezo-actuated jaws of the cell [Fig. 15(a)]. A custom thermal link to the cryostat base was mounted onto the strain cell, from which two copper wires were used to heat sink the sample via EPO-TEK H20E silver epoxy. Two additional wires were routed from the sample to a Cernox thermometer and resistive heater, respectively, both of which were mounted onto the strain cell body and isolated from the thermal link by an air gap. The CS130 was equipped with a capacitive sensor to monitor the displacement of the jaws, which we measured using a Keysight E4980AL LCR meter.

To explore the temperature-strain phase diagram of $EuIn_2As_2$, we measured the birefringence as a function of both the temperature and strain across the two transitions. Figure 15(b) shows the temperature dependence of the *T*-mod birefringence for different strains at a characteristic spot on the sample. For each curve, the sample was cooled to 16 K in zero applied strain, after which the voltage applied to the piezos was ramped to a fixed value at approximately 0.5 V/s. The birefringence was measured while warming under the applied voltage, allowing the reading on the strain cell capacitor to stabilize at each temperature. Because the voltage-strain relationship is temperature dependent, strictly speaking, the strain changes with temperature for each curve; however, we report the average strain and find that the variance is negligible over



FIG. 15. (a) Sample loaded into strain cell. (b) Strain dependence of birefringence vs temperature.

this small temperature range. Importantly, the transition temperature and overall shape of the birefringence with the temperature is unmodified by strain, indicating that the essential character of the phase transition remains intact with strain.

Furthermore, to demonstrate strain tunability of the magnetic structure, we swept the strain at a fixed temperature of 16 K (Fig. 10). After cooling the sample in zero applied strain, we swept the voltage on the piezos from $0 \rightarrow 20 \text{ V} \rightarrow -20 \text{ V} \rightarrow 0$, measuring the capacitance at each point.

2. Calibration

The methods for calculating the strain applied to a sample mounted in a CS130 strain cell are described in detail in the Razorbill Instruments application notes [57]. The strain cell incorporates a capacitive displacement sensor to quantify the strain in the sample for a given applied voltage to the peizo stacks. In the ideal scenario, the displacement of the capacitor plates is equivalent to the displacement of the crystal, from which the strain can be calculated as $\epsilon = \Delta L/L$. However, several caveats must be taken into account: (1) The capacitance of the sensor exhibits a temperature dependence independent of the true gap between the jaws of the cell, primarily due to a contraction of the distance between the capacitor plates at lower temperatures, (2) the differential thermal expansion between the sample and the titanium cell leads to an offset to the displacement of the sample, and (3) the finite stiffness of the epoxy and bolting plates securing the sample to cell manifests as an additional reduction to the sample displacement. Obtaining an accurate strain measurement thus requires careful consideration. Here, we describe the specific procedure used toward these corrections.

The displacement of the piezo stacks corrected for temperature-dependent contractions in the capacitor can be parametrized as

$$\Delta L_{\text{piezo}}(T,C) = \frac{\alpha}{C(T) - C_p(T)} - d_0, \qquad (C1)$$

where *T* and *C* are the temperature and measured capacitance, respectively, α and d_0 are provided by Razorbill, and $C_p(T)$ represents a temperature-dependent parallel capacitance. $C_p(T)$ can be obtained by loading a stiff titanium dummy sample, which isolates the temperature dependence of the measured capacitance to changes in the capacitor itself, and measuring the capacitance with no applied voltage $C_{\text{dummy}}(T)$ as a function of the temperature:

$$C_p(T) = C_{\text{dummy}}(T) - \frac{\alpha}{d_0}.$$
 (C2)



FIG. 16. Measured $C_{\text{dummy}}(T)$ vs T.

The experimental values for $C_{\text{dummy}}(T)$ measured in our system are shown in Fig. 16.

As mentioned above, ΔL_{piezo} ought to be corrected to obtain the true sample displacement ΔL . However, as neither the thermal expansion nor the Young's modulus are available for EuIn₂As₂, we chose to disregard these corrections in reporting strain with the understanding that the values are relative and should be taken as an upper bound to the true strain. These need not be small corrections; simulations for the case of the iron-pnictide superconductors show strain transmission of 70% due to the pliability of the epoxy, and the thermal expansion between titanium and a test sample can differ by several multiples [58,59]. However, the qualitative observation that the optical axis is tunable under strain is independent of these corrections.

APPENDIX D: MICRODOMAIN SCENARIO

In any symmetry-breaking transition, energetically equivalent but distinguishable configurations of the order parameter, i.e., domains, can be obtained by the application of the broken-symmetry operators. In EuIn₂As₂, the transition into phase I breaks the C_{3z} symmetry of the paramagnetic state, so three domains related by C_{3z} rotation are generally expected. However, our experiment does not show the expected three domains and reveals a broad orientation continuum instead (Fig. 4).

We consider a collection of small domains within the area of our laser spot. Each domain is described by a nematic director with its principle axis oriented along one of the three C_{3z} -equivalent directions of the crystal, which we label i = 0, 1, 2; taking orientation 0 to point along the *x* axis, these orientations correspond to the nematic director pointing along $i\pi/3$. We denote the population fraction of domains at orientation *i* as x_i , such that $\sum_i x_i = 1$. The reflectivity matrix for a domain at orientation *i* can be expressed as



FIG. 17. (a) Birefringence amplitude and (b) principle axis orientation as a function of x_0 and x_1 with x_2 set by the normalization condition. The ellipsoid indicates a possible phase-space trajectory that yields a constant amplitude and changing orientation. (c) Population fraction phase-space trajectories around the constant amplitude ellipsoid versus expected principal axis.

$$r_i = R(-i\pi/3) \begin{pmatrix} 1+\delta r & 0\\ 0 & 1-\delta r \end{pmatrix} R(i\pi/3), \quad (D1)$$

where $R(\theta) = \begin{pmatrix} \cos(\theta) - \sin(\theta) \\ \sin(\theta) & \cos(\theta) \end{pmatrix}$ is a typical rotation matrix.

As discussed in Appendix B, each domain at orientation *i* contributes a polarization rotation as a function of the incident polarization angle of $d\phi_i = \delta r \sin[2(\phi - i\pi/3)]$ such that the total signal is $d\phi = \sum_i x_i d\phi_i$. To first order in δr , this evaluates to

$$d\phi = A(\mathbf{x}) \sin\{2[\phi + \alpha(\mathbf{x})]\},\tag{D2}$$

where the effective birefringence amplitude $A(\mathbf{x})$ and principle axis orientation $\alpha(\mathbf{x})$ are given by

$$A(\mathbf{x}) = \delta r \sqrt{3(x_1 - x_2)^2 + (x_1 + x_2 - 2x_0)^2}, \quad (D3)$$

$$\alpha(\mathbf{x}) = \frac{1}{2} \tan^{-1} \left(\frac{\sqrt{3}(x_1 - x_2)}{x_1 + x_2 - 2x_0} \right), \tag{D4}$$

$$\mathbf{x} = (x_0, x_1, x_2).$$
 (D5)

Having developed a model for the microdomain scenario, we want to determine whether a continuously changing principle axis orientation can be consistent with a constant amplitude as observed in the experiments. To this end, we explore both quantities in the \mathbf{x} phase space, which represents the possible spread of domains at different spots on the sample. The normalization constraint $\sum_i x_i =$ 1 allows us to rewrite Eqs. (D3) and (D4) in terms of just two components of **x**, arbitrarily taken as x_0 and x_1 . Figures 17(a) and 17(b) show the effective birefringence amplitude and principle axis within this reduced parameter space. As indicated by the red ellipses, we find that the only trajectories that are constant in amplitude with changing principle axis are very fine-tuned, involving contrived changes in populations of all three domains [Fig. 17(c)]. While this scenario is technically feasible, the strict restriction on the phase space has seemingly no physical basis. On these grounds, we discard the microdomain scenario as a viable explanation of the birefringence in $EuIn_2As_2$.

