Origin of Incommensurate Magnetic Order in the RAlSi Magnetic Weyl Semimetals (R = Pr, Nd, Sm)

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We investigate rare-earth magnetic Weyl semimetals through first-principles simulations, analyzing the connection between incommensurate magnetic order and the presence of Weyl nodes in the electronic band structure. Focusing on PrAlSi, NdAlSi, and SmAlSi, we demonstrate that the reported helical ordering does not originate from the nesting of topological features at the Fermi Surface or the Dzyaloshinskii-Moriya interaction. Instead, the helical order arises from frustrated isotropic short-range superexchange between the 4f moments facilitated by pd-hybridization with the main group elements. Employing a spin Hamiltonian with isotropic exchange and single-ion anisotropy we replicate the experimentally observed helical modulation.

Magnetic Weyl semimetals form an exciting class of topological materials [1], owing to the possibility they offer of combining nontrivial topology in both reciprocal momentum space and the space of magnetic order parameters. This raises the prospect of identifying new topological invariants which characterize novel and intriguing physical response properties. In momentum space, Weyl nodes emerge as topologically nontrivial points of contact or crossings between two Kramers' degeneracy-lifted bands, acting as sinks and sources of diverging Berry curvature [2, 3]. These correspond to magnetic monopoles strongly affecting electronic response and transport properties, such as anomalous Hall [4] or Nernst effects, magnetoresistance [5, 6] or optical properties [7], if they are in the vicinity of the Fermi surface (FS). These emergent Weyl fermions are either a consequence of time-reversal symmetry or spatial-inversion symmetry breaking (ISB) in presence of the spin-orbit interaction (SOI) [8].

Weyl materials are of great interest magnetically owing to the complex magnetic textures such as spin-spirals they can exhibit which can be turned into topological nontrivial textures such as skyrmions with the application of external magnetic fields. ISB, SOI, and magnetism are also the necessary ingredients for chiral Dzyaloshinskii-Moriya magnetic interactions (DMI) [9, 10]. DMI can compete with exchange interactions, giving rise to chiral spin-spiral ground states [11] and chiral magnetic skyrmions [12] when magnetic fields are applied. This hints at a relation between the Weyl points and skyrmion formation.

All the more surprising is the recent observation of spiral magnetism in magnetic rare-earth (R) RAlSi Weyl semimetals and an apparent link to nesting between topologically nontrivial Fermi surface pockets [13–18]. This suggests that the Weyl points are directly related to the much stronger exchange interaction, rather than the weaker DMI, and are the determinants for the magnetic phases. A correlation between band structure topology,

i.e. the Weyl points, and the emergence of an incommensurate magnetic order has been found for NdAlSi [13]. The observed helical magnetic order is characterized by a wave vector q that matches the vector connecting the topological features observed in the FS. The incommensurate order in NdAlSi transforms to a commensurate ferrimagnetic one [13] at low temperatures, attributed to the magnetic anisotropy originating from crystal field effects [19]. For SmAlSi [14] helical magnetic order has been identified to persist to lower temperatures and include a topological Hall effect characteristic of the Aphase [20] in skyrmion materials. PrAlSi exhibits both ferromagnetic and also possibly spin glass or ferromagnetic cluster glass behavior [15]. Thus, RAISi compounds have garnered significant interest to be ideal systems for the exploration of the interplay between nontrivial valence band structures and chiral magnetic textures.

The relation between the FS nesting vectors and the helical incommensurate order is suggestive, this assumption is valid in presence of an indirect exchange or Ruderman-Kittel-Kasuya-Yosida (RKKY) interactions [21–23]. In general, several exchange mechanisms are present in complex magnetic materials, including superexchange (SX) [24, 25] and double exchange [26]. A quantitative evidence including all the aforementioned exchange mechanisms which shows a correlation between the topological features at the FS and the magnetic order in the RAlSi family is still missing.

