

Anisotropic fractal magnetic domain pattern in bulk Mn 1.4 Pt Sn

A. S Sukhanov, B. E Zuniga Cespedes, P. Vir, A. S Cameron, A. Heinemann, N. Martin, G. Chaboussant, V. Kumar, P. Milde, L. M Eng, et al.

► To cite this version:

A. S Sukhanov, B. E Zuniga Cespedes, P. Vir, A. S Cameron, A. Heinemann, et al.. Anisotropic fractal magnetic domain pattern in bulk Mn 1.4 Pt Sn. Physical Review B, 2020, 102 (17), 10.1103/Phys-RevB.102.174447 . hal-03092082

HAL Id: hal-03092082 https://hal.science/hal-03092082

Submitted on 1 Jan 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Anisotropic fractal magnetic domain pattern in bulk Mn_{1.4}PtSn

A. S. Sukhanov,^{1,2} B. E. Zuniga Cespedes,^{1,3} P. Vir,¹ A. S. Cameron,² A. Heinemann,⁴ N. Martin⁰,⁵ G. Chaboussant,⁵

V. Kumar[®],⁶ P. Milde[®],³ L. M. Eng[®],^{3,7} C. Felser,^{1,7} and D. S. Inosov[®],^{2,7}

¹Max Planck Institute for Chemical Physics of Solids, D-01187 Dresden, Germany

²Institut für Festkörper- und Materialphysik, Technische Universität Dresden, D-01069 Dresden, Germany

³Institute of Applied Physics, Technische Universität Dresden, D-01069 Dresden, Germany

⁴German Engineering Materials Science Centre at Heinz Maier-Leibnitz Zentrum,

Helmholtz-Zentrum Geesthacht GmbH, D-85747 Garching, Germany

⁵Université Paris-Saclay, CEA, CNRS, Laboratoire Léon Brillouin, CEA Saclay, F-91191 Gif-sur-Yvette, France

⁶Physik-Department, Technische Universität München, D-85748 Garching, Germany

⁷Dresden-Würzburg Cluster of Excellence on Complexity and Topology in Quantum Matter (ct.qmat),

TU Dresden, D-01062 Dresden, Germany

(Received 18 June 2020; revised 5 November 2020; accepted 5 November 2020; published 30 November 2020)

The tetragonal compound $Mn_{1.4}$ PtSn with D_{2d} symmetry recently attracted attention as the first known material that hosts magnetic antiskyrmions, which differ from the skyrmions known so far by their internal structure. The latter have been found in a number of magnets with the chiral crystal structure. In previous works, the existence of antiskyrmions in Mn_{1.4}PtSn was unambiguously demonstrated in real space by means of Lorentz transmission electron microscopy on thin-plate samples (~ 100 nm thick). In the present study, we used small-angle neutron scattering and magnetic force microscopy to perform reciprocal- and real-space imaging of the magnetic texture of bulk $Mn_{1.4}$ PtSn single crystals at different temperatures and in applied magnetic field. We found that the magnetic texture in the bulk differs significantly from that of thin-plate samples. Instead of spin helices or an antiskyrmion lattice, we observe an anisotropic fractal magnetic pattern of closure domains in zero field above the spin-reorientation transition temperature, which transforms into a set of bubble domains in high field. Below the spin-reorientation transition temperature the strong in-plane anisotropy as well as the fractal self-affinity in zero field is gradually lost, while the formation of bubble domains in high field remains robust. The results of our study highlight the importance of dipole-dipole interactions in thin-plate samples for the stabilization of antiskyrmions and identify criteria which should guide the search for potential (anti)skyrmion host materials. Moreover, they provide consistent interpretations of the previously reported magnetotransport anomalies of the bulk crystals.

DOI: 10.1103/PhysRevB.102.174447

I. INTRODUCTION

Solids exhibiting topological properties are promising for future applications, in particular for spintronics. In the case of magnetic materials with the ferromagnetic spin-spin exchange coupling, the presence of the antisymmetric Dzyaloshinskii-Moriya interaction (DMI) may twist the otherwise homogeneous collinear spin texture into a two-dimensional lattice of densely packed nanometer-sized whirls. Each of these whirls is formed by a spatial distribution of the regularly canted magnetic moments that wrap a whole unit sphere, if mapped out onto it. The mutual noncoplanar orientation of the neighboring spins can be described by the topological charge (or the skyrmion winding number) $N_{\rm sk}$ that takes values ± 1 and differentiates two types of the topologically protected magnetic structures: skyrmions and antiskyrmions [1–6].

While there are an increasing number of the discovered skyrmion-hosting compounds, such as B20-type chiral magnets (e.g., MnSi, FeGe, Fe_xCo_{1-x}Si) [7–12], β -Mn-type Co-Zn-Mn alloys [13–17], Cu₂OSeO₃ [18–23], which support Bloch-type skyrmions, and lacunar spinels (e.g., GaV_4S_8 , GaV₄Se₈) [24–26], which support Néel-type skyrmions, antiskyrmions have been observed only in thin plates of Mn_{1.4}Pt(Pd)Sn [27–30] and Mn₂Rh_{0.95}Ir_{0.05}Sn [31] to date. In contrast to the skyrmion materials with cubic $(P2_13$ and $P4_132$ space groups) or rhombohedral C_{3v} crystal structures (space group R3m), tetragonal Mn_{1.4}PtSn belongs to the D_{2d} symmetry class (space group $I\overline{4}2d$), which is a prerequisite of antiskyrmions [4,27,32,33]. In agreement with the symmetry-based theoretical predictions, the first Lorentz transmission microscopy (LTEM) measurements of Mn_{1.4}PtSn demonstrated nucleation of the triangular lattice of magnetic antiskyrmions in a magnetic field of ~ 0.2 T, applied perpendicular to the surface of a thin lamella sample and

Published by the American Physical Society under the terms of the Creative Commons Attribution 4.0 International license. Further distribution of this work must maintain attribution to the author(s) and the published article's title, journal citation, and DOI. Open access publication funded by the Max Planck Society.

parallel to the [001] crystallographic direction (the tetragonal *c* axis). The antiskyrmions are \sim 200 nm in diameter and were observed in a wide temperature range below $T_{\rm C}$ of \sim 400 K down to $T \approx 150$ K [27,28].