The predictions based on this model in the main text show the case of only two competing domain orientations. Specifically, we define the orientation with maximum weight in Fig. 4(c) to be $\theta_0 = 0$ with respect to the crystal with population x_0 and consider the broad distribution to be a consequence of mixing in some of the other two domains; for orientations clockwise to θ_0 , we allow a population x_2 of $\phi_2 = 2\pi/3$ domains, and for orientations counterclockwise to θ_0 , we allow a population x_1 of $\phi_1 = \pi/3$ domains, taking the population of the third domain to zero. For example, $x_2 = 0$ corresponds to the bottom edges of the phase space in Fig. 17. For each point in Fig. 5(b), we then use Eqs. (D3) and (D4) to uniquely specify the amplitude expected for the measured orientation.

APPENDIX E: SYMMETRY ANALYSIS

In this section, we provide details of the symmetry analysis and group-theory methods that we applied to pinpoint the magnetic configuration for phase I and phase II of $EuIn_2As_2$.

Two ingredients are important to characterize a magnetically ordered state: the propagation vector(s) \mathbf{Q} , which determines the periodicity of the magnetic structure, and the symmetries that they preserve. The spatial dependence of the expectation values of the magnetic moments in the ordered state is given by

$$\mathbf{M}_{\alpha}(\mathbf{r}_{i}) = \sum_{\mathbf{Q}} e^{i\mathbf{Q}\cdot\mathbf{r}_{i}} \mathbf{M}_{\alpha}(\mathbf{Q}).$$
(E1)

Here, α denotes the sublattice index, and \mathbf{r}_i is the position of the atom *i* of the underlying lattice. $\mathbf{M}_{\alpha}(\mathbf{r}_i)$ transforms as one or more irreps of the parent paramagnetic space group (also called gray group). The gray group is formed by the crystal space group *G* combined with time reversal (\mathcal{T}), $G \oplus \mathcal{T}G$. The development of a magnetic order breaks \mathcal{T} alone and certain spatial operations $g \in G$, [60] but may



FIG. 18. Representative moment configurations of the class of magnetic structures listed in Tables I and II. The moments in each Eu layer are in plane and ferromagnetically aligned, pointing along the direction indicated by the arrows. The top view of each structure is shown in the top row of the figure [panels (a)–(e)]. In panels (e) and (f), the broken-helix structure fulfills the equal-moment condition in Eq. (H10).

preserve certain combinations of spatial operations and time-reversal $\mathcal{T}g$. Therefore, the magnetic order lowers $G \oplus \mathcal{T}G$ down to one of its subgroups that define the MSG G_M of the ordered state.

For EuIn₂As₂, the gray group is $P6_3/mmc1'$ (No. 194.264), $\mathbf{Q} = \mathbf{Q}_1 = \pm(0, 0, 1/3)$ for phase I and $\mathbf{Q} = \mathbf{Q}_1, \mathbf{Q}_2 = \pm(0, 0, 1)$ for phase II. In addition, $\alpha = 1, 2$ since there are two nonequivalent Eu atoms per crystallographic unit cell. For the discussion that comes later in this section, we can focus on the dependence of $\mathbf{M}_{\alpha}(z_i)$ on the *z* component of the position of the Eu layer *i*, since the moments are ferromagnetically aligned within each plane and the ordering vectors are perpendicular to the direction \hat{z} of stacking of the Eu planes (see Fig. 18).

Note that throughout the paper, we write **Q** in terms of the reciprocal lattice vectors $\mathbf{Q} = (h,k,l) = h\mathbf{b}_1 + k\mathbf{b}_2 + l\mathbf{b}_3$,

where $\mathbf{b}_1 = (4\pi/\sqrt{3}a)\hat{x}$, $\mathbf{b}_2 = (2\pi/\sqrt{3}a)(-\hat{x}+\sqrt{3}\hat{y})$, and $\mathbf{b}_3 = (2\pi/c)\hat{z}$.

Magnetic structures with the same \mathbf{Q} but distinct $\mathbf{M}_{\alpha}(\mathbf{Q})$ have different symmetry properties and transform as distinct irreps of the parent gray group. In this section, we explain how to get a list of all magnetic structures and corresponding $\mathbf{M}_{\alpha}(\mathbf{Q})$ that are candidates for phases I and II from a symmetry perspective only. We further narrow it down based on experimental evidence. Our multimodal protocol can be summarized as follows:

(i) Step 1. Given the paramagnetic space group and the ordering vector(s) for the phase in case, the list of all magnetic space groups that are subgroups of the paramegnetic groups and can be reached by a magnetic phase transition can be readily obtained using the software ISOTROPY [61]. The corresponding irreps and order parameters are also listed.

- (ii) Step 2. For each irrep, there are multiple magnetic structures belonging to different magnetic space groups. Coupled irreps are also possible. We constrained which irrep to focus on by requiring that the magnetic structure has magnetic moments lying on the Eu planes and magnetic Bragg peaks consistent with scattering experiments [16,29]. These constraints singled out irrep $m\Delta_6$ for phase I and $m\Delta_6$ coupled with $m\Gamma_5^+$ for phase II.
- (iii) Step 3. All the magnetic structures that transform as the irrep(s) selected in step 2 are listed in Table I for phase I and Table II for phase II.
- (iv) Step 4. The symmetries of each magnetic structure from step 3 are compared with the constraints set by the optical measurements, which are used to rule out a subset of them.

For phase I, our multimodal approach allowed us to identify three states that are consistent with both scattering and optical experiments. They correspond to nodal amplitude-modulated states with different orientations of the magnetic moments. One of the possibilities is that the moments are parallel to $[100] = \hat{x}$, as illustrated in Fig. 18(b) and characterized by an order parameter $\Psi_I = (a, a, 0, 0)$. Because of the high-symmetry direction of the moments, this structure preserves the twofold rotation around \hat{y} ($C_{2[010]}$), as well as a nonsymmorphic twofold rotation around $\hat{z}(C_{27})$ and the product between time reversal and inversion (TP), and belongs to the MSG Cm'c'm' (No. 65.564). Note that any structure obtained from $\Psi_I = (a, a, 0, 0)$ by a threefold rotation around $z(C_{3z})$ is a related domain and belongs to the same MSG. The second allowed nodal structure is characterized by the order parameter $\Psi_I = (0, 0, a, -a)$ and has moments parallel to \hat{y} . This is also a high-symmetry direction in the Eu plane. The moments break $C_{2[010]}$ but preserve a mirror symmetry $M_{[010]} = \mathcal{P}C_{2[010]}$. The third allowed nodal structure is parametrized by $\Psi_I =$ (a, a, b, -b) and is a linear superposition of the two structures mentioned before. As a result, the magnetic moments point along a direction that is not a high-symmetry one, and neither $M_{[010]}$ nor $C_{2,[010]}$ are preserved. The only symmetries left in the generic orientation nodal structure are the nonsymmetric C_{2z} and \mathcal{PT} , lowering the MSG to $P2_1/m'$ (No. 11.53).