In this letter, for three specific rare-earth semi-metallic candidates, PrAlSi, NdAlSi, and SmAlSi, we explore to what extent the RKKY interactions and the topology of the threadbare FSs can contribute to the materials' magnetic orders. Moreover, we also assess whether short-ranged but frustrated SX interactions are preeminent owing to the presence of a quasi-band gap. To this end, we examine the magnetic interactions between the rare-earth atoms employing a first-principles approach, which is unbiased as to the exchange mechanism.

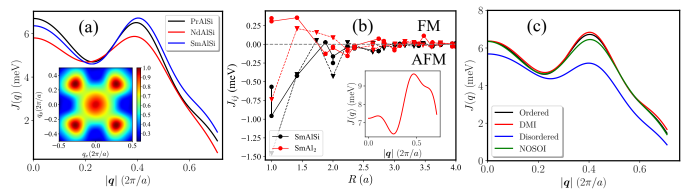


FIG. 1. (a) The maximal eigenvalues J(q), indicating a competition between ferromagnetic (q = 0) and helical order at q_p in [110] direction, for PrAlSi, NdAlSi, and SmAlSi. The inset highlights the 4-fold degenerate peak for SmAlSi. (b) Real-space isotropic exchange interactions in SmAlSi and SmAl₂ between 4f atoms at distance R, showing different regimes of SX and RKKY. The full (dotted) lines represent intra-layer (inter-layer) interactions, the inset depicts the J(q) for SmAl₂. (c) A comparison of J(q) for different approximations: SmAlSi compound with ordered Si and Al atoms (black), including the DMI (red), without SOI (green), and with a disordered Si and Al distribution, restoring centrosymmetry, with SOI (blue).

In order to extract some generic insights about the magnetic interactions prevalent in these 4f magnetic Weyl semi-metal materials, we study a GdAlSi prototype (see Supplementary Material [27]) whose lattice parameters are set to match each RAlSi material [21, 22]. This enables us to discriminate between RKKY-like magnetic interaction mechanisms, reliant on electronic structure near the Fermi energy in metals, or SX-like mechanisms inherent in magnetic insulators. We find generic competing ferromagnetic and incommensurate helical order interactions. We then perform ab initio crystal field theory calculations of single-ion anisotropies [28] to determine the specific magnetic ordered structures.

We employ Hubbard U-corrected density functional theory (DFT+U) [29] calculations to investigate the magnetic interactions and electronic band structures. The calculations are performed with the all-electron full-potential Korringa-Kohn-Rostoker (KKR) Green function method [30], including scalar relativistic effects and spin-orbit coupling self-consistently [31]. We compute the magnetic interactions between the 4f magnetic atoms using the infinitesimal rotation method [32–34]. The real space exchange interactions tensor and the corresponding lattice Fourier transforms are carefully inspected, unveiling the different exchange mechanisms at play. The lattice constants employed in the calculations are taken from experiment (Supplementary Material [27]).

A minimal spin Hamiltonian, \mathcal{H}_{m} , for a non-centro-symmetric crystal, with the magnetic interactions, which we have calculated *ab initio*, is given by:

$$\mathcal{H}_{m} = -\frac{1}{2} \sum_{i \neq j} J_{ij} \, \boldsymbol{m}_{i} \cdot \boldsymbol{m}_{j} + \frac{1}{2} \sum_{i \neq j} \boldsymbol{D}_{ij} \cdot (\boldsymbol{m}_{i} \times \boldsymbol{m}_{j})$$

$$- \sum_{i} K_{i} \, (\boldsymbol{e}_{n} \cdot \boldsymbol{m}_{i})^{2} \quad .$$

$$(1)$$

 m_i is the direction of the magnetic moment at a R-site i, and e_n is the direction of the (effective) easy axis speci-