Subsequent LTEM experiments revealed that the antiskyrmions in $Mn_{1,4}$ PtSn can also arrange in a square lattice in some particular temperature and field regions of the phase diagram, which may be affected by the sample thickness [29,30]. Moreover, elliptically distorted skyrmions of both handednesses and the nontopological bubble lattice were shown to appear when a symmetry-breaking in-plane magnetic field is applied in combination with the out-of-plane field [29,30]. This makes $Mn_{1,4}$ PtSn a unique compound hosting a rich variety of controllable topological magnetic objects.

Because the previous studies on thin lamellae of Mn1.4PtSn pointed out that the sample geometry and the sample preparation process can alter some aspects of the material properties [29,30], it is essential to characterize the magnetic structure of the compound in the bulk. Due to a very high penetration depth of the neutron radiation, neutron scattering techniques allow investigations of the magnetic structure of bulk samples. Taking into account the long-periodic modulations of the magnetic texture in Mn_{1.4}PtSn, small-angle neutron scattering (SANS) is a suitable probe, which enables reciprocal-lattice imaging of magnetic structures with periods ranging from ~ 2 to ~ 400 nm [34]. For real-space investigations of the magnetic texture of bulk samples, mostly, surface-sensitive techniques are available. Magnetic force microscopy (MFM) proved to be a valuable tool when studying complex spin textures such as helices and skyrmions as well as complex domain patterns on length scales between ~ 20 nm and $\sim 100 \ \mu m$ and can be applied at various temperatures as well as in external magnetic fields [11,21,24,35].

In the present study, we employ both SANS and MFM to resolve the nanometer-scale magnetic texture of Mn_{1.4}PtSn in the bulk single-crystalline form and observe how it changes when the sample temperature and the applied magnetic field are varied. We discuss the characteristic features of the obtained reciprocal-space and real-space patterns and demonstrate that the magnetic structure of the bulk Mn_{1.4}PtSn differs dramatically from the previously reported LTEM observations obtained using samples in thin-plate geometry. Instead of helices or an antiskyrmion lattice, we observe an anisotropic fractal magnetic pattern of closure domains in zero field above the spin-reorientation transition temperature T_{SR} , with characteristic hints for the DMI inherent in the D_{2d} symmetry of the crystal, which transforms into a set of bubble domains in high field. Below T_{SR} the strong in-plane anisotropy and the fractal self-affinity in zero field are gradually lost, while the formation of bubble domains in high field remains robust. The results of our study highlight the importance of dipoledipole interactions in thin-plate samples for the stabilization of antiskyrmions and identify search criteria for potential (anti)skyrmion host materials.

This paper is organized as follows. After a description of the experimental details, we first discuss the experimental results obtained in zero field and above T_{SR} . Then, we describe the field dependency above T_{SR} . Finally, we discuss the temperature dependence.

II. EXPERIMENTAL DETAILS

Our SANS measurements were performed at the instruments SANS-1 (FRM-II, Garching, Germany) and PA20 (LLB-Orphée, CEA Saclay, France) [36]. In both experiments, we used the same sample, which consisted of 17 crystals coaligned together with a relative misalignment not worse than 3°, mounted on an aluminum plate holder. The crystals were coaligned to increase the total volume of the sample and the resulting experimental signal-to-noise ratio (see the Supplemental Material [37] for a photograph of the sample). All the crystals were grown by the self-flux method, as described in [38,39]. The high quality of the crystals was confirmed by means of magnetic susceptibility, resistivity, and x-ray diffraction measurements. Their stoichiometry was examined by energy-dispersive x-ray spectroscopy, which did not reveal any variation of the chemical composition between the crystals of the same batch or within a surface of each crystal within the precision of this method.

MFM measurements were performed with two instruments. For room-temperature measurements without external fields we used the Park Systems NX10 [40] with PPP-MFMR probes from Nanosensors [41] at lift heights between 100 and 150 nm. Low-temperature measurements with external field were performed in an Omicron cryogenic ultrahigh-vacuum scanning tunneling microscopy/atomic force microscopy instrument [42] using the RHK R9s electronics [43] for scanning and data acquisition. We employed PPP-QMFMR probes from Nanosensors driven at mechanical oscillation amplitudes $A \approx 20$ nm at lift heights between 400 and 800 nm. All data analysis was performed with the GWYDDION [44] software. Two samples have been investigated. Sample A is a single crystal of Mn_{1.4}PtSn, whose native surface was gently polished with a focused ion beam using Xe ions at currents below 10 nA. After the polishing, we checked with MFM that the resulting amorphous surface layer did not alter the domain pattern. Sample B is a polycrystalline sample of Mn_{1.4}Pt_{0.9}Pd_{0.1}Sn that was carefully polished. Measurements of sample A are presented in the main text, while measurements of sample B can be found in the Supplemental Material.

III. RESULTS

A. Magnetic texture at $T > T_{SR}$

Figure 1(a) shows a SANS pattern collected at the sample temperature T = 250 K and zero magnetic field (no prior field history). The sample was oriented with its tetragonal c axis parallel to the incident neutron beam. In this scattering geometry, the reciprocal (HK0) plane is imaged at the position-sensitive detector. The in-plane orientation of the sample corresponds to the momentum component q_x aligned with the [110] direction. Thus, the scattering pattern in Fig. 1(a) represents spin-texture modulations in the ab plane of Mn_{1.4}PtSn. As can be seen, there is a clear scattering intensity distribution that covers almost the whole imaged reciprocal space in the momentum range 0.002 < $Q < 0.017 \,\text{\AA}^{-1}$. The scattering is diffuse and does not have sharp features, such as Bragg peaks one would expect in SANS of helimagnets [7,8,12,13,19,23] for a helical state with a period λ of 37–315 nm. This is in strong contrast to the

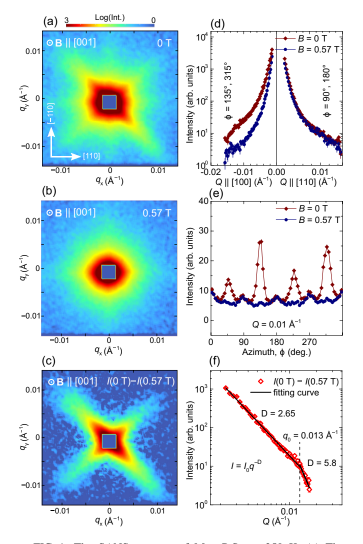


FIG. 1. The SANS patterns of Mn_{1.4}PtSn at 250 K. (a) The SANS pattern at zero field. The sample orientation is depicted by white arrows. (b) The scattering in the field-saturated state at 0.57 T. (c) The result of the subtraction of the SANS map recorded at 0.57 T from the map at 0 T. (d) The comparison of the intensity profiles for the momenta along [100] and [110]. (e) The intensity profiles as a function of the azimuthal angle ϕ . (f) The radial intensity profile of the background-subtracted data (symbols) and the fit by the power function (solid line).