TABLE I. Candidate magnetic structures for phase I that are consistent with the experimentally observed Bragg peaks $\mathbf{Q}_1 = (0, 0, 1/3)$ and magnetic moments lying within the Eu planes. For each structure, we specify the form of the order parameter Ψ_I written in the basis of the representation space of $m\Delta_6$ adopted by ISOTROPY (helical basis). The transformation matrix between this basis and the collinear basis that we adopt in this work is shown in Appendix F 3. The MSG and its generators are also listed. 1 denotes the identity operation, \mathcal{P} denotes spatial inversion, and \mathcal{T} represents time reversal. $C_{n,[uvw]}$ denotes a rotation of $2\pi/n$ around the axis $[uvw] = u\mathbf{a}_1 + v\mathbf{a}_2 + \mathbf{a}_3$, where \mathbf{a}_j are primitive crystal axes. We set $[001] \|\hat{z}$. The mirror operations are $M_{[uvw]} = \mathcal{P}C_{2,[uvw]}$. When any of these operations (generically denoted by g) is combined with a translation $\tau = \tau_1 \mathbf{a}_1 + \tau_2 \mathbf{a}_2 + \tau_3 \mathbf{a}_3$ by a fraction of the primitive unit cell, we have a nonsymmorphic operation $\{g|\tau_1\tau_2\tau_3\}$. The last column shows the two optical responses considered in this work. \checkmark (×) signifies that the response is symmetry allowed (forbidden).

				Optical r	responses
Magnetic structure	Order parameter	MSG	Generators of the MSG	<i>T</i> -mod signal	<i>H</i> -mod signal
60° helix	(0, 0, 0, a)	<i>P</i> 6 ₁ 2'2' (178.159)	1, $\{C_{2z} 00\frac{3c}{2}\}, \{C_{3z} 00c\}, \mathcal{T}C_{2,[010]}$	×	×
	(0, 0, a, 0)	<i>P</i> 6 ₅ 2'2' (179.165)	1, $\{C_{2z} 00\frac{3\tilde{c}}{2}\}, \{C_{3z} 002c\}, \mathcal{T}C_{2,[010]}$	×	×
	(a, 0, 0, 0)	<i>P</i> 6 ₅ 22 (179.161)	$1, \{C_{2z} 00\frac{3c}{2}\}, \{C_{3z} 002c\}, C_{2,[010]}$	×	×
	(0, a, 0, 0)	<i>P</i> 6 ₁ 22 (178.155)	1, $\{C_{2z} 00\frac{3c}{2}\}, \{C_{3z} 00c\}, C_{2,[010]}$	×	×
	(a, 0, b, 0)	$P6_5$ (170.117)	1, $\{C_{2z} 00\frac{3c}{2}\}, \{C_{3z} 002c\}$	×	×
Colinear nodal amplitude modulated	(a, a, 0, 0)	<i>Cm'c'm'</i> (63.465)	1, $\{C_{2z} 00\frac{3c}{2}\}, C_{2,[010]}, TP$	\checkmark	×
Colinear nodeless amplitude modulated	(0, 0, a, a)	<i>Cm</i> ′ <i>c</i> ′ <i>m</i> (63.462)	1, $\{C_{2z} 00\frac{3c}{2}\}, \mathcal{T}C_{2,[010]}, \mathcal{P}$	\checkmark	\checkmark
Colinear nodal amplitude modulated	(0, 0, a, -a)	<i>Cmcm</i> ′ (63.461)	1, $\{C_{2z} 00\frac{3c}{2}\}, M_{[010]}, TP$	\checkmark	×
Colinear nodeless amplitude modulated	(a, -a, 0, 0)	<i>Cmcm</i> (63.457)	1, $\{C_{2z} 00\frac{3c}{2}\}, C_{2,[010]}, \mathcal{P}$	\checkmark	\checkmark
Colinear nodeless amplitude modulated	(a, a, b, b)	$Cm'c'2_1$ (36.176)	1, $\{C_{2z} 00\frac{3c}{2}\}, TM_{[010]}$	\checkmark	\checkmark
Colinear nodeless amplitude modulated	(a, -a, b, -b)	$Cmc2_1$ (36.172)	1, $\{C_{2z} 00\frac{3c}{2}\}, M_{[010]}$	\checkmark	\checkmark
Distorted helix	(0, 0, a, b)	$C2'2'2_1$ (20.33)	1, $\{C_{2z} 00\frac{3c}{2}\}, \mathcal{T}C_{2,[010]}$	\checkmark	\checkmark
Distorted helix	(a, b, 0, 0)	C222 ₁ (20.31)	1, $\{C_{2z} 00\frac{3c}{2}\}, C_{2,[010]}$	\checkmark	\checkmark
Colinear nodal amplitude modulated	(a, a, b, -b)	$P2_1/m'$ (11.53)	1, $\{C_{2z} 00\frac{3c}{2}\}, TP$	\checkmark	×
Colinear nodeless amplitude modulated	(a, -a, b, b)	$P2_1/m$ (11.50)	1, $\{C_{2z} 00\frac{3c}{2}\}, \mathcal{P}$	\checkmark	\checkmark
Distorted helix	(a, b, c, d)	<i>P</i> 2 ₁ (4.7)	1, $\{C_{2z} 00\frac{3c}{2}\}$	1	~

TABLE II. Candidate magnetic structures for phase II that are consistent with the experimentally observed Bragg peaks $\mathbf{Q}_1 = (0, 0, 1/3)$ and $\mathbf{Q}_2 = (0, 0, 1)$. All of the structures are allowed by symmetry to have finite *T*-mod and *H*-mod optical responses. For each structure, we specify the form of the order parameter Ψ_I and the antiferromagnetic component Ψ_L written in the helical basis (see Appendixes F 1 and F 3). The last column shows the MSG for each structure, whose generators can be found in Table I.

Magnetic structure	Ψ_L	Ψ_I	MSG
Colinear nodeless amplitude modulated	$(0,a) \\ (a,\sqrt{3}a)$	$(0,0,b,b)\ (-rac{b}{2},-b,-rac{\sqrt{3}b}{2},0)$	<i>Cm'c'm'</i> (No. 63.462) <i>Cmcm</i> (No. 63.457)
	$(a,\sqrt{3}a)$	$(\frac{-b+\sqrt{3}c}{2}, -b, -\frac{\sqrt{3}b+c}{2}, -c)$	<i>Cm</i> ′ <i>c</i> ′2 ₁ (No. 36.176)
	(0, a)	(b,b,c,c)	<i>Cmc</i> 2 ₁ (No. 36.172)
Broken helix	$(0,a) \\ (a,\sqrt{3}a)$	(0, 0, b, c) $(\frac{-b}{2}, c, -\frac{\sqrt{3}b}{2}, 0)$	<i>C</i> 2′2′2 ₁ (No. 20.33) <i>C</i> 222 ₁ (No. 20.31)
	(a,b) (a,b)	(c, -c, d, d) (c, d, e, f)	$P2_1/m$ (No. 11.50) $P2_1$ (No. 4.7)

Note that the sixth-order terms in the Landau functional (see Appendix H) enforce the moments to point along a high-symmetry direction, reflecting the underlying crystalline anisotropy. However, in the regime where built-in strain dominates over such a crystal anisotropy, the nodal structure with lower symmetry is generically favored by uniaxial strain along an arbitrary direction. Motivated by the observed broad distribution of crystal axes, we will consider this regime.