fied with respect to the crystal structure. The first term refers to the isotropic Heisenberg interactions, the second the anti-symmetric Dzyaloshinskii-Moriya interactions (DMI) [9, 10], which promote chiral spin textures [35], and the third the crystal field single-ion anisotropy at the rare-earth sites (K_i) . We can safely neglect the twoion anisotropy due to its small size compared to K_i . In order to find the origin of the helical order observed in the RAISi family, we inspect the Fourier transform $\mathcal{J}_{mn}^{\alpha\beta}(q)$ of the magnetic exchange interactions. m and n denote the atomic indices in the unit cell labelling the two magnetic atoms in RAlSi, α and β indicate the $\{x, y, z\}$ components. The maximal eigenvalue J(q) of the Fourier transform matrix $\mathcal{J}_{mn}^{\alpha\beta}(\boldsymbol{q})$ provides information on the magnetic order and an estimate of the transition temperature T_N . J(q) is depicted in Fig. 1(a) for our Gd prototype with the lattice structures of PrAlSi, NdAlSi and SmAlSi, respectively (supplemental note 3). In each case, we see two peaks of roughly comparable magnitude at q = 0, indicative of intra-layer ferromagnetic correlations, and at $q = \frac{2\pi}{a}(q,q,0)$ with $q \approx 0.3$ (a being the in-plane lattice constant), which describes single-qspiroidal magnetic correlations [22, 23]. The dominant peak determines the magnetic order that will form below the transition temperature.

For PrAlSi, the maximum peak occurs at $\mathbf{q}_p = (0,0,0)$ and the ferromagnetic order within each layer is favored as observed experimentally in Ref. [36], while for the NdAlSi case, the maximum peak occurs at $\mathbf{q}_p = \frac{2\pi}{a}(q_p,q_p,0)$ with $q_p = 0.273$ in agreement with the experimental value of $q_p^{\text{exp}} = \frac{1}{3} + \delta$ [13]. However, the small energy difference $\Delta E = 0.07$ meV between the spiral and ferromagnetic states, $J(\mathbf{q}_p) - J(\mathbf{0})$, is not sufficient to overcome the magnetic anisotropy energy caused by the crystal field, suppressing the helical ordering at low temperature and enforcing a ferromagnetic order intralayer [13]. For the case of SmAlSi, the maximum peak occurs once more at a finite-q with $\mathbf{q}_p = \frac{2\pi}{a}(q_p, q_p, 0)$

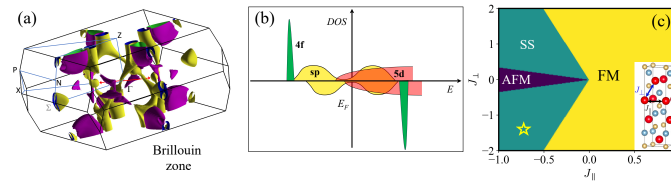


FIG. 2. (a) The complex FS of semi-metallic SmAlSi, with the high-symmetry points and Brillouin zone indicated. The red arrow indicates the non-trivial nesting vector at the FS discussed in Ref. [14]. (b) Schematic representation of the low semimetallic density of states (DOS) at $E_{\rm F}$ displaying the sp-band and the hybridization with spin-polarized d-electrons. (c) A simplified two-parameter phase diagram incorporating nearest neighbor intra-layer (J_{\parallel}) and inter-layer (J_{\perp}) interactions (see inset in Fig. 2c), illustrating the emergence of spin-spiral (SS) order within each rare-earth layer under antiferromagnetic $(J_{\parallel} < 0)$ intra-layer exchange interactions. Phases with (anti)ferromagnetic order within each layer are labeled with FM (AFM). The yellow star indicates the value of the nearest neighbors' exchange interactions calculated ab initio.

and $q_p=0.283$, very close to the experimental value $q_p^{\rm exp}=0.33$ [14]. Compared to the NdAlSi case, the energy difference here is much larger, $\Delta E=0.36$ meV, and together with the reduction of the crystal field effects owing to the lanthanide lattice contraction of heavier 4f elements (see discussion below), this results in a persisting incommensurate order at low temperatures [14]. At these temperatures quantum effects play a role in determining the transition temperatures. Nonetheless, estimating $T_{\rm N}$ using a mean-field, classical spin prescription, $T_{\rm N}=J(q_p)/3k_{\rm B}$, with $k_{\rm B}$ being the Boltzmann constant, we find $T_{\rm N}\simeq 25$ K, which is of the same order of magnitude as the experimentally measured $T_{\rm N}$ [14].