LTEM observations of the helical spin structure with a period of ~ 100 nm in the thin lamellae samples, including at T = 250 K and B = 0 T [27–30], and implies that the magnetic structure of the bulk Mn_{1.4}PtSn is not a spin helix, as was previously anticipated.

The observed SANS pattern is diffuse yet strongly anisotropic and can be viewed as eight streaks (stripes) of intensity, which can be considered a sum of two four-pointed stars (crosses), one of which is higher in intensity and points to the $\langle 100 \rangle$ directions, whereas the other is weaker and oriented with respect to $\langle 110 \rangle$.

It should be noted that a presence of the spiral state with $\lambda > 315 \text{ nm}$ in bulk Mn_{1.4}PtSn would result in the Bragg peaks in SANS that are not resolved within the direct beam in our experimental setup. However, a possibly unresolved

helical state cannot cause the observed diffuse scattering. Moreover, as shown below, the MFM imaging did not reveal the presence of a helical state with $\lambda > 315$ nm, which further supports the conclusions drawn from the SANS measurements.

The origin of this scattering can be understood when a saturating magnetic field is applied parallel to the c axis (i.e., along the neutron beam). Figure 1(b) depicts the SANS pattern at B = 0.57 T, which is well above the saturation field of $\sim 0.5 \text{ T}$ [38]. As one can see, the stripes of intensity along (100) disappeared, but the weaker stripes along (110)remained unchanged, which implies that only the former are of magnetic origin. Hence, the pattern in Fig. 1(b) can be used as a background, and we subtracted the intensity $I(\mathbf{Q}, 0.57 \,\mathrm{T})$ from the pattern $I(\mathbf{Q}, 0 \mathrm{T})$. Both the foreground data and the background data were normalized by the monitor counts for the correct subtraction. The resulting pattern is shown in Fig. 1(c), which represents the pure magnetic intensity. To highlight the difference between the zero-field and the field-saturated states, intensity profiles are plotted for the momentum directions (H00) and (HH0) in Fig. 1(d). Figure 1(e) demonstrates the anisotropy of the intensity distribution cut at $Q = 0.01 \text{ Å}^{-1}$ as a function of the in-plane (azimuthal) angle ϕ . As can be seen from the azimuthal profile, each stripe has a base width of $\sim 45^{\circ}$.

The observation of diffuse anisotropic scattering suggests that Mn_{1.4}PtSn develops a magnetic texture of rectangularshaped domains with domain walls oriented strictly perpendicular to the crystallographic [100] and [010] axes. The well-defined orientation of the magnetic domains follows from the cross-shaped scattering within the reciprocal (HK0)plane. The domains, however, do not feature any regularity either in their sizes or in the domain-wall spacing, which can be concluded from the smooth radial profile of the diffuse scattering. Furthermore, the SANS patterns collected when the sample is oriented by its [110] axis along the incident neutron beam showed the absence of scattering intensity along the reciprocal L direction, which confirms that the spin correlations are homogeneous (ferromagnetic) along the tetragonal c axis (see Fig. S3 in the Supplemental Material [37]).

For further analysis, the radial profile along the stripe of the background-subtracted SANS pattern taken at T = 250 K and B = 0 T is plotted in Fig. 1(f) on a log-log scale. I(q) obeys a power-function trend $I \propto q^{-D}$ with two different exponents below and above the crossover momentum $q_0 = 0.013 \text{ Å}^{-1}$, where the slope changes. The fitting yields D = 2.65 for momenta below q_0 down to the lowest accessible momentum of $\sim 0.002 \text{ Å}^{-1}$, which is a signature of the scattering from fractal objects [45-48]. This implies a complex intertwined arrangement of the domain walls of the rectangular domain pattern. The momentum q_0 then determines the lowest real-space scale down to which the fractal self-affinity holds, which is here \sim 48 nm. The upper limit of the fractal structure cannot be reached within the accessible q range and lies above 315 nm. Above q_0 the exponent D changes to D = 5.8, which can be attributed to scattering from density profiles without sharp (D = 4) contrast (either due to roughness along the profile or a smoothness of the profile) [45,46]. A similar analysis was applied in a SANS study of Nd₂Fe₁₄B [49], where anisotropic

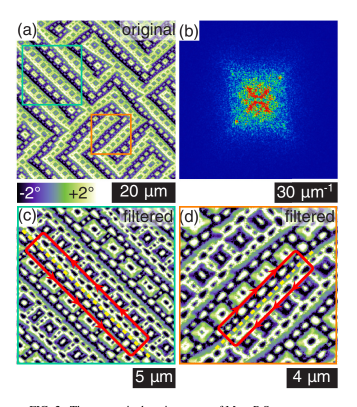


FIG. 2. The magnetic domain pattern of $Mn_{1.4}$ PtSn at room temperature. (a) MFM phase image showing the domain pattern in the *ab* plane at zero field and (b) its fast Fourier transform. (c) and (d) depict the zoomed area marked in (a) by the turquoise and orange squares, respectively. They are filtered for enhanced contrast of the smaller domains. The red loops highlight the chiral sense of the pattern of triangular-shaped nested domains at the domain walls (yellow dashed line) of the lamellar stripe domains. The numbers written in the bars correspond to the length of the bars. Note that the reciprocal-space scale in (b) is much smaller than the momentum range covered in the SANS patterns.

diffuse scattering described by D = 3.7 (D = 3.1) was observed below (above) the spin-reorientation transition.