For phase II, on the other hand, all structures in Table II are consistent with both scattering and optical experiments. Among them, the three broken-helix structures are more likely to describe phase II since they are favored by the equal-moment condition, which in turn is expected if the magnetism of EuIn₂As₂ arises from localized S = 7/2 Eu²⁺ moments (see Appendix H4). Similar to the three nodal amplitude-modulated structures possible for phase I, the three broken helices in phase II are distinguished by the orientation of the Néel component with respect to the crystal axis. Once again, crystal anisotropy favors moments parallel to the in-plane high-symmetry direction, but the dominant built-in strain favors a less symmetric orientation of the moments [MSG $P2_1$ (No. 4.7)]. Given the broad distribution of built-in strain direction suggested by our optical experiments, the broken helix in phase II is essentially unpinned from the crystal axes, presumably due to built-in strain.

Representatives of each type of state appearing in Tables I and II are shown in Fig. 18. Note that in phase I there is a class of structures labeled "distorted helix," which is obtained by a superposition of an amplitude-modulated structure and a 60° helix. The moments get distorted in comparison with the more symmetric 60° counterpart, but there is still a well-defined single-handed circulation of the moments' direction around \hat{z} . In phase II, an AFM component is also added to the magnetic structure, in agreement with the development of the additional $Q_2 = (0, 0, 1)$ ordering vector. When strong enough, this AFM component leads to the interchange of four moment

directions, leading to a broken helix similar to that originally introduced in Ref. [16] [compare Figs. 18(d) and 18(e)]. However, as mentioned in the main text, a key difference is that the broken helix proposed in this work is not pinned to a high-symmetry direction in the Eu plane due to the strong effect of the built-in strain.

APPENDIX F: REPRESENTATIONS OF THE ORDER PARAMETER

In this section, we address equivalent ways of representing the order parameters $\mathbf{M}_{\alpha}(\mathbf{Q}_1)$ and $\mathbf{M}_{\alpha}(\mathbf{Q}_2)$ for phases I and II of EuIn₂As₂. $\mathbf{M}_{\alpha}(\mathbf{Q})$ are the Fourier components of the magnetic structure $\mathbf{M}_{\alpha}(z_i)$ in each phase and, therefore, is a pseudovector in the Euclidean space. $\mathbf{M}_{\alpha}(z_i)$ transforms as one or more irreps of the parent gray space group. Within each irrep of the space group, the spatial operations in the group $g \in G \oplus TG$ can be represented by a matrix. These matrices are defined in a vector space called representation space and have a dimension given by the product of the dimension of the corresponding irrep of the little group of \mathbf{Q} and the number of legs in the star of \mathbf{Q} . For the irreps relevant for EuIn₂As₂, $m\Delta_6$ of $P6_3/mmc1'$ associated with the ordering vector \mathbf{Q}_1 is four dimensional, and the irrep $m\Gamma_5^+$ associated with \mathbf{Q}_2 is two dimensional.

It is useful to represent $M_{\alpha}(\mathbf{Q}_1)$ and $M_{\alpha}(\mathbf{Q}_2)$ in the representation space of $m\Delta_6$ and $m\Gamma_5^+$, respectively. We start with $M_{\alpha}(\mathbf{Q}_1)$ in the next section.

1. Order parameter for phase I

The order parameter for phase I takes the form of a fourcomponent vector in the representation space of $m\Delta_6$,

$$\Psi_I = (\Delta_1, \Delta_2, \Delta_3, \Delta_4), \tag{F1}$$

where Δ_j are real numbers. The basis of the representation space is defined in terms of four basis functions. The transformation of the basis functions upon the action of the symmetry operations of the space group is then used to construct the matrix representation of each operation within the irrep. The choice of basis function is arbitrary since there are many possible choices of basis vectors for a vector space. The order parameters Ψ_I obtained in ISOTROPY [61] are written in a basis that we call the helical basis. This means that the basis functions correspond to four independent helical magnetic structures that we denote by $\mathbf{f}_{x\pm}(z_i)$ and $\mathbf{f}_{y\pm}(z_i)$:

$$(1, 0, 0, 0) \to \mathbf{f}_{x-}(z_i),$$
 (F2)

$$(0, 1, 0, 0) \to \mathbf{f}_{x+}(z_i),$$
 (F3)

$$(0, 0, 1, 0) \to \mathbf{f}_{y-}(z_i),$$
 (F4)

$$(0, 0, 0, 1) \to \mathbf{f}_{y+}(z_i),$$
 (F5)

where

$$\mathbf{f}_{x\mp}(z_i) = \left[\pm \cos\left(\frac{2\pi z_i}{3c}\right), -\sin\left(\frac{2\pi z_i}{3c}\right), 0\right]^T, \quad (F6)$$

$$\mathbf{f}_{y\mp}(z_i) = \left[\mp \sin\left(\frac{2\pi z_i}{3c}\right), -\cos\left(\frac{2\pi z_i}{3c}\right), 0\right]^T.$$
(F7)

Here, the superscript *T* denotes the transpose of the vector. $\mathbf{f}_{x\pm}(z_i)$ is a vector function in Euclidian space and describes the 60° helix with clockwise (–) or counterclockwise (+) helicity and moments parallel to \hat{x} every third Eu layer. Recall that $z_i = 0, c/2, c, ...$ Similarly, $\mathbf{f}_{y\pm}(z_i)$ describes 60° helices with opposite helicities with moments parallel to \hat{y} in every third Eu layer. A generic magnetic structure associated with the ordering vector Ψ_I in Eq. (F1) can thus be written as

$$\mathbf{M}(z_i^{\alpha}) = \Delta_1 \mathbf{f}_{x-}(z_i^{\alpha}) + \Delta_2 \mathbf{f}_{x+}(z_i^{\alpha}) + \Delta_3 \mathbf{f}_{y-}(z_i^{\alpha}) + \Delta_4 \mathbf{f}_{y+}(z_i^{\alpha}).$$
(F8)

Here, we define $M_{\alpha}(z_i) \equiv M(z_i^{\alpha})$, where z_i^{α} denotes the position of the Eu layer belonging to the sublattice α . Note that $z_i^{\alpha} = 0, c, 2c, ...$ for $\alpha = 1$ and $z_i^{\alpha} = c/2$, 3c/2, 5c/2, ... for $\alpha = 2$.

We now have all the ingredients needed to relate Ψ_I and $\mathbf{M}_{\alpha}(\mathbf{Q}_1)$. Inverting Eq. (E1), substituting into it Eq. (F8), and recalling that $z_i = 0, c, 2c, ...$ for sublattice $\alpha = 1$ and $z_i = c/2, 3c/3, 5c/c, ...$ for sublattice $\alpha = 2$, we obtain

$$\frac{\mathbf{M}_1(\mathbf{Q}_1)}{N_m} = \frac{\mathbf{M}_2(\mathbf{Q}_1)}{N_m} = \frac{3}{2} \begin{pmatrix} \Delta_1 - \Delta_2 + i(\Delta_3 - \Delta_4) \\ -\Delta_3 - \Delta_4 + i(\Delta_1 + \Delta_2) \\ 0 \end{pmatrix},$$
(F9)

where N_m denotes the number of magnetic unit cells.

In Appendix F 3, we define another basis which is constructed from orthogonal collinear structures rather than helical. We call it the collinear basis, and this is the basis adopted in the discussion carried out in the main text. The helical-to-collinear basis transformation is provided in Appendix F 3.

