Evolution of Magnetism in Magnetic Topological Semimetal NdSb_xTe_{2-x+δ}

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Abstract

Magnetic topological semimetals LnSbTe (Ln = Lanthanide) have attracted intensive attention because of the presence of interplay between magnetism, topological, and electron correlations depending on the choices of magnetic Ln elements. Recently, varying Sb-Te composition has been found to effectively control the electronic and magnetic states in LnSb_xTe_{2x}. With this motivation, we report the evolution of magnetic properties with Sb-Te substitution in NdSb_xTe_{2-x+δ}, ($0 \le x \le 1$). Our work reveals the interesting non-monotonic change in magnetic ordering temperature with varying composition stoichiometry. In addition, reducing the Sb content x drives the reorientation of moments from in-plane (ab-plane) to out-of-plane (c-axis) direction that results in the distinct magnetic structures for two end compounds NdTe₂ (x = 0) and NdSbTe (x = 1). Furthermore, the moment orientation in NdSb_xTe_{2-x+δ} is also found to be strongly tunable upon application of weak magnetic field, leading to rich magnetic phases depending on the composition stoichiometry, temperature, and magnetic field. Such strong tuning of magnetism in this material establishes it as a promising platform for investigating tunable topological states and correlated topological physics.

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I. Introduction

Topological semimetals (TSMs) such as Dirac or Weyl semimetals feature symmetryprotected linearly dispersed Dirac or Weyl cones in their electronic structures, which host relativistic fermions with low-energy excitations can be described by Dirac or Weyl equations respectively [1,2]. In contrast to Dirac or Weyl nodes at discrete points in the momentum space in Dirac and Weyl semimetals, another class of TSMs i.e., nodal-line semimetals exhibit interesting linear band crossings along one-dimensional loops or lines. Various exotic properties have been discovered, such as large magnetoresistance [3], ultrahigh mobility [3], chiral anomaly [4,5], and surface Fermi arcs [6–9], providing deeper understanding of fundamental topological physics as well as opportunities for future technological applications. Within these categories of TSMs, there has recently been rapidly growing interest in magnetic TSMs such as Co₂Mn(Al/Ga) [10–14], Co₃Sn₂S₂ [15,16], FeSn [17], Fe₃Sn₂ [18], Fe₃(Al/Ga) [19], Fe₃GeTe₂ [20], Mn₃(Ge/Sn) [21–24], and GdPtBi [25]. These compounds offer a rare platform to investigate interplay between magnetism and non-trivial band topology, which can generate novel exotic quantum phenomena such as large intrinsic anomalous Hall effect [26] and anomalous Nernst effect [27].

As described above, the magnetism in the majority of magnetic TSMs reported so far originates from 3*d* transition metal elements. In addition to transition metal-based compounds, magnetic lanthanide (*Ln*)-based TSMs are also highly desired because of strong correlation effects brought by the 4*f* electrons. The *Ln*SbTe compounds represent one such model example [28–44]. *Ln*SbTe belongs to ZrSiS-type nodal-line semimetal family that can be represented by a general chemical formula *WHM* (*W*= Zr/Hf/lanthanides; *H*=Si/Ge/Sn/Sb, *M* = S, Se,Te). Those materials crystallize in a layered PbFCl-type crystal structure (space group *P*4/*nmm*), characterized by square or nearly square net layers of *H*-atoms which harbor relativistic fermions [28,31,45–60]. In

*Ln*SbTe, the presence of magnetic *Ln* element such as Ce [28–30], Nd [37,38,44], Sm [35,36], Gd [31–33], Tb [42,43], Dy [41,44], Ho [39,40,43], and Er [44] activates the spin degree of freedom that leads to diverse antiferromagnetic (AFM) ground states depending on the choice of *Ln* [28,31,35,37,39,41,44]. Varying *Ln* element is also found to tune the topological states generated by Sb layers [28,31,35,39–41,57,58,61,62], providing opportunity to engineer band topology via coupling between magnetism and electronic bands [28]. Furthermore, rich quantum phenomena such as Kondo effect, charge density waves (CDWs), and correlation enhancement have been reported in various *Ln*SbTe compounds [28,29,33,35,37,61,63,64], which are also strongly dependent on the choices of *Ln* elements.