For comparison, we show a MFM measurement on the ab plane of the Mn_{1.4}PtSn single crystal (sample A) at room temperature in Fig. 2(a). The domain pattern is highly reminiscent of the fractal magnetic closure domains observed in the low-temperature easy-cone anisotropy phase of Nd₂Fe₁₄B [49-51]. It consist of lamellar domains with smaller nested closure domains arranged in a fractal pattern resembling the Sierpinski carpet [52]. The width of the lamellar domains sets the upper boundary of $\sim 3 \mu m$, which is one order of magnitude higher than the upper fractal scale accessed in SANS. The in-plane orientation of the domain walls is highly anisotropic with pinning to two perpendicular directions within the *ab* plane. In the Fourier transform [see Fig. 2(b)], the same cruciform pattern as in SANS is visible. Hence, we can safely assume that our SANS and MFM results describe the same anisotropic fractal domain pattern. Such domain patterns arise from a competition of uniaxial ferromagnetic exchange interaction favoring collinear domains in the easy axis without domain walls and dipolar interactions at the surfaces of the material adding stray-field energy. The

latter is minimized by nucleation of closure domains at the surface at the expense of additional domain walls, whose orientation is defined by the anisotropy within the ab plane. Thus, the nested domains are present in a region below the surface and form a fractal tree structure along the c axis, also known as branching domains [35]. Additional measurements on the polycrystalline sample B confirm that the fractal pattern indeed belongs to such closure domains. Details can be found in the Supplemental Material.

Moreover, we find characteristic hints for the DMI in the material. As already mentioned, lamellar stripe domains appear in two orientations within the *ab* plane. Examples are shown in Figs. 2(c) and 2(d). Note that these two images have been filtered for enhanced contrast of the smaller domains. The nested domains within these stripes partly appear with an arrowhead shape. The direction defined by the arrows along the stripe domain walls (highlighted by yellow dashed lines) defines a certain chirality, which is solely set by the in-plane orientation of the domain wall, as schematically shown in Figs. 2(c) and 2(d) by the red loops. This reflects the D_{2d} symmetry of the crystal; namely, in order to transform from one loop to the other, one has to apply the combination of a 90° rotation and an inversion, like for the chirality of helices and the nontopological bubbles observed by LTEM in thin-plate samples [30]. This very peculiar feature of the domain pattern is so far unique to Mn₁₄PtSn and, to the best of our knowledge, has not been reported before for any other fractal magnetic domain pattern. It may be related to the Dzyaloshinskii-Moriya interaction, which is responsible for the existence of antiskyrmions in the material in the first place [27].

B. Domain structure in applied field

Next, we discuss the magnetic-field response of the magnetic texture of Mn_{1.4}PtSn. Figure 3(a) shows a series of SANS patterns collected at different field magnitudes applied parallel to the c axis at T = 250 K. For all patterns, the B =0.57 T scattering (the fully polarized state) was subtracted as background in analogy to Fig. 1(c). The azimuthal intensity profiles were extracted from each SANS pattern and are plotted in Fig. 3(b). The pattern of B = 0.09 T looks very similar to the SANS pattern at zero field; namely, it has the same anisotropic cruciform scattering distribution with the same intensity. The pattern recorded at B = 0.21 T retains the anisotropy with approximately two times lower overall intensity. The SANS pattern at B = 0.33 T demonstrates very low intensity, which is seen only in the vicinity of the center $Q \approx 0$, where Q denotes the momentum transfer defined as the difference between the momentum of the scattered neutrons \mathbf{k}_{f} and the momentum of the incident neutrons \mathbf{k}_{i} , $\mathbf{Q} = \mathbf{k}_f - \mathbf{k}_i$. Nevertheless, the characteristic cross shape remains quite distinguishable in the pattern. At a higher field of 0.45 T, only a very weak isotropic scattering is observed at small momenta in the vicinity of the direct beam, which indicates a transformation from rectangular towards isotropic domains. Not only is the symmetry of the diffuse scattering preserved in increased magnetic field up to B = 0.33 T, but also the radial I(Q) profiles approximately retain the initial slope, as evidenced in Fig. 3(c), where the profiles are shown

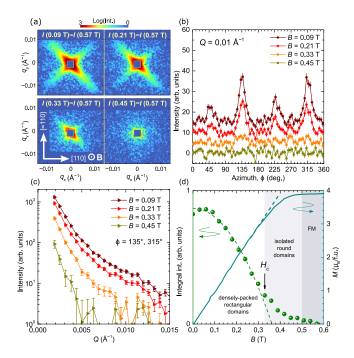


FIG. 3. Magnetic field dependence of the SANS signal at T = 250 K. (a) A series of SANS patterns (after background subtraction) collected in different magnetic fields applied along the *c* axis. (b) The azimuthal profiles of the scattering extracted from each pattern. (c) The radial intensity profiles at the same fields. (d) The integral intensity of the magnetic scattering as a function of the applied field (symbols) and the magnetization M(H) curve measured at the same temperature and field orientation. The dashed lines are the approximations by parabolic and linear functions (see the text). The black arrow marks the crossover between the two field-polarizing regimes.

for the same set of fields. The data collected at higher fields up to 1.4 T did not reveal any scattering, which confirms that the sample is uniformly magnetized already at 0.57 T.

There are two possible scenarios of the magnetic-structure polarization process that would cause the observed smooth decrease in the intensity of the anisotropic diffuse scattering. The first one implies that in the applied magnetic field the rectangular-domain texture is gradually dissolved into the homogeneous ferromagnetic background. In other words, the sample breaks into coexisting domains of the fully polarized state, growing in volume with an increasing field, and the volume occupied by the densely packed small rectangular domains. In this case, the intensity of the SANS should decrease in accord with the modulated-texture volume, $I = I_0(1 - M_z/M_s)$, where M_z/M_s is the normalized net magnetization and I_0 is the intensity in zero field. In the opposite scenario, the modulated magnetic texture occupies the entire volume in finite applied fields, but the magnitude of the modulated component of the local magnetization (its in-plane projection) is reduced in favor of the homogeneous M_{τ} component. Since the SANS intensity $I \propto M_{xy}(x, y)^2$, where $M_{x,y}(x, y)$ is the magnetization component modulated in the basal plane, the field dependence of I should read $I = I_0 [1 - (M_z / M_s)^2].$

The integral intensities (integrated along the stripes) of the diffuse scattering were extracted from the SANS patterns,

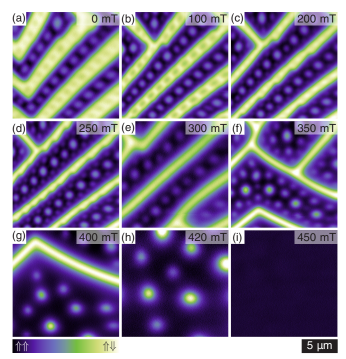


FIG. 4. Magnetic field dependence of the domain pattern in MFM at T = 240 K. Sample magnetization parallel and antiparallel to the external field is shown in purple and green, respectively. The frame size is equal for all images; the span of the color scale has been adapted individually.