2. Order parameter for phase II

We now repeat the analysis for phase II, which is characterized by both Ψ_I and by a Néel component to the magnetic order. The latter takes the form of a twocomponent vector in the representation space of $m\Gamma_5^+$,

$$\Psi_L = (l\cos\theta, l\sin\theta). \tag{F10}$$

l and θ are real numbers, and their meaning will become clear shortly. As before, we need to specify the basis functions for $m\Gamma_5^+$. We adopt the same basis used as ISOTROPY [61], which consists of two orthogonal in-plane AFM configurations:

$$(1,0) \to \mathbf{g}_x(z_i), \tag{F11}$$

$$(0,1) \to \mathbf{g}_{y}(z_{i}), \tag{F12}$$

with

$$\mathbf{g}_{x}(z_{i}) = (-1)^{\frac{2z_{i}}{c}}(1,0,0)^{T}, \qquad (F13)$$

$$\mathbf{g}_{y}(z_{i}) = (-1)^{\frac{2z_{i}}{c}}(0, 1, 0)^{T}.$$
 (F14)

 $\mathbf{g}_x(z_i)$ and $\mathbf{g}_y(z_i)$ are thus vector functions in Euclidean space that describe an AFM order with moments pointing along \hat{x} and \hat{y} , respectively. A generic Néel order with order parameter Ψ_L in Eq. (F10) can thus be written as

$$\mathbf{M}(z_i^{\alpha}) = l\cos\theta \mathbf{g}_x(z_i^{\alpha}) + l\sin\theta \mathbf{g}_y(z_i^{\alpha}).$$
(F15)

Recall that $z_i^{\alpha} = 0, c, 2c, ...$ for $\alpha = 1$ and $z_i^{\alpha} = c/2, 3c/2, 5c/2, ...$ for $\alpha = 2$. From this equation, we can readily see that *l* and θ denote, respectively, the amplitude and orientation with respect to the lattice axes of the moments in the Néel state.

The relation between $\mathbf{M}_{\alpha}(\mathbf{Q}_2)$ and Ψ_L is obtained by inverting Eq. (E1) and using Eq. (F15):

$$\frac{\mathbf{M}_1(\mathbf{Q}_2)}{N_m} = \frac{\mathbf{M}_2(\mathbf{Q}_2)}{N_m} = 3 \begin{pmatrix} l\cos\theta\\l\sin\theta\\0 \end{pmatrix}.$$
(F16)

3. Basis choice for the representation space

A more convenient choice of basis functions for the magnetic order $M_{\alpha}(\mathbf{Q}_1)$ consists of using the following

four orthogonal collinear magnetic structures obtained by linear combinations of the helical basis functions [Eqs. (F6) and (F7)]:

$$\mathbf{f}_{A_{1,x}}(z_{i}) = \frac{1}{\sqrt{2}} (\mathbf{f}_{x+}(z_{i}) - \mathbf{f}_{x-}(z_{i})), \qquad (F17)$$

$$\mathbf{f}_{A_{1},y}(z_{i}) = \frac{1}{\sqrt{2}} (\mathbf{f}_{y-}(z_{i}) + \mathbf{f}_{y+}(z_{i})), \qquad (F18)$$

$$\mathbf{f}_{A_{2},x}(z_{i}) = \frac{1}{\sqrt{2}} (\mathbf{f}_{y+}(z_{i}) - \mathbf{f}_{y-}(z_{i})), \qquad (F19)$$

$$\mathbf{f}_{A_{2},y}(z_{i}) = -\frac{1}{\sqrt{2}} (\mathbf{f}_{x+}(z_{i}) + \mathbf{f}_{x-}(z_{i})).$$
(F20)

These combinations result in amplitude-modulated structures, where the moments in all Eu layers point along the same direction, while the norm of the moments vary sinusoidally from layer to layer. In the structure $\mathbf{f}_{A_1,x}(z_i)$ $[\mathbf{f}_{A_1,y}(z_i)]$ referred as A_1 , the moments point along \hat{x} (\hat{y}), and the node of the sinusoidal amplitude modulation lies in between Eu planes. These are the nodeless amplitudemodulated structures defined in the main text. Similarly, $\mathbf{f}_{A_2,x}(z_i)$ $[\mathbf{f}_{A_2,y}(z_i)]$ are nodal amplitude-modulated structures denoted A_2 with moments pointing along x (y) and nodes at every third Eu layer.

The order parameter for phase I [Eq. (F1)] in this colinear basis takes the form shown in the main text:

$$\Psi_I = (a_1 \cos \phi_1, a_1 \sin \phi_1, a_2 \cos \phi_2, a_2 \sin \phi_2).$$
 (F21)

Here, $a_{1,2}$ and $\phi_{1,2}$ set the amplitude and in-plane orientation of the moments in $A_{1,2}$. The transformation between the helical and the colinear basis is given by

$$\begin{pmatrix} a_1 \cos \phi_1 \\ a_1 \sin \phi_1 \\ a_2 \cos \phi_2 \\ a_2 \sin \phi_2 \end{pmatrix} = \frac{1}{\sqrt{2}} \begin{pmatrix} -1 & 1 & 0 & 0 \\ 0 & 0 & 1 & 1 \\ 0 & 0 & -1 & 1 \\ -1 & -1 & 0 & 0 \end{pmatrix} \begin{pmatrix} \Delta_1 \\ \Delta_2 \\ \Delta_3 \\ \Delta_4 \end{pmatrix}.$$
(F22)

As a result, Eq. (F9) can be reexpressed as

$$\frac{\mathbf{M}_{1}(\mathbf{Q}_{1})}{N_{m}} = \frac{\mathbf{M}_{2}(\mathbf{Q}_{1})}{N_{m}} = -\frac{3}{\sqrt{2}} \begin{pmatrix} a_{1}\cos\phi_{1} + ia_{2}\cos\phi_{2} \\ a_{1}\sin\phi_{1} + ia_{2}\sin\phi_{2} \\ 0 \end{pmatrix}.$$
(F23)

APPENDIX G: NEMATIC DIRECTOR FOR PHASE II

In phase II, we have the coexistence of two Potts-nematic order parameters, η which is related to Ψ_I , and ξ that is

related to the Néel state that emerges in phase II. Therefore, the general form of the nematic order parameter associated with the magnetic structure in phase II is $\eta + \kappa \xi$, where κ is a real number. In this section, we show that for the forms of η and ξ adopted in the main text [Eqs. (6) and (8)], $\kappa = 1$ is the most natural choice.

The nematic director can be calculated directly from the magnetic structure $\mathbf{M}(z_i)$ by

$$\boldsymbol{n} = \frac{1}{N_l} \sum_{z_i=0}^{5c/2} \binom{M_x^2(z_i) - M_y^2(z_i)}{2M_x(z_i)M_y(z_i)}.$$
 (G1)

Here, x, y denote the Cartesian components of $M(z_i)$. In addition, $N_l = 6$ corresponds to the number of Eu layers in the magnetic unit cell. Finite birefringence requires a nonzero n.