Besides magnetism from *Ln*, tuning Sb-Te composition stoichiometry has been established as an effective approach to engineer electronic and magnetic phases in *Ln*Sb_xTe_{2-x} [33,36,61,62,62,62,63,65]. Given that the square-net lattices are inherently unstable [66], doping electrons by substituting Te for Sb causes distortion of Sb-square net and the subsequent formation of CDWs, which has been found to modify electronic band structures and intrinsic magnetism [33,36,61–63,65]. In *Ln*Sb_xTe_{2-x}, the magnetism has been found to be effectively tunable with varying Sb composition *x*, however the Sb-Te substitution induces distinct evolution of magnetic properties in few *Ln*SbTe (*Ln* = Ce [30], Sm [36], and Gd [33,63]) compounds despite of similar crystal symmetry and structure evolution from tetragonal (space group *P4/nmm*) to orthorhombic (space group *Pmmm*) phase in off-stoichiometric compositions. While numerous stoichiometric *Ln*SbTe compounds [28,29,33,35,37,61,63,64] have been discovered, the evolution of magnetism by tuning composition in off-stoichiometric compounds is still in an early stage. Only recently, a neutron diffraction study has determined the modification of magnetic structure with varying Sb-Te composition in NdSb_{0.94}Te_{0.92} and NdSb_{0.48}Te_{1.37} [65]. Such promising results demand a complete understanding of magnetism and possible tuning of topological states over entire Sb-Te composition in $NdSb_xTe_{2-x}$. In this work, we present the evolution of magnetic properties from x = 0 to 1 in $NdSb_xTe_{2-x}$. Our work reveals an interesting non-monotonic variation of magnetic ordering temperature (T_N) and the reorientation of Nd moments with Sb-Te substitution. These results provide a rich platform for tunable topological states and further studying the correlated topological physics.

II. Experiment

The NdSb_xTe_{2-x+ δ} ($0 \le x \le 1$, δ represents possible vacancies) single crystals used in this work were synthesized by a chemical vapor transport (CVT) method using I₂ as the transport agent. The pristine NdTe₂ was grown by a direct CVT method with elementary Nd and Te powders as source materials. For each of the other compositions with Sb, a polycrystalline precursor is necessary to minimize vacancies (as discussed below). Each precursor was prepared by heating the mixture of Nd with different ratios of Sb and Te powders (shown in Table I) at 850 °C for 2 days. The single crystals were obtained via CVT with a temperature gradient from 1000 to 850 °C for two weeks. The elemental compositions and crystal structures of the obtained crystals were examined by energy-dispersive x-ray spectroscopy (EDS) and x-ray diffraction (XRD), respectively. Magnetization measurements up to 9 T were performed by using a physical property measurement system (PPMS, Quantum Design). Magnetization measurements up to 7 T and angular dependent magnetization were performed by using a magnetic property measurement system (MPMS, Quantum Design) equipped with a rotator.

III. Result and discussion

Recent surge of interests in TSMs featuring a square net of atoms has motivated the study of LnSbTe family [28–44]. While the square net planes in majority of WHM compounds are formed by group-IV elements H = Si, Ge, and Sn [45–56,67], a Sb (group-V) square net sandwiched by Ln-Te bilayers is present in LnSbTe compounds. Previous studies have demonstrated that the synthesis of ideal stoichiometric LnSbTe compounds is challenging [29,37,65] and often yield various off-stoichiometric $LnSb_xTe_{2-x}$ (0 < x < 1) compositions consisting of distorted Sb plane [33,36,61,62,62,62,63,65]. Such off-stoichiometry is also accompanied by vacancies in the Sb [33,36,63–65] and Te [33,61,62,65] layers that enhance with reducing Sb content [36,63,65], and eventually after complete substitution of Te for Sb produces a strong Te vacancy in structurally similar $LnTe_2$ compounds [68,69]. The chalcogen vacancy widely occurs in LnX_2 (X = S, Se, and Te), which has been ascribed to the presence of mixed anions $(X_2)^{2-}$ and $(X)^{2-}$ in chalcogen layer [69,70] and favored by decreasing Ln^{3+} cation radius along the lanthanide series [70]. Therefore, chalcogen vacancy is usually seen in compounds with smaller Ln^{3+} cations such as NdTe_{1.89} [68], SmTe_{1.84} [71], GdTe_{1.80} [72], TbTe_{1.80} [72], and DyTe_{1.80} [72]. Replacing Sb for Te in these compounds induces hole doping and consequently reduces the number of vacancies required for charge balance in Te layers [63].