To determine the dominant exchange contribution in RAlSi, we examine Fig. 1(b), where the black curves represent the isotropic exchange interactions J_{ij} as functions of inter-atomic distance. The full (dotted) line represents short-range, antiferromagnetic intra-layer (interlayer) interactions indicating a SX mechanism [24] over a weak RKKY exchange. Despite the coexistence with RKKY-like interactions, the scarcity of electronic states near $E_{\rm F}$ and the threadbare FS (Fig. 2(a)) favor the dominance of SX interactions.

Figure 2(a) displays the FS of the Gd-prototype (with the SmAlSi lattice constant) obtained using the FLEUR code [37] (see Supplementary Material [27]). The FS occupies only a small portion of the Brillouin zone owing to the low density of states near the Fermi energy $(E_{\rm F})$ i.e. the semi-metallic nature of SmAlSi. The suggested nesting between the non-trivial Fermi pockets is indicated by a red arrow in Fig. 2(a). However, the FS sheets are not parallel and exhibit a three-dimensional dispersion, which does not fulfill the nesting condition required to stabilize incommensurate ordering. This observation aligns with recent findings in Ref. [38], where the computation of the Lindhard susceptibility, based on the FS, does not indicate a finite-q helical ordering.

To further elaborate on this balance between RKKY

and SX interactions, we analyze a related but metallic compound GdAl₂ with the same lattice constants (SmAlSi). Substituting Si with Al removes one electron from the system, shifting $E_{\rm F}$ below the semi-metallic gap (see Supplementary Material [27]). The resulting magnetic interactions are shown in Fig. 1(b). In contrast to GdAlSi, they exhibit an oscillatory long-range behavior, indicating that the RKKY interactions dominates over SX in this metallic regime, favoring a spiroidal state (see inset of Fig. 1(b)). The SX mechanism in RAlSi compounds can be understood through an analysis of the density of states, as depicted schematically in Fig. 2(b). In these compounds, the 4f electrons induce a local Zeeman magnetic field, causing spin polarization of the rare-earth's 5d-electrons. These induced 5d magnetic moments interact with the p electrons at different sites through the non-magnetic Si and Al atoms. The SX interactions can be described in terms of a charge-transfer model similar to the transition-metal oxides one [39]: $J_{ij} \propto -t_{pd}^4/\Delta^3$, where t_{pd} represents the hopping integral between the d and p orbitals, and Δ is the charge-transfer energy. A comparable scenario arises in Gd monopnictides where SX competes with RKKY interactions [40]. Lastly, while maintaining the same crystalline configuration but substituting Gd with Eu (EuAlSi), the R valence 5d electrons are removed (the rare earth atoms are divalent rather than trivalent), leaving only sp-electrons that scatter off the localized 4f electrons. This ultimately leads to an RKKY interaction among the 4f moments (Supplementary Material [27]), once again emphasizing the significant role of 5d-electrons in generating the SX mechanism.

Although our analysis thus far indicates a dominant role for SX, we now investigate the interplay between the Weyl points and the incommensurate helical ordering by considering two distinct cases: firstly excluding SOI, where the band structure contains nodal lines [38, 41], and secondly including SOI which introduces gaps in the

nodal lines, and generates Weyl points at specific symmetry dictated positions in k-space [41]. Fig. 1(c) depicts J(q) of SmAlSi for these two cases - including (black curve) and excluding the SOI (green curve). While J(q) differs slightly near $(q_p,q_p,0)$, the finite-q peak structure remains and incommensurate order is still favored over ferromagnetism with an energy barrier $\Delta E = 0.10$ meV. This demonstrates that a stable helical order can form in the absence of Weyl points. Moreover, despite the presence of Weyl nodes in the electronic band structure of PrAlSi, J(q) has the maximal value at q = (0,0,0) favoring a collinear order even without the inclusion of single-ion anisotropy (see Fig. 1(a)).