In phase II, $M(z_i)$ takes the following form in the collinear basis:

$$\boldsymbol{M}(z_i) = \sum_{j=1}^{2} \left[a_j \cos \phi_j \boldsymbol{f}_{A_j, x}(z_i) + a_j \sin \phi_j \boldsymbol{f}_{A_j, y}(z_i) \right] + l \cos \theta \boldsymbol{g}_x(z_i) + l \sin \theta \boldsymbol{g}_y(z_i), \qquad (G2)$$

where $g_{\mu}(z_i)$, $f_{A1,\mu}(z_i)$, and $f_{A2,\mu}(z_i)$ (with $\mu = x, y$) are the basis functions defined in Appendix F. Substituting Eq. (G2) into Eq. (G1), we obtain

$$\boldsymbol{n} = 2(\boldsymbol{\eta} + \boldsymbol{\xi}). \tag{G3}$$

APPENDIX H: FREE-ENERGY FUNCTIONAL

In this section, we derive the analytic form of the Landau functional discussed in the main text. The generic Landau function for EuIn₂As₂ that captures a sequence of phase transitions where \mathbf{Q}_1 and \mathbf{Q}_2 emerge takes the form of a power series on the components of the order parameters Ψ_I and Ψ_L defined in the main text (see also Appendix F). Ψ_I is the order parameter for phase I [Eq. (5)], and Ψ_L is the order parameter corresponding to the AFM component of the magnetic state that develops in phase II [Eq. (F10)]. The terms in the power series are constrained by the symmetries of the paramagnetic space group $P6_3/mmc1'$ (No. 194.264) of EuIn₂As₂ and can be readily obtained using the tool INVARIANTS in the ISOTROPY software [61] with the condition of coupled $m\Delta_6$ and $m\Gamma_5^+$ irreps. They can be grouped into three classes of terms,

$$F(\Psi_I, \Psi_L) = F_A(\Psi_I) + F_L(\Psi_L) + F_{AL}(\Psi_I, \Psi_L).$$
(H1)

We now show the expressions for each of them.

1. The form of F_A

We start with F_A , which involves only the components of Ψ_I . In the helical basis [Eq. (F1)] and up to order $\mathcal{O}(\Delta_j^6)$, it reads

$$F_{A} = \alpha_{1} \left(\sum_{j} \Delta_{j}^{2} \right) + \frac{\beta_{1}}{2} \left(\sum_{j} \Delta_{j}^{2} \right)^{2} + \alpha_{2} \boldsymbol{\eta} \cdot \boldsymbol{\eta} - \frac{\beta_{2}}{2} \eta_{1} (\eta_{1}^{2} - 3\eta_{2}^{2}) + \frac{\beta_{4}}{2} \left(\sum_{j} \Delta_{j}^{2} \right)^{3} + \gamma_{3} \left(\sum_{j} \Delta_{j}^{2} \right) \boldsymbol{\eta} \cdot \boldsymbol{\eta} + \gamma_{1} [\Delta_{3}^{2} (\Delta_{3}^{2} - 3\Delta_{1}^{2})^{2} + \Delta_{4}^{2} (\Delta_{4}^{2} - 3\Delta_{2}^{2})^{2}] - \frac{\gamma_{2}}{2} (\Delta_{1} \Delta_{2} - \Delta_{3} \Delta_{4}) [\Delta_{1}^{2} (\Delta_{2}^{2} - 3\Delta_{4}^{2}) - 8 \Delta_{1} \Delta_{2} \Delta_{3} \Delta_{4} + \Delta_{3}^{2} (\Delta_{4}^{2} - 3\Delta_{2}^{2})], \qquad (H2)$$

where

$$\mathbf{\eta} = \begin{pmatrix} \eta_1 \\ \eta_2 \end{pmatrix} = \begin{pmatrix} -\Delta_1 \Delta_2 - \Delta_3 \Delta_4 \\ \Delta_2 \Delta_3 - \Delta_1 \Delta_4 \end{pmatrix}$$
(H3)

is a three-state Potts-nematic composite order parameter [35] allowed to develop in phase I. In the main text, η is expressed in the collinear basis (see Appendix F 3). We emphasize that the sixth-order terms are essential to remove the degeneracy between different amplitude-modulated phases.

Note that $\mathbf{\eta}$ vanishes for choices of Δ_j resulting in magnetic structures that preserve the threefold rotation symmetry around \hat{z} (C_{3z}). For instance, $\mathbf{\eta} = 0$ if $\Delta_2 = \Delta_3 = \Delta_4 = 0$ in Eq. (F1), which is the order parameter for a 60° helix. Importantly, $\eta \neq 0$ is required for a finite *T*-mod signal. Therefore, all of the magnetic structures listed in Tables I and II with an allowed *T*-mod optical signal have a nonzero nematic component.

Although the values for the parameters in Eq. (H2) could be determined through a microscopic theory for magnetism in EuIn₂As₂, this is beyond the scope of this work, and a phenomenological approach suffices to capture the sequence of observed phase transitions in this material. The parameter $\alpha_1 = \alpha_0(T - T_{N1})$ with $\alpha_0 > 0$ changes sign at the onset of phase I, at temperature T_{N1} . Moreover, $\beta_1 > 0$ and $\beta_4 > 0$ guarantee that F_A is bounded. The parameters α_2 and γ_3 , when chosen to be negative, energetically favor nematic magnetic structures.

Rewriting F_A in the collinear basis (see Appendix F 3), we find that β_2 and γ_1 are the coefficients of terms involving combinations of $\sin \phi_{1,2}$ and $\cos \phi_{1,2}$ and, therefore they favor specific orientations of the magnetic moments within the Eu planes. Throughout this work, motivated by the experimental observation that the magnetic moments are not pinned to the crystal axes, we focus on the limit where β_2 and γ_1 are much smaller than the energy scale associated with built-in strain. In practice, to simplify the analysis, we will set these coefficients to zero and choose an arbitrary direction for the moment in phase I. Of course, these coefficients are not identically zero, and the direction is set by the local strain, as discussed in the main text.

The coefficient γ_2 distinguishes between the nodeless $(a_1 \neq 0)$ and nodal $(a_2 \neq 0$ and $a_1 = 0)$ amplitudemodulated structures. This can be readily seen in the collinear basis, where the term proportional to γ_2 in Eq. (H2) simplifies to

$$\frac{1}{16}(a_1^2 - a_2^2)[(a_1^2 - a_2^2)^2 - 12a_1^2a_2^2\cos^2(\phi_1 - \phi_2)].$$
(H4)

2. The form of F_L

The second term on the right-hand side of Eq. (H1) involves only components of the emergent AFM order parameter in phase II. Up to order $O(l^6)$, we find

$$F_{L}(\Psi_{L}) = \alpha_{L}l^{2} + \frac{1}{2}\beta_{L}l^{4} + \frac{1}{3}\gamma_{L}l^{6}\cos 6\theta.$$
(H5)

We emphasize that α_3 does not need to change the sign for the AFM component to become nonzero. Having two order parameters with independent transition temperatures requires more fine-tuning of the model parameters. Here, instead, we argue that $\alpha_3 > 0$ and that the nonzero Ψ_L is induced by Ψ_I through the coupling term $F_{AL}(\Psi_I, \Psi_L)$, as we show in the next section. Note that γ_3 enforces the moments to point along high-symmetry directions of the lattice. Similar to what we discussed above, we set this term to zero to capture the fact that the moment direction is selected locally by the strain.