In NdSb_xTe_{2-x} studied in this work, both slight [65] and the lack [44] of Te vacancy are observed in a nearly-stoichiometric composition. Although an earlier study on CeSb_xTe_{2-x- δ} [62] has claimed less effect of Te vacancy on magnetization, we selected NdSb_xTe_{2-x} samples with minimum vacancy to ensure the systematic tuning of magnetization with varying composition and without the interference of vacancy effects. The millimeter-size single crystals in the whole compositional range from NdTe₂ (*x* = 0) to NdSbTe_{1.08} (*x* = 1) were obtained using CVT [Fig. 1(a)],

similar to previous NdSbTe growths [37,38,44]. Our EDS results have revealed the nearly 1:2 stoichiometric composition ratio between Nd and (Sb+Te) (Table I), from which we conclude the absence of vacancies within the resolution limit of our instrument. Such observation is in contrast to the reported Ce, Sm, and Gd sibling compounds [33,36,62-64], which might be due to the slight difference in synthesis method and sample screening. However, we observed a slight excess of Te in our composition analysis. Therefore, our samples can be represented as $NdSb_xTe_{2-x+\delta}$, where δ = 0.0.08 denotes the excess Te that might be due to the instrument error or partial accommodation into the Sb layer, as seen in CeSbTe [29] and GdSbTe [63]. As shown in Table I, the nominal composition in source materials is found to yield significantly different composition (determined by EDS) in the grown crystals, consistent with previous Sb-Te substitution studies in this family [36,63]. Earlier work on $GdSb_xTe_{2-x-\delta}$ has adopted an strategy of adding more Sb in the starting materials to obtain crystals with increasing x [63]. In contrast, we did not observe a systematic correlation between nominal and final compositions among the grown crystals [Table I], indicating the difficulty to control composition stoichiometry in NdSb_xTe_{2-x+ δ}. This could be the reason for the lack of a complete Sb-Te evolution study for this compound although NdSbTe [37] was one of the earliest studied *Ln*SbTe compounds.

The stoichiometric *Ln*SbTe compounds crystallize in tetragonal (space group *P4/nmm*) structure [28–44], which on substituting Te for Sb results in structure transition to orthorhombic phase at around x = 0.7 to 0.85 in *Ln*Sb_xTe_{2-x} (*Ln* = Ce [30], Sm [36], and Gd [63]). The structural information determined by our structure refinement using powder XRD spectra presented in Fig. 1(b) also reveals an orthorhombic distortion at around $x \approx 0.70$ in NdSb_{x- δ}Te_{2-x+ δ} that is accompanied by shrinking *c*-axis and expanding *ab*-plane [Fig. 1(c)], consistent with other *Ln*Sb_xTe_{2-x} (*Ln* = Ce, Sm, and Gd) members [36,62,63]. The tetragonal crystal lattice is retained

on further decreasing the Sb content below $x \approx 0.18$ leading to tetragonal structure for NdTe₂ (x = 0). Both tetragonal and orthorhombic structures have been reported in Te-deficient compounds NdTe_{1.80} [73] and NdTe_{1.89} [68], respectively. The tetragonal structure has been identified in analogous compounds CeTe₂ [74] and PrTe₂ [75] whereas LaTe₂ crystallizes in monoclinic structure [76]. The tetragonal and orthorhombic structures are very similar and could be difficult to distinguish during crystal structure refinement. In addition, because of the presence of CDW in this material family (especially in the Sb-less compositions), the refinement results could be influenced by the additional CDW satellite peaks. Unfortunately, the impact of CDW in the crystal structure refinement for our $NdSb_{x-\delta}Te_{2-x+\delta}$ samples is difficult to clarify due to instrument limitation of our x-ray diffractometer. For this, a synchrotron light source is needed, which could be the scope for future studies that might also clarify the evolution of CDW with composition stoichiometry in NdSb_xTe_{2-x+ δ}. Though we are not able to examine CDW due to instrumental limitations, the re-emergence of tetragonal structure for x < 0.18 in NdSb_xTe_{2-x+ δ} can be understood by the evolution of XRD spectra with varying composition stoichiometry. As shown in Fig. 1b, the XRD spectra display clear peak splitting for samples with intermediate Sb content (indicated by the red arrows), which is consistent with the lowering symmetry from tetragonal to orthorhombic.