We now inspect the role of inversion-symmetry breaking on the magnetic order of SmAlSi. The centrosymmetry can be restored using a 50%-50% alloy of Al and Si species at each site, i.e. $Sm(Al_{0.5}Si_{0.5})_2$. This is achieved computationally using the coherent potential approximation [42]. The electronic band structure obtained for this centrosymmetric SmAlSi alloy is given in the Supplementary Material [27]. Centrosymmetry leads to the removal of the Weyl points (note that the DMI is suppressed as well) and the disorder smears out the electronic bands near the Fermi energy [43]. The resulting J(q), shown in Fig. 1(c) (blue curve), supports ferromagnetic order prevailing in the alloy ($\Delta E = -0.17 \text{ meV}$), but the finite-q peak persists indicating that the helical order does not originate from inversion symmetry breaking or the Wevl points. Lastly, inclusion of DMI (red curve) is found to have a minimal effect owing to its weak magnitude - while it breaks the $\pm q$ reciprocity reducing the four-fold degeneracy (inset Fig. 1(a)), it does not alter the position or the magnitude of the finite-q peak in J(q) which is purely driven by isotropic exchange.

The crystal field effects play an important role in the RAlSi family. The 4f electrons' electronic configuration in RAlSi has an atomic-like behavior in accordance with Hund's rules [14], which in turn determines the shape of the 4f charge cloud. This charge is subjected to the crystal field (CF) originating from the valence electrons and surrounding ions. Considering that the noncollinear order is driven by the isotropic exchange interactions and not crystal fields effects, the tetragonal uniaxial magnetic anisotropy constant K_i is computed from fitting the classical CF energy differences [44] while rotating the 4f moment from the c-axis to the a-axis. The crystal field parameters are obtained within the yttrium analogue approach [28] (see Supplementary Material [27]). Both PrAlSi and NdAlSi display an outof-plane anisotropy, in agreement with the experimental observation [13]: PrAlSi has a high value of $K_i = 2.593$ meV, while the NdAlSi constant is one order of magnitude smaller $K_i = 0.218$ meV. On the other hand, SmAlSi prefers a canted easy anisotropy axis e_n along the $(\theta_n, \phi_n) = (60^\circ, 45^\circ)$ direction with respect to tetragonal basis vectors, with an anisotropy constant of $K_i = 0.13$

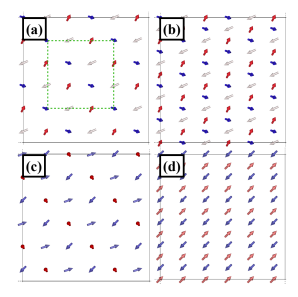


FIG. 3. (a) Helical magnetic structure of SmAlSi in the presence of frustrated isotropic exchange interactions and a canted magnetic anisotropy, the colors indicates the m_z component, blue: $m_z=-1$, red: $m_z=1$, white: $m_z=0$). The green dashed square denotes magnetic unit cell. (b) Same as in (a), including sub-lattice two. (c) Same as in (a) including an external magnetic field (B=4 Tesla) along the easy axis ($60^{\circ},45^{\circ}$). (d) Stacked antiferromagnetic state along the c-axis with higher energy compared to (b).

meV.

We extract the following minimal atomistic spin model in reciprocal space to understand the formation of a helical order in RAlSi from the short-ranged, antiferromagnetic interactions (J < 0)

$$\begin{cases} J_{\rm nn}^{\parallel}(\mathbf{q}) = 2J_{\parallel}(\cos q_x a + \cos q_y a) & , \\ J_{\rm nn}^{\perp}(\mathbf{q}) = 2J_{\perp}(\cos(q_x a/2) + \cos(q_y a/2)) & . \end{cases}$$
 (2)

Hereby, we consider an isotropic Heisenberg model with nearest-neighbor intra-layer (J_{\parallel}) and inter-layer (J_{\perp}) interactions as depicted in the inset of Fig. 2(c). We construct the phase diagram shown in Fig. 2(c) by varying J_{\parallel} and J_{\perp} , and identifying the in-plane ${\bf q}$ which maximizes the eigenvalue $J({\bf q})$. Within each rare-earth layer three phases emerge, namely a ferromagnetic (FM), antiferromagnetic (AFM) and spin spiral (SS) phase. For $J_{\parallel} < 0$, the AFM phase switches to a SS phase as the magnitude of J_{\perp} increases. For SmAlSi, the J_{\parallel} and J_{\perp} parameters lie in the region of the SS phase as can be seen from the first-principles results in Fig. 1(b).