which were measured in applied fields of different magnitudes, and are plotted in Fig. 3(d) (symbols) along with the isothermal magnetization curve (solid line) obtained with a superconducting quantum interference device magnetometer. The magnetization demonstrates a linear dependence up to $B \approx 0.33$ T, where it reaches $\sim 3/4$ of the saturated moment $M_{\rm s} = 3.9 \mu_{\rm B}$. Notably, the intensity I(B) can be well approximated by a parabolic function in the same field range, which agrees with the second scenario. Above B = 0.33 T, the intensity starts deviating from the quadratic field dependence and switches to the $1 - M_z/M_s$ behavior, as predicted by the first scenario, until it vanishes at a saturating field of ~ 0.5 T. The field $B_{\rm c} = 0.33 \,{\rm T}$ can therefore be denoted as the crossover point at which the partially polarized rectangular domain texture becomes diluted by the regions of the fully polarized state or isotropic domains.

In Fig. 4, we show a series of MFM measurements obtained at T = 240 K on the *ab* plane of the Mn_{1.4}PtSn single crystal with magnetic field applied parallel to the *c* axis, i.e., perpendicular to the plane of view. Areas with sample magnetization pointing parallel (antiparallel) to the applied field are shown in purple (green). In the region between zero field and B = 100 mT there are no qualitative changes in the domain pattern [see Figs. 4(a) and 4(b)], which agrees with only a small change in SANS intensity in this field range. With increasing field up to B = 200 mT, shown in Fig. 4(c), the nested domains within the antiparallel magnetized stripe domains (shown in green) disappear, yet the edges of those obey a sawtooth shape. In turn, in the stripe domains magnetized parallel to the field, the number of nested domains increases. Between B = 250 and 350 mT, the sawtooth shape of the domain walls is lost, and the nested domains appear with both a more rounded shape and less ordering, shown in Figs. 4(d) to 4(f). For further increasing field magnitudes, the antiparallel magnetized domains start to disappear, and only a few bubble domains or very large antiskyrmions are left over for B > 400 mT [see Fig. 4(h)]. The last of those switches into the field-polarized state when B = 450 mT is reached, consistent with the isothermal magnetization curve, taking into account the limited field of view of the MFM. Overall, there is perfect agreement between the real-space images and the behavior of the diffuse scattering pattern described before.

C. Magnetic texture at $T < T_{SR}$

Finally, it is important to understand how the magnetic texture changes with temperature. The crucial point is whether or not the rectangular domain structure persists at any T < $T_{\rm C}$, including temperatures below the spin-reorientation (SR) transition $T_{SR} = 170$ K. The SR transition in Mn_{1.4}PtSn can be detected by a step in the temperature-dependent magnetic susceptibility and by a kink in the resistivity [38,39]. Powder neutron diffraction measurements showed that the local (on the scale of the unit cell) magnetic structure of Mn_{1.4}PtSn is collinear ferromagnetic above T_{SR} and becomes canted (noncollinear) ferromagnetic below T_{SR} [38,53]. The correlations between the local (within the unit cell) magnetic configuration and the topology of the large-scale magnetic texture seemed controversial. On the one hand, the magnetotransport measurements of bulk single crystals revealed a topological Hall effect (THE) below T_{SR} [38], which is widely associated with the skyrmion phase in other materials [54–60] (antiskyrmions are expected to give rise to a THE similar to that of skyrmions [6]). On the other hand, the LTEM measurements [27–30] demonstrated that the antiskyrmions nucleate only at $T > T_{SR}$, which suggests that the THE is related to the local noncollinear structure.

Again, we first describe the SANS measurements. Figure 5(a) shows SANS patterns (no background subtraction) collected after zero-field cooling to two temperatures above $T_{\rm SR}$, namely, T = 280 and 180 K, as well as to two temperatures below T_{SR} , namely, T = 160 K and T = 90 K. The corresponding azimuthal profiles and the radial profiles of the intensity are plotted in Figs. 5(b) and 5(c), respectively. The patterns at 280 and 180 K exhibit the same anisotropy. Despite the fact that the magnetic moment is reduced at higher temperatures due to thermal fluctuations (which reduces the intensity of magnetic scattering at elevated temperatures), the intensity at 280 K is significantly higher. At T = 160 K, which is just below T_{SR} , the stripes of intensity can still be distinguished but have much weaker intensity than at T = 180 K. Far below T_{SR} at 90 K, the cross-shaped scattering disappears completely. The remaining diamond-shaped scattering is similar to the pattern at the field-polarized state at 250 K [Fig. 1(b)]. Finally, in Fig. 5(d), the scattering intensity of the (100) streaks is plotted as a function of temperature. It decreases upon cooling with a kink at $T = T_{SR}$. This shows that the fractal magnetic domain pattern is inherent in the high-temperature phase with the locally collinear magnetic order. However, it does not transform to the homogeneous state immediately below the

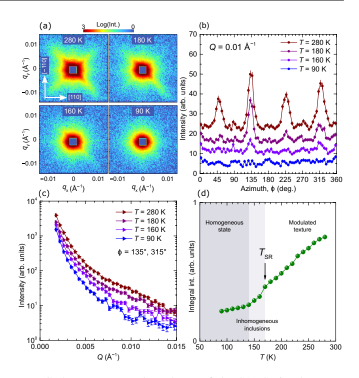


FIG. 5. Temperature dependence of the SANS signal at zero field. (a) A series of SANS patterns (no background subtraction) collected at different temperatures in zero-field cooling. (b) The azimuthal profiles of the scattering extracted from each patter. (c) The radial intensity profiles at the same temperatures. (d) The integral intensity of the magnetic scattering as a function of T. The black arrow marks the spin-reorientation transition.

SR transition. Instead, traces of the anisotropic scattering are present over a 20–30 K wide region below T_{SR} , which correlates with the gradual change in the spin canting within the unit cell [38,53].

The loss of the fractal self-affinity as well as the absence of lamellar stripe domains in zero field is apparent from the MFM measurements obtained below at T_{SR} both T = 140 and 80 K, which we show in Figs. 6(a) and 6(c), respectively. This is also supported by the fitting of the low-temperature SANS data (Fig. S4 in the Supplemental Material [37]), which is described by the exponent *D* that is close to simple Porod scattering. Moreover, also the strong anisotropic pinning of the domain walls to two perpendicular directions within the *ab* plane is lost. Interestingly, in higher fields, again, round-shaped domains appear before the field-polarized state is reached [see Figs. 6(b) and 6(d)]. In comparison to the reported LTEM measurements [27–30], where antiskyrmions were present only above T_{SR} , it is very likely that the domains here are (closure) bubble domains rather than antiskyrmions.