With the orthorhombically distorted lattice, the 2D Sb square net characterized by identical Sb-Sb bonding length and 90° bonding angles in stoichiometric has been found to undergo deviation of bonding angles from 90° with reducing Sb content that indicates the distorted Sb square plane [36,62,63,65]. Distorted Sb-square nets in orthorhombic crystal lattice depending on the composition stoichiometry in *Ln*SbTe are reported to drive tunable CDWs [33,36,61–63,65]. Even the formation of CDW has been revealed in tetragonal CeTe₂ [77–79] and PrTe₂ [77] as well as monoclinic LaTe₂ [80]. The CDW is found to strongly interplay with magnetism in *Ln*SbTe

leading to complex magnetic phases in off-stoichiometric compositions [33,62,65] and even a modification of a collinear AFM structure in a nearly stoichiometric composition to a more complex elliptical cycloid magnetic structure for a Sb-less composition in NdSb_xTe_{2-x-δ} [65]. To investigate the evolution of magnetic properties over entire composition range in $NdSb_xTe_{2-x+\delta}$, we have measured the temperature dependence of susceptibility $\chi(T)$ under in-plane (H//ab) and outof-plane (*H*//*c*) magnetic fields of $\mu_0 H = 0.1$ T [Fig. 2(a)] in order to investigate the variation of and moment orientation with Sb-Te substitution. Similar the $T_{\rm N}$ to previous studies [35,37,41,44,63], T_N for each sample, except for x = 0.29, is extracted from the susceptibility peak or anomaly in in-plane (χ_{ll} , measured with field parallel to the *ab*-plane) and out-of-plane (χ_{\perp} ; measured with field parallel to the *c*-axis) susceptibility measurements, as indicated by the solid triangles in Fig. 2a. The obtained transition temperatures are consistent with that determined from the derivative susceptibility dx/dT [Fig. 2(b)]. Furthermore, for the x = 0.29sample which does not display clear feature in susceptibility measurement, the derivative susceptibility reveals the possible magnetic transition temperature, as shown in Fig. 2b. The extracted T_N of 2.04 K for the end compounds NdTe₂ (x = 0) is distinct from the lack of magnetic ordering down to 2 K in NdTe_{1.89} with significant Te vacancy [68]. Such difference might be attributed to the suppression of Te-mediated magnetic interaction between Nd moments. For the other end compound NdSbTe_{1.08} (x = 1), $T_N \approx 2.74$ K is consistent with the nearly-stoichiometric composition NdSb_{0.94}Te_{0.92} [65] but slightly lower than the ideal stoichiometry NdSbTe ($T_N \approx 3.1$ K) [44]. In Fig. 3(a) we summarized the magnetic transition temperatures for our samples, which exhibit a non-monotonic composition dependence. Both monotonic [62] and non-monotonic [63] composition-dependent T_N have been observed in $LnSb_xTe_{2-x}$. In CeSb_xTe_{2-x} [62], the T_N systematically increases with decreasing Sb content x, while a similar non-monotonic composition

dependence of T_N is seen in GdSb_xTe_{2-x} [63]. The monotonic variation of T_N in CeSb_xTe_{2-x}, which is in contrast to Nd and Gd samples, might be attributed to difference in interplay between CDWs and magnetism [65] arising from the distinct moment orientation in CeSbTe [62] as compared to NdSbTe [65] and GdSbTe [33]. In *Ln*Sb_xTe_{2-x}, the CDWs exhibit single modulation wave-vector q within the Sb plane in the intermediate Sb-composition range i.e., (0.21-0.34) < x < (0.74-0.85) whereas below x < (0.21-0.34) these compounds host multiple q-vectors along different crystallographic axes [62,63]. The out-of-plane moment orientation in CeSb_xTe_{2-x} [62] does not align with CDW q-vector, especially in single q-vector (aligned along the *ab*-plane) region for intermediate Sb-composition of T_N . On the other side, the in-plane orientation of moments in NdSbTe [65] and GdSbTe [33] might strongly couple with in-plane CDW q-vector leading to non-monotonic dependence of T_N with Sb-Te substitution.