3. The form of F_{AL}

The last term in Eq. (H1) includes both linear and quadratic couplings between Ψ_I and Ψ_L :

$$F_{AL} = \delta_1 F_{AL}^{(1)} + \delta_2 F_{AL}^{(2a)} + \delta_3 F_{AL}^{(2b)}.$$
 (H6)

We omitted the explicit dependence on Ψ_I and Ψ_L to shorten the notation. In the collinear basis, the coupling that is linear in l is given by

$$F_{AL}^{(1)} = \frac{a_1 l}{\sqrt{2}} [(a_1^2 - 2a_2^2)\cos(\theta - \phi_1) - a_2^2\cos(\theta + \phi_1 - 2\phi_2)].$$
(H7)

In the presence of a nonzero $F_{AL}^{(1)}$, when Ψ_I becomes nonzero it immediately triggers the development of Ψ_L , and the two transitions happen at the same temperature, except for fine-tuned values of the three angles. We know from the scattering experiments that this is not the case in EuIn₂As₂ where phases I and II are separated in temperature. From Eq. (H7), we note that $F_{AL}^{(1)}$ vanishes identically if $a_1 = 0$. This is an important result as it tells us that the nodal collinear amplitude-modulated phase ($a_2 \neq 0$ and $a_1 = 0$, as well as the symmetry-related structures) is the only candidate magnetic structure for phase I that allows for the development of Ψ_I and Ψ_L at distinct temperatures. This is in agreement with the results of Table I for the structures (expressed in the helical basis) that display a *T*-mod signal but not an *H*-mod signal.

The quadratic couplings, also in the collinear basis, take the form

$$F_{AL}^{2a} = -\frac{l^2}{2} [a_1^2 \cos\left(2(\theta - \phi_1)\right) + a_2^2 \cos\left(2(\theta - \phi_2)\right)], \quad (\text{H8})$$

$$F_{AL}^{2b} = (a_1^2 + a_2^2)l^2.$$
(H9)

These are the terms that can trigger the second phase transition at a temperature $T_{N,2}$ lower than $T_{N,1}$ depending on the signs of the coefficients δ_2 and δ_3 . Indeed, $\delta_3 < 0$ favors a coexistence between a_2 and l, whereas the sign of δ_2 sets the relative angle between θ and ϕ_2 , with $\delta_2 > 0$ favoring collinear directions and $\delta_2 < 0$, orthogonal directions. To summarize, the free-energy analysis gives us the following picture for the phase transitions in EuIn₂As₂: At T_{N1} , a transition into a nodal collinear amplitudemodulated state takes place. As the temperature is further lowered, a_2 increases and through the couplings $F_{AL}^{(2a)}$ and $F_{AL}^{(2b)}$ it can renormalize the coefficient $\alpha_3 > 0$ in F_L , reducing its value. When the renormalized coefficient of l^2 eventually changes sign, a finite Ψ_L develops, triggering the transition to phase II. A nonzero Ψ_L makes F_{AL} also nonzero, and together with the quadratic couplings, also change the form of Ψ_I .

4. Equal-moment condition

The qualitative analysis of the free energy presented above demonstrates that the phenomenology of the magnetic phases of $EuIn_2As_2$ can be captured by the Landau free-energy introduced here, provided that the coefficients satisfy certain conditions. In order to be able to perform a more quantitative analysis, additional information about the Landau coefficients is necessary, otherwise the parameter space is intractably large. Ultimately, these coefficients should be derived from a microscopic model which, as we discussed in the main text, is itself challenging. To achieve some progress, we instead restrict the Landau coefficients such that configurations with equal-amplitude moments are favored by minimization of the free energy (away from the transition).

To motivate our choice of parameters, we start by considering a magnetic structure in phase II with order parameters given by Eqs. (F10) and (F21).

Calculating $|M_{\alpha}(z_i)|$ for this generic structure, we find that the norm of the magnetic moments are layer independent if the following conditions are fulfilled:



FIG. 19. The free-energy landscape given by Eq. (H12), for $A_2 = 1$, $\phi_2 = 0$, $\theta = \pi/2$, and two values of *L*, as noted on the figures. The dashed lines denote the equal spin length condition [Eq. (H10)] and clearly coincide with the minima of this free-energy term.

$$a_1 = \sqrt{a_2^2 + 2l^2} - \sqrt{2}l, \quad \phi_1 = \phi_2 \pm \frac{\pi}{2}, \quad \theta = \phi_2 \mp \frac{\pi}{2}.$$
(H10)

The form of Eq. (H10) suggests that for angles $\phi_1 = \phi_2 \pm \pi/2$ and $\theta = \phi_2 \mp \pi/2$, the equal-moment condition would be favored by a term of the Landau functional of the form

$$F_{N0} = \left((a_1 + \sqrt{2}l)^2 - (a_2^2 + 2l^2) \right)^2.$$
(H11)

We find by inspection that such a term is allowed by the symmetries of the paramagnetic group and, in fact, can be obtained directly from the action by constraining the Landau coefficients. In particular, by choosing $\delta_2 = \delta_1$, $\delta_3 = -(\delta_1/2)$ in F_{AL} , $\alpha_2 = -(\delta_1/2)$ in F_A , and enforcing the relationships between ϕ_1 , ϕ_2 , and θ given in Eq. (H10), we reproduce a term proportional to F_{N0} . This motivates us to keep the angles ϕ_1 , ϕ_2 , and θ arbitrary and enforce the same relationship between the parameters δ_1 , δ_2 , δ_3 , and α_2 . We obtain

$$F_{N} = -\frac{\delta_{1}}{8} \{a_{1}^{4} + a_{2}^{4} + 4a_{1}^{2}l^{2}\cos[2(\theta - \phi_{1})]\} + 4\sqrt{2}a_{1}l[-(a_{1}^{2} - 2a_{2}^{2})\cos(\theta - \phi_{1}) + a_{2}^{2}\cos(\theta + \phi_{1} - 2\phi_{2})] + 4l^{2}\{a_{1}^{2} + a_{2}^{2} + a_{2}^{2}\cos[2(\theta - \phi_{2})]\} + 2a_{1}^{2}a_{2}^{2}\cos[2(\phi_{1} - \phi_{2})].$$
(H12)

It turns out that F_N is minimized when condition (H10) is met. To show that, in Fig. 19 we plot F_N as a function of a_1 and ϕ_1 , for two different values of l. In both cases, we find minima of F_N when the equal-moment condition is satisfied.

Thus, to perform the quantitative analysis shown in Fig. 9 of the main text, we minimize the following free energy:

$$\begin{split} \tilde{F} &= (T - T_{N1})a^2 + \frac{\beta_1}{2}a^4 + \frac{\beta_4}{2}a^6 - \frac{\delta_1}{8}(a_1^4 + 2a_1^2a_2^2\cos[2(\phi_1 - \phi_2)] + a_2^4) \\ &+ \frac{\gamma_2}{16}(a_1^2 - a_2^2)[(a_1^2 - a_2^2)^2 - 12a_1^2a_2^2\cos(\phi_1 - \phi_2)^2] + \alpha_3l^2 + \frac{1}{2}\beta_3l^4 - \frac{1}{2}\delta_1\{a_1^2l^2\cos(2(\theta - \phi_1)) + \sqrt{2}a_1l[-(a_1^2 - 2a_2^2)\cos(\theta - \phi_1) + a_2^2\cos(\theta + \phi_1 - 2\phi_2)] + l^2[a^2 + a_2^2\cos(2(\theta - \phi_2))]\}, \end{split}$$
(H13)

where we introduced the notation $a^2 = a_1^2 + a_2^2$.

For the results shown in Fig. 9, we introduced a temperature dependence for δ_1 , such that $\delta_1 = \chi_0 - \chi_1(T_{N1} - T)\Theta(T_{N1} - T)$, with $\chi_1 = 0.075$ and $\chi_0 = 0.05$. Here, $\Theta(x)$ is the Heaviside step function, which is 0 for $x \le 0$ and 1 for x > 0. Such a temperature-dependent term plays a similar role to higher-order terms in the free energy that are only relevant farther from the transition, since the order parameter scales as $a^2 \sim (T_{N1} - T)\Theta(T_{N1} - T)$.