IV. DISCUSSION AND CONCLUSIONS

To conclude, we used a combination of SANS and MFM to study the bulk magnetic structure of $Mn_{1.4}$ PtSn and showed that it differs drastically from what was previously reported from LTEM measurements of thin-plate samples. The bulk $Mn_{1.4}$ PtSn does not support antiskyrmions or any other type of regular long-periodic single-*q* or multi-*q* structures but develops ferromagnetic lamellar stripe domains combined

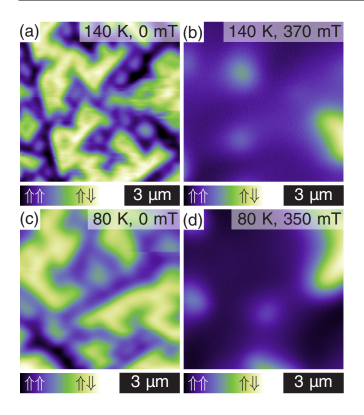


FIG. 6. Magnetic domain pattern in MFM below T_{SR} at (a) and (b) T = 140 K and (c) and (d) T = 80 K both (a) and (c) in zero field and (b) and (d) in high field. Sample magnetization parallel and antiparallel to the external field is shown in purple and green, respectively. The span of the color scale has been adapted individually.

with an anisotropic fractal surface domain pattern that has a characteristic scale with a lower boundary of ~ 48 nm and an upper boundary of $\sim 3 \ \mu m$ defined by the width of the lamellar stripe domains. This magnetic pattern gives rise to an anisotropic diffuse intensity distribution in SANS and, to the best of our knowledge, was previously observed only in the low-temperature phase of Nd₂Fe₁₄B. Yet it differs from the latter in the presence of characteristic hints of the DMI inherent in the D_{2d} symmetry of the crystal, which manifest in the orientation of arrowhead-shaped nested domains at the domain walls of the lamellar stripe domains. Our measurements showed that the magnetic texture of bulk Mn14PtSn is polarized by the applied magnetic field in a multistep process. During the first step, the spins gradually cant towards the field direction, keeping the overall domain pattern unaffected. During the second step, the fractal domain pattern softens, and the nested domains transform into an assembly of bubble domains. Finally, in the third step, only the bubble domains persist and switch individually into the field-polarized state. The latter two steps resemble the metamagnetic phases of the dense antiskyrmion lattice and the isolated antiskyrmions in the ferromagnetic background, respectively, observed in the thin plates [27-30]. Like the antiskyrmions in the thin-plate geometry, the fractal magnetic pattern of the bulk Mn_{1.4}PtSn appears for $T > T_{SR}$ and shows an enhanced stability at elevated temperatures, which further demonstrates the intimate connection between the magnetic structure in the bulk and in the thin plates.

To the best of our knowledge, Mn_{1,4}PtSn is therefore the first known material where the magnetic texture can be unambiguously tuned between ferromagnetic domains with an anisotropic fractal closure domain pattern in zero field or bubble domains in high field in bulk samples and helices or antiskyrmions in the thin-plate geometry. A sizable anisotropic DMI allowed in the D_{2d} symmetry, if present, will result in a modulated texture with a period defined by the ratio between the exchange interaction J and the DMI constant D, $\lambda \propto J/D$, independent of the sample thickness. Contrary to the previous interpretations of the experiments on the thin plates, the observed complex multiscale domain-wall structure of bulk Mn_{1.4}PtSn suggests that the DMI plays only a minor role in this material. The other relevant magnetic energy terms include the second-order uniaxial and fourthorder in-plane magnetocrystalline anisotropies, as well as the sample thickness-dependent dipole-dipole energy, the competition of which is known to influence the width of domain walls [35,61–63]. Thus, our findings highlight the importance of the dipolar interaction as an essential ingredient to describe the magnetic texture of Mn_{1.4}PtSn. This was also realized to be important for explaining all the observations in recent LTEM experiments [29,30].

Moreover, we may speculate that new materials may be (anti)skyrmion hosts in the thin-plate geometry even if they have not been identified as such in bulk experiments. Possible candidates would be uniaxial materials with fractal closure domains in the bulk and with D_{2d} symmetry. Since a noncentrosymmetric crystal symmetry is the only necessary condition for the possible existence of (anti)skyrmions, the range of materials that support (anti)skyrmions in the thin-plate geometry could show more complicated domain patterns in the bulk. For example, this could be the case for $Cr_{11}Ge_{19}$, which hosts biskyrmions in thin-plate geometry [64].

Note added. Recently, we became aware of a thicknessdependent study in the range up to $4 \mu m$ of the magnetic texture in Mn_{1.4}PtSn [65]. The authors explained the crucial role of dipolar forces in the material by rigorous simulations, which fully agree with the conclusions drawn in our study.

ACKNOWLEDGMENTS

A.S.S. thanks S. E. Nikitin for many stimulating discussions. B.E.Z.C. thanks U. Burkhardt and S. Kostmann for electron backscatter diffraction measurements and polishing of the polycrystalline sample. This project was funded by the German Research Foundation (DFG) under Grants No. FE 633/30-1, No. IN 209/9-1, No. EN 434/38-1, and No. MI 2004/3-1, as well as Grants No. EN 434/40-1 and No. IN 209/7-1 as part of the Priority Program SPP 2137 "Skyrmionics"; via Projects C03 and C05 of the Collaborative Research Center SFB 1143 (Project-id 247310070) at TU Dresden; and the Würzburg-Dresden Cluster of Excellence on Complexity and Topology in Quantum Matter-ct.qmat (EXC 2147, Project-id 390858490). A.S.S. and B.E.Z.C. acknowledge support from the International Max Planck Research School for Chemistry and Physics of Quantum Materials (IMPRS-CPQM).