As seen in Fig. 3(a), the non-monotonic composition dependence of T_N in NdSb_xTe_{2-x+\delta} involves three distinct regions featuring different orientations for Nd moments: (1) within *ab*-plane (represented as AFM_{*ab*}) for Sb-rich compositions, (2) canted configuration (cAFM) for intermediate Sb-compositions, and (3) along *c*-axis (AFM_{*c*}) for Sb-less compositions. These moment orientations were determined by $\chi(T)$ and were further confirmed by field dependence of magnetization M(H) measurements shown in Fig. 4. Starting with pristine NdSbTe_{1.08} (x = 1), the χ_{II} gradually decreases below $T_N \approx 2.74$ K, in contrast to weakly temperature-dependent χ_{\perp} . This is suggestive of an in-plane AFM configuration, consistent with the reported magnetic structure for nearly-stoichiometric NdSb_{0.94}Te_{0.92} [65]. Decreasing Sb content to x = 0.82, similar $\chi(T)$ trend with unchanged T_N (≈ 2.78 K) is observed [Fig. 3(a)], indicating a similar magnetic ordering to that of the x = 1 compound. The in-plane easy axis is also supported by the M(H) measurements

under in-plane (*H*//*ab*) and out-of-plane (*H*//*c*) magnetic fields at T = 2 K. As shown in Fig. 4, for both x = 1 and 0.82, the in-plane (*M*//; magnetization parallel to the *ab*-plane) isothermal magnetization is larger than the out-of-plane (*M*_⊥; magnetization parallel to the *c*-axis) magnetization, which implies that the easy axis should align towards the *ab*-plane. The magnetic anisotropy in these samples is further manifested in the angular dependence of susceptibility [$\chi(\theta)$] measurements for samples representing three different AFM regions [AFM_{*ab*}, cAFM, and AFM_{*c*} in Fig. 3(a)] at T = 2 K and $\mu_0 H = 0.1$ T [Fig. 3(c)]. For x = 1, the susceptibility is maximum and minimum along the in-plane and out-of-plane fields respectively, which is in line with $\chi(T)$ and *M*(*H*) measurements.

This scenario completely changes after entering into the orthorhombically distorted regime (x < 0.70). As shown in Fig. 2(a), in contrast to the Sb rich samples that display strong drop in χ // while roughly constant χ_{\perp} , both χ // and χ_{\perp} exhibit clear peaks in the x = 0.60 sample followed by a sharp drop and then remain relatively flat down to the lowest measured temperature. Such a similar temperature-dependent behavior down to the lowest measured temperature along the both field directions suggests a spiral spin texture with a canted spin component [33]. The moment canting scenario is consistent with the reported elliptical cycloid magnetic structure propagating along the *b*-axis but also with a non-zero out-of-plane component determined by neutron diffraction experiment for NdSb_{0.48}Te_{1.48} [65]. The elliptical cycloid magnetic structure with both in-plane and out-of-plane moment components can also be understood by relatively less anisotropy between field-dependent $M_{l/}$ and M_{\perp} [81] at T = 2 K for x = 0.60 and 0.45 (Fig. 4). Furthermore, the weaker magnetic anisotropy in orthorhombic regime as compared to x > 0.70 tetragonal samples is clearly demonstrated by the lack of significant anisotropy in the $\chi(\theta)$ measurement in x = 0.60 [Fig. 3(c)]. Despite the well-defined peaks in both χ // and χ_{\perp} for x = 0.60, the drop of χ // below T_N is more