Now, for the case of SmAlSi, considering (short range) magnetic interactions J_{ij} in real space up to 2.5 a, with $K_i = 0.13$ meV, $\boldsymbol{e}_n = (60^\circ, 45^\circ)$, and neglecting the DMI, we minimize the Hamiltonian $\mathcal{H}_{\rm m}$ (1) by solving the Landau-Lifshitz-Gilbert equation as implemented in the Spirit code [45]. The resulting helical order for the first sub-lattice is depicted in Fig. 3(a) (top view), featuring a propagation vector $\boldsymbol{q} = \frac{2\pi}{a}(q,q,0)$ with q = 0.33 consistent with the maximum of $J(\boldsymbol{q})$ for SmAlSi. Fig. 3(b)

illustrates the helimagnetic structure where both sublattices show a similar magnetic order but with a positional shift owing to the stacking along the c-axis. Besides the helical order, a solution slightly higher in energy ($\Delta E = 0.05 \text{ meV}$) features moments antiferromagnetically coupled along the c-axis, as depicted in Fig. 3(d). The helical order can transform to this antiferromagnetic state under high-field conditions or with thermal fluctuations. To explore the emergence of non-collinear spin textures, we apply a magnetic field along the easy-axis [23] e_n . The resulting state is shown in Fig. 3(c). We observe a canting of the moments towards the +z direction, but no skyrmion lattice phases can form owing to the short period of the magnetic structure (3a). The helical order remains the most stable one.

Coming back to the above mentioned non-conventional contribution to the Hall signal observed experimentally for SmAlSi when a magnetic field is applied [14], we conjecture that this signal can be interpreted in terms of the recently introduced non-collinear Hall effect [46]. This effect emerges from the interference between non-collinear magnetism and spin-orbit interactions in a non-centrosymmetric environment, without invoking the presence of magnetic skyrmions. Likely, this unconventional Hall signal originates from the scattering of Weyl fermions on the helical magnetic background.

In summary, we have examined three members from the RAlSi family, recently proposed as materials where Weyl-mediated RKKY interactions generate helical ordering. Our findings demonstrate that neither the Weyl points nor the RKKY interactions provide the predominant mechanism owing to the presence of a low density of states near the Fermi energy in these semi-metals. Instead, we find a significant p-d antiferromagnetic SX contribution. The competition of these isotropic SX interactions between different atoms leads to a helical order with a short period of approximately three lattice constants, all without the assistance of DMI. Our ab initio calculations reveal a strong magnetic anisotropy arising from crystal field effects for PrAlSi and NdAlSi, locking the moments into a collinear configuration. In contrast, the magnetic anisotropy in SmAlSi is much lower, which allows the formation of a helical incommensurate order at low temperatures. Lastly, the short period of the spin spiral does not permit the emergence of a skyrmion lattice when external magnetic fields are applied.

For future prospects, our *ab initio* computed crystal field coefficients serve as a theoretical input for ongoing experimental investigations using inelastic neutron scattering to understand the complex magnetic structures and crystal field excitations of the RAlSi compounds. Meanwhile, our analysis of the Fermi surface and electronic band structure in different configurations, including chemically ordered, paramagnetic, and chemically disordered, provides very useful insights to interpret experimental photo-emission data [38]. Finally, alloying Al

and Si with other elements from the same family or applying strain may drive the exchange parameters of the compound closer to the SS-FM phase boundaries of the phase diagram (Fig. 2(b)), leading to longer-period helical orders which may be turned into a skyrmion lattice by applying an external magnetic field. This provides an ideal platform to study the interplay between the topology of the magnetic textures in real space and that of Weyl fermions in reciprocal space.

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