At T_{N1} , the amplitude-modulated phase is favored by $\alpha_2 < 0$. Note that this condition is satisfied by our parameters, since we set $\alpha_2 = -\delta_1/2$ and $\delta_1 = \chi_0 > 0$ at T_{N1} . As the temperature is lowered, the assumption is that higher-order terms (i.e., beyond sixth order) effectively renormalize δ_1 and make it switch sign below T_{N1} , which favors the equal-moment condition.

The other parameters were set to $\alpha_3 = 0.025$, $\beta_1 = 0.5$, $\beta_3 = 0.01$, $\beta_4 = 0.2$, $\gamma_2 = 0.1$. Recall that, as explained above, β_2 , γ_1 , and γ_3 were set to zero to model the dominant role played by the local strain in setting the moments direction as compared to the intrinsic crystalline anisotropy.

The starting point for energy minimization in Fig. 9(a) at the lowest temperature was a series of random moment configurations. Since all relative orientations of the moments and the lattice are degenerate when setting $\beta_2 = \gamma_1 = \gamma_3 = 0$, the moment orientation in the optimized structure was determined by the choice of the random initial conditions (and could be changed by a change of the random number generator). At every temperature *T*, the structure from the last temperature step (T - dT) was used as the initial condition, mimicking the experimental evolution of the structure.

APPENDIX I: SPIN HAMILTONIAN

1. Heisenberg term

In the following, we construct the exchange Hamiltonian that captures the tendency toward the order at two wave vectors, as observed in $EuIn_2As_2$. Let us suppose the Heisenberg exchange between *n*th neighbors is

$$\mathcal{J}_{n}(J_{0}, k, n, d) = J_{0} \frac{\cos(k(n-1))}{(kn)^{d}},$$
 (I1)

where J_0 corresponds to nearest-neighbor exchange. For d = 0, the interaction does not decay at all and is represented by a function in k space. d = 1, 2, 3 correspond

to leading distance-dependent terms of the RKKY interaction in one, two, and three dimensions [62]. Let us assume a single-q structure of the form

$$\mathbf{S}_{\mathbf{n}} = S_q(\cos{(\pi q n)}\hat{\mathbf{x}} + \sin{(\pi q n)}\hat{\mathbf{y}}), \quad (I2)$$

and calculate the total energy of such a state, assuming Heisenberg interactions, with the distance-dependent exchange given by Eq. (I1). The total exchange energy is

$$E_{\rm ex} = J_0 S_q^2 \sum_{n=1}^N \frac{\cos{(k(n-1))}}{(kn)^d} \cos{nq}, \qquad (I3)$$

where N corresponds to the number of neighbors considered in the interaction.

In Fig. 20, we plot the exchange energy [Eq. (I3)] for $k = \pi/3$, J = -1, N = 100, and $S_q = 1$ as a function of q, for d = 0, 1, 2, 3. It is interesting to see that only for d = 0 and d = 1 the exchange energy is minimized for a $q \neq 0$; in other cases, the interaction drops off too quickly to overcome the nearest-neighbor ferromagnetic tendency.



FIG. 20. The exchange energy [Eq. (I3)] for $k = \pi/3$, J = -1, N = 100, and $S_q = 1$ as a function of q, for d = 0, 1, 2, 3. The dashed line marks Q_1 .



FIG. 21. Exchange energy of a state with periodicity q if the bilinear terms are given by Eq. (I4) for (a) $J_1 = -0.1$, $k_1 = \pi/3, J_2 = 0.12, k_2 = \pi, N = 100$; (b) $J_1 = -0.1, k_1 = 0.4\pi$, $J_2 = 0.125, k_2 = \pi, N = 10$.

We now have all the tools needed to choose the distancedependent exchange interaction, which will favor both the Q_2 and the Q_1 order. Our Heisenberg term will be constructed as a superposition of terms favoring the Q_2 and Q_1 orders:

$$\mathcal{J}_n = J_1 \frac{\cos\left(k_1(n-1)\right)}{k_1 n} + J_2 \frac{\cos\left(k_2(n-1)\right)}{k_2 n}, \quad (\text{I4})$$

where $k_1 = \pi/3$ and $k_2 = \pi$. In Figs. 21(a) and 21(b), we plot the total exchange energy as a function of q for two sets of parameters. In Fig. 21(a), we show the exchange energy for N = 100 neighbors, showing that the minimum is indeed found at $q_z = 1/3$ for $k_1 = \pi/3$ in the limit of a large number of neighbors. Since it is not practical for a real calculation to include 100 neighbors, in Fig. 21(b) we modify the parameters to obtain the two minima at q =1, 1/3 for N = 10. By controlling J_2/J_1 , we can choose the relative depth of the two energy minima, and therefore pick the lowest energy q; for the rest of our modeling, we will use $J_2/J_1 = -1.25$, which ensures that the lowest-energy states are found for Q_1 , but also that the Q_2 state is close to it in energy.

To summarize, we find that a long-range RKKY-like exchange is needed to capture the susceptibility toward order at two values of q. However, the result of energy minimization of such a model is always a single-q state: The deepest minimum in Fig. 21 is chosen.

2. Atomistic simulations

Atomistic simulations shown in Fig. 12 were performed with the SPIRIT code [50]. For simplicity, we simulated a cubic lattice with $6 \times 6 \times 12$ sites. The Hamiltonian used for simulations was

$$H = \sum_{ij} \mathcal{J}_{ij} \mathbf{S}_i \cdot \mathbf{S}_j + \sum_{ijkl} \mathcal{J}_{ijkl} (\mathbf{S}_i \cdot \mathbf{S}_j) (\mathbf{S}_k \cdot \mathbf{S}_l), \quad (\mathbf{I5})$$

TABLE III. Heisenberg terms used for atomistic simulations. da, db, and dc denote the spin-spin distances along the three orthogonal directions in units of lattice length. All exchanges are given in meV. The signs in the table are consistent with the SPIRIT convention [50].

da	db	dc	$-{\cal J}_{ij}$
1	0	0	1
0	1	0	1
0	0	1	0.25
0	0	2	0.202 254
0	0	3	-0.218 169
0	0	4	-0.038 627 1
0	0	5	-0.019 098 3
0	0	6	0.125
0	0	7	-0.013 641 6
0	0	8	-0.0193136
0	0	9	-0.0727232
0	0	10	-0.040 450 8

TABLE IV. Fourth-order exchange terms used for atomistic simulations. da_j , db_j , and dc_j denote the distances between spins *i* and *j* along the three orthogonal directions in units of lattice length. All exchanges are given in meV. The signs in the table are consistent with the SPIRIT convention [50].

da_j	db_j	dc_j	da_k	db_k	dc_k	da_l	db_l	dc_l	$-\mathcal{J}_{ijkl}$
0	0	1	0	0	0	0	0	1	-0.2
0	0	1	0	0	1	0	0	2	-0.2

TABLE V. LLG parameters.

Parameter	Value
llg_seed llg_n_iter llg_n_iter llg_dampi llg_beta llg_dt	20 006 ations 5 000 000 ations_log 5000 ing 0.3 0.1 1 0 <i>E</i> -5
llg_force_	convergence 10 <i>e</i> -9

with the Heisenberg parameters given in Table III and the fourth-order terms in Table IV. All other parameters needed to reproduce the simulations are given in Table V.

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