pronounced than χ_{\perp} , which implies the nearly in-plane orientation for canted moments that agrees well with the higher (about 3 times) in-plane than the out-of-plane component in elliptical cycloid magnetic structure for NdSb_{0.48}Te_{1.48} [65]. The moments remain canted for x = 0.45, as demonstrated by the similar magnetic transitions in both χ_{\parallel} and χ_{\perp} at a slightly higher $T_{\rm N}$ (≈ 2.9 K) but reducing the Sb content from x = 0.60 to 0.45 pushes the moments away from the *ab*-plane because the decrease in χ_{ll} and χ_{\perp} below T_N is much more comparable that that of x = 0.60. The spin reorientation in the intermediate Sb-regime has been attributed to the interplay between CDW and magnetism [65], which is in line with the fact that both $T_{\rm N}$ and moment orientation essentially remain unchanged in the tetragonal regime (x > 0.70) where the CDW is absent [62,63]. Such coupling between CDW and magnetism has also been proposed to relieve the magnetic frustration arising due to the competition between nearest-neighbor AFM and ferromagnetic (FM) interactions in a nearly-stoichiometric NdSb_{0.94}Te_{0.92} [65]. In fact, in our x = 1 sample, the PM to AFM transition lacks a sharp peak but is reflected by the broad transition in $\chi_{//}$, indicating frustrated magnetic ordering consistent with a similar composition NdSb_{0.94}Te_{0.92} [65]. The broad T_N peak in x = 1 starts to become sharper with decreasing x leading to sharp magnetic transitions in x = 0.60and 0.45, suggesting the suppression of frustration when CDW is functional. Furthermore, relieving the magnetic frustration would strengthen the magnetic interactions, which explains the systematic rise of T_N with reducing Sb content from x = 1 until reaching a maximum value for x =0.45 [Fig. 3(a)]. The maximal T_N around x = 0.45 is also seen in GdSb_xTe_{2-x-δ} [63], suggesting a similar scenario of coupling between CDW and in-plane magnetic moments in GdSbTe [33].

The T_N reduces on further decreasing the Sb content from x = 0.45 to 0.29. The drop of T_N for x = 0.29 nearly coincides with the phase boundary between single and multiple CDW *q*-vectors identified in CeSb_xTe_{2-x- δ} [62] and GdSb_xTe_{2-x- δ} [63], therefore further decreasing the Sb content

modifies the CDW which is expected to tune the magnetic ordering [65]. In fact, entering into multiple CDW *q*-vectors regime on reduction of Sb content below x < 0.29, a clear magnetic transition featuring a drop in χ_{\perp} below $T_N \approx 2.01$ K for x = 0.10 and $T_N \approx 2.04$ K for x = 0 (NdTe₂) is observed. For $\chi_{//}$, the magnetic transitions in both samples manifest into weak features as shown in the insets in Fig. 2. This susceptibility behavior is distinct than the stronger susceptibility drop observed below T_N in $\chi_{//}$ for Sb-rich compositions and in both $\chi_{//}$ and χ_{\perp} for intermediate Sb-compositions. Such susceptibility behavior showing drop in χ_{\perp} but a weak transition in $\chi_{//}$ below T_N is suggestive of the moment reorientation towards the out-of-plane direction or *c*-axis, consistent with a much larger M_{\perp} than $M_{//}$ in M(H) measurement for x = 0.10 and 0 (Fig. 4). In fact, the switching of magnetic anisotropy from in-plane direction in high-Sb compositions x = 1 and 0.82 to out-of-plane direction in Sb-less samples x = 0.10 and 0 can be directly observed in the $\chi(\theta)$ measurement. As shown in Fig. 3(c), the $\chi(\theta)$ for x = 0 exhibits completely opposite trend in comparison to x = 1 sample with maximum and minimum value along the out-of-plane and in-plane fields respectively.