- [1] W. Koshibae and N. Nagaosa, Nat. Commun. 7, 10542 (2016).
- [2] S. Huang, C. Zhou, G. Chen, H. Shen, A. K. Schmid, K. Liu, and Y. Wu, Phys. Rev. B 96, 144412 (2017).
- [3] L. Camosi, N. Rougemaille, O. Fruchart, J. Vogel, and S. Rohart, Phys. Rev. B 97, 134404 (2018).
- [4] M. Hoffmann, B. Zimmermann, G. P. Müller, D. Schürhoff, N. S. Kiselev, C. Melcher, and S. Blügel, Nat. Commun. 8, 308 (2017).
- [5] X. Zhang, J. Xia, Y. Zhou, X. Liu, H. Zhang, and M. Ezawa, Nat. Commun. 8, 1717 (2017).
- [6] N. Nagaosa and Y. Tokura, Nat. Nanotechnol. 8, 899 (2013).
- [7] S. Mühlbauer, B. Binz, F. Jonietz, C. Pfleiderer, A. Rosch, A. Neubauer, R. Georgii, and P. Böni, Science 323, 915 (2009).
- [8] J. Kindervater, T. Adams, A. Bauer, F. X. Haslbeck, A. Chacon, S. Mühlbauer, F. Jonietz, A. Neubauer, U. Gasser, G. Nagy, N. Martin, W. Häussler, R. Georgii, M. Garst, and C. Pfleiderer, Phys. Rev. B 101, 104406 (2020).
- [9] X. Z. Yu, N. Kanazawa, Y. Onose, K. Kimoto, W. Z. Zhang, S. Ishiwata, Y. Matsui, and Y. Tokura, Nat. Mater. 10, 106 (2011).
- [10] H. Wilhelm, M. Baenitz, M. Schmidt, U. K. Rössler, A. A. Leonov, and A. N. Bogdanov, Phys. Rev. Lett. 107, 127203 (2011).
- [11] P. Milde, D. Köhler, J. Seidel, L. M. Eng, A. Bauer, A. Chacon, J. Kindervater, S. Mühlbauer, C. Pfleiderer, C. Schütte, and A. Rosch, Science 340, 1076 (2013).
- [12] E. Moskvin, S. Grigoriev, V. Dyadkin, H. Eckerlebe, M. Baenitz, M. Schmidt, and H. Wilhelm, Phys. Rev. Lett. 110, 077207 (2013).
- [13] Y. Tokunaga, X. Z. Yu, J. S. White, H. M. Rønnow, D. Morikawa, Y. Taguchi, and Y. Tokura, Nat. Commun. 6, 7638 (2015).
- [14] K. Karube, J. S. White, N. Reynolds, J. L. Gavilano, H. Oike, A. Kikkawa, F. Kagawa, Y. Tokunaga, H. M. Ronnow, Y. Tokura, and Y. Taguchi, Nat. Mater. 15, 1237 (2016).
- [15] V. Ukleev, Y. Yamasaki, D. Morikawa, K. Karube, K. Shibata, Y. Tokunaga, Y. Okamura, K. Amemiya, M. Valvidares, H. Nakao, Y. Taguchi, Y. Tokura, and T. Arima, Phys. Rev. B 99, 144408 (2019).
- [16] K. Karube, J. S. White, D. Morikawa, M. Bartkowiak, A. Kikkawa, Y. Tokunaga, T. Arima, H. M. Ronnow, Y. Tokura, and Y. Taguchi, Phys. Rev. Mater. 1, 074405 (2017).
- [17] K. Karube et al., Sci. Adv. 4, eaar7043 (2018).
- [18] S. Seki, X. Z. Yu, S. Ishiwata, and Y. Tokura, Science 336, 198 (2012).
- [19] T. Adams, A. Chacon, M. Wagner, A. Bauer, G. Brandl, B. Pedersen, H. Berger, P. Lemmens, and C. Pfleiderer, Phys. Rev. Lett. 108, 237204 (2012).
- [20] S. L. Zhang, A. Bauer, D. M. Burn, P. Milde, E. Neuber, L. M. Eng, H. Berger, C. Pfleiderer, G. van der Laan, and T. Hesjedal, Nano Lett. 16, 3285 (2016).
- [21] P. Milde, E. Neuber, A. Bauer, C. Pfleiderer, H. Berger, and L. M. Eng, Nano Lett. 16, 5612 (2016).
- [22] A. Stefancic, S. H. Moody, T. J. Hicken, M. T. Birch, G. Balakrishnan, S. A. Barnett, M. Crisanti, J. S. O. Evans, S. J. R. Holt, K. J. A. Franke, P. D. Hatton, B. M. Huddart, M. R. Lees, F. L. Pratt, C. C. Tang, M. N. Wilson, F. Xiao, and T. Lancaster, Phys. Rev. Mater. 2, 111402(R) (2018).