These results clearly demonstrate the complex interaction between CDW and magnetism proposed in earlier *Ln*SbTe studies [33,62,65], which can also be understood by the evolution of magnetic parameters such as Curie-Weiss temperature (θ_{cw}) and effective magnetic moment (μ_{eff}) [Fig. 3(b)]. These parameters are extracted by fitting the $\chi(T)$ data in the paramagnetic phase using a modified Curie-Weiss model $\chi_{mol} = \chi_0 + C/(T - \theta_{cw})$, where χ_0 is the temperature independent part of susceptibility and *C* is Curie constant. From the fitting, we found negative θ_{cw} for all the samples as expected for their AFM ground state. As shown in Fig. 3(b), the θ_{cw} lacks the systematic dependence on Sb content *x* with significantly different θ_{cw} for two end compounds x = 0 ($\theta_{cw} \approx$ -29.5 K) and 1 ($\theta_{cw} \approx$ -7.8 K), which is distinct than the systematic variation in CeSb_xTe_{2-x-8} [62] and GdSb_xTe_{2-x- δ} [63]. Higher (more negative) θ_{cw} for x = 0 in comparison to x = 1 sample implies stronger AFM interaction in NdTe₂ (x = 0), however its T_N is lower than that of NdSbTe_{1.08} (x = 1). Between these two end compounds, the variation of θ_{cw} that gives the information about magnetic exchange interactions is also unable to explain the evolution of T_N with Sb-Te substitution. Such discrepancy between the composition dependence of θ_{cw} and T_N suggests the additional contribution affecting the formation of long-range AFM ordering in NdSb_xTe_{2-x+ δ}, which has also been proposed in GdSb_xTe_{2-x- δ} [63]. Thus, the role of CDW in tuning magnetism seems plausible in this family.

In addition, from the Curie constant we have obtained the effective moments by μ_{eff} = $\sqrt{\frac{3k_BC}{N_A}}$, where N_A is the Avogadro's number and k_B is the Boltzmann constant. The obtained μ_{eff} also exhibits a non-monotonic dependence on Sb content x with a value of 3.70 $\mu_{\rm B}$ for NdSbTe_{1.08} (x = 1) that is slightly different than the theoretically expected value of 3.62 $\mu_{\rm B}$ [shown by a dashed line in Fig. 3(b)] for a Nd³⁺ ion with a 4f³ configuration. The μ_{eff} is further deviated from the theoretical value with increasing substitution and becomes minimum for x = 0.82 ($\mu_{eff} \approx 2.85 \mu_B$, which is consistent with tetragonal to orthorhombic phase boundary that activates the CDW. On reducing the Sb content to x = 0.60, the μ_{eff} start to approach the theoretical value which is slightly surpassed on further decreasing Sb content to x = 0.45 and reaching a maximum $\mu_{eff} \approx 4.05 \ \mu_{B}$ for x = 0.29 that appears to align with single to multiple CDW *q*-vectors transition below which μ_{eff} decreases to a value of $\mu_{\rm eff} \approx 3.38 \ \mu_{\rm B}$ for x = 0. Such variation of $\mu_{\rm eff}$ also indicates the interplay between CDW and magnetism, which is in stark contrast to $CeSb_xTe_{2-x-\delta}$ [62] and $GdSb_xTe_{2-x-\delta}$ [63] where μ_{eff} has been reported to be close to the theoretical values for Ce³⁺ and Gd³⁺ ions respectively over entire Sb-Te composition. The deviation from theoretical μ_{eff} with Sb-Te substitution in

 $NdSb_{x}Te_{2-x+\delta}$ might be attributed to a few reasons such as varying spin-orbit coupling (SOC) [82], crystal electric field effect (CEF) [83] and/or the hybridization between the 4f moments and conductions electrons [84-87]. The SOC is less likely to play a significant role given the fact that the Sb-Te substitution in CeSb_xTe_{2-x- δ} [62] and GdSb_xTe_{2-x- δ} [63], which would cause a similar variation of SOC, has less effect on μ_{eff} . In addition, the CEF on 4*f* electrons is negligible because they are well-screened by the electrons of 5s and 5p orbitals [88,89]. This implies that the coupling of 4f moments and conductions electrons generated by Sb bands could cause the variation of $\mu_{\rm eff}$ from the theoretical value. This further supports the interplay between magnetism and CDW induced by distorted Sb-square net. Such coupling of magnetism and CDW might also be the origin for the reorientation of Nd moments that leads to the change in magnetic structure with Sb-Te substitution in $NdSb_{x}Te_{2-x+\delta}$, which is again distinct than the similar AFM configuration over an entire composition range in CeSb_xTe_{2-x- δ} [62] and GdSb_xTe_{2-x- δ} [33]. Further neutron scattering experiment on a wide range of $NdSb_{x}Te_{2-x+\delta}$ compositions similar to a recent study on two Ndbased samples NdSb_{0.94}Te_{0.92} and NdSb_{0.48}Te_{1.37} [65] is needed to clarify the interplay between magnetism