- [23] A. S. Sukhanov, P. Vir, A. S. Cameron, H. C. Wu, N. Martin, S. Mühlbauer, A. Heinemann, H. D. Yang, C. Felser, and D. S. Inosov, Phys. Rev. B 100, 184408 (2019).
- [24] I. Kézsmárki, S. Bordács, P. Milde, E. Neuber, L. M. Eng, J. S. White, H. M. Rønnow, C. D. Dewhurst, M. Mochizuki, K. Yanai, H. Nakamura, D. Ehlers, V. Tsurkan, and A. Loidl, Nat. Mater. 14, 1116 (2015).
- [25] Y. Fujima, N. Abe, Y. Tokunaga, and T. Arima, Phys. Rev. B 95, 180410(R) (2017).
- [26] S. Bordács, A. Butykai, B. G. Szigeti, J. S. White, R. Cubitt, A. O. Leonov, S. Widmann, D. Ehlers, H.-A. Krug von Nidda, V. Tsurkan, A. Loidl, and I. Kézsmárki, Sci. Rep. 7, 7584 (2017).
- [27] A. K. Nayak, V. Kumar, T. Ma, P. Werner, E. Pippel, R. Sahoo, F. Damay, U. K. Rössler, C. Felser, and S. S. P. Parkin, Nature (London) 548, 561 (2017).
- [28] R. Saha, A. K. Srivastava, T. Ma, J. Jena, P. Werner, V. Kumar, C. Felser, and S. S. P. Parkin, Nat. Commun. 10, 5305 (2019).
- [29] J. Jena, B. Göbel, T. Ma, V. Kumar, R. Saha, I. Mertig, C. Felser, and S. S. P. Parkin, Nat. Commun. 11, 1115 (2020).
- [30] L. Peng, R. Takagi, W. Koshibae, K. Shibata, K. Nakajima, T. Arima, N. Nagaosa, S. Seki, X. Yu, and Y. Tokura, Nat. Nanotechnol. 15, 181 (2020).
- [31] J. Jena, R. Stinshoff, R. Saha, A. K. Srivastava, T. Ma, H. Deniz, P. Werner, C. Felser, and S. S. P. Parkin, Nano Lett. 20, 59 (2020).
- [32] O. Meshcheriakova, S. Chadov, A. K. Nayak, U. K. Rössler, J. Kübler, G. Andre, A. A. Tsirlin, J. Kiss, S. Hausdorf, A. Kalache, W. Schnelle, M. Nicklas, and C. Felser, Phys. Rev. Lett. 113, 087203 (2014).
- [33] A. N. Bogdanov, U. K. Rössler, M. Wolf, and K.-H. Müller, Phys. Rev. B 66, 214410 (2002).
- [34] S. Mühlbauer, D. Honecker, E. A. Perigo, F. Bergner, S. Disch, A. Heinemann, S. Erokhin, D. Berkov, C. Leighton, M. R. Eskildsen, and A. Michels, Rev. Mod. Phys. 91, 015004 (2019).
- [35] A. Hubert and R. Schäfer, Magnetic Domains: The Analysis of Magnetic Microstructures (Springer, Berlin, 1998).
- [36] G. Chaboussant, S. Desert, P. Lavie, and A. Brulet, J. Phys.: Conf. Ser. 340, 012002 (2012).
- [37] See Supplemental Material at http://link.aps.org/supplemental/ 10.1103/PhysRevB.102.174447 for additional information about the samples and complementary MFM measurements.
- [38] P. Vir, J. Gayles, A. S. Sukhanov, N. Kumar, F. Damay, Y. Sun, J. Kübler, C. Shekhar, and C. Felser, Phys. Rev. B 99, 140406(R) (2019).
- [39] P. Vir, N. Kumar, H. Borrmann, B. Jamijansuren, G. Kreiner, C. Shekhar, and C. Felser, Chem. Mater. 31, 5876 (2019).
- [40] Park Systems Corp. KANC 15F, Suwon, Korea.
- [41] Nanosensors, Neuchatel, Switzerland.
- [42] Omicron NanoTechnology Gmbh, Taunusstein, Germany.
- [43] RHK Technology, Inc., Troy, Michigan.
- [44] D. Nečas and P. Klapetek, Cent. Eur. J. Phys. 10, 181 (2012).
- [45] P. W. Schmidt, in *The Modern Aspects of Small-Angle Scattering*, edited by H. Brumberger (Kluwer Academic, Dordrecht, 1995), p. 1.
- [46] D. F. R. Mildner and P. L. Hall, J. Phys. D 19, 1535 (1986).
- [47] J. E. Martin and A. J. Hurd, J. Appl. Crystallogr. 20, 61 (1987).
- [48] E. G. Iashina, E. V. Velichko, M. V. Filatov, W. G. Bouwman, C. P. Duif, A. Brulet, and S. V. Grigoriev, Phys. Rev. E 96, 012411 (2017).

- [49] A. Kreyssig, R. Prozorov, C. D. Dewhurst, P. C. Canfield, R. W. McCallum, and A. I. Goldman, Phys. Rev. Lett. **102**, 047204 (2009).
- [50] Yu. G. Pastushenkov, A. Forkl, and H. Kronmüller, J. Magn. Magn. Mater. 174, 278 (1997).
- [51] Yu. G. Pastushenkov, Bull. Russ. Acad. Sci.: Phys. 74, 1423 (2010).
- [52] B. B. Mandelbrot, *The Fractal Geometry of Nature* (Holt, New York, 1983).
- [53] V. Kumar, N. Kumar, M. Reehuis, J. Gayles, A. S. Sukhanov, A. Hoser, F. Damay, C. Shekhar, P. Adler, and C. Felser, Phys. Rev. B 101, 014424 (2020).
- [54] A. Neubauer, C. Pfleiderer, B. Binz, A. Rosch, R. Ritz, P. G. Niklowitz, and P. Böni, Phys. Rev. Lett. **102**, 186602 (2009).
- [55] S. X. Huang and C. L. Chien, Phys. Rev. Lett. **108**, 267201 (2012).
- [56] J. C. Gallagher, K. Y. Meng, J. T. Brangham, H. L. Wang, B. D. Esser, D. W. McComb, and F. Y. Yang, Phys. Rev. Lett. 118, 027201 (2017).
- [57] N. Kanazawa, Y. Onose, T. Arima, D. Okuyama, K. Ohoyama, S. Wakimoto, K. Kakurai, S. Ishiwata, and Y. Tokura, Phys. Rev. Lett. **106**, 156603 (2011).

- [58] T. Yokouchi, N. Kanazawa, A. Tsukazaki, Y. Kozuka, M. Kawasaki, M. Ichikawa, F. Kagawa, and Y. Tokura, Phys. Rev. B 89, 064416 (2014).
- [59] Y. Li, N. Kanazawa, X. Z. Yu, A. Tsukazaki, M. Kawasaki, M. Ichikawa, X. F. Jin, F. Kagawa, and Y. Tokura, Phys. Rev. Lett. 110, 117202 (2013).
- [60] C. Franz, F. Freimuth, A. Bauer, R. Ritz, C. Schnarr, C. Duvinage, T. Adams, S. Blügel, A. Rosch, Y. Mokrousov, and C. Pfleiderer, Phys. Rev. Lett. 112, 186601 (2014).
- [61] M. Ezawa, Phys. Rev. Lett. 105, 197202 (2010).
- [62] A. Bernand-Mantel, C. B. Muratov, and T. M. Simon, Phys. Rev. B 101, 045416 (2020).
- [63] X. Yu, M. Mostovoy, Y. Tokunaga, W. Zhang, K. Kimoto, Y. Matsui, Y. Kaneko, N. Nagaosa, and Y. Tokura, Proc. Natl. Acad. Sci. USA 109, 8856 (2012).
- [64] R. Takagi, X. Z. Yu, J. S. White, K. Shibata, Y. Kaneko, G. Tatara, H. M. Rønnow, Y. Tokura, and S. Seki, Phys. Rev. Lett. 120, 037203 (2018).
- [65] T. Ma, A. K. Sharma, R. Saha, A. K. Srivastava, P. Werner, P. Vir, V. Kumar, C. Felser, and S. S. P. Parkin, Adv. Mater. 32, 2002043 (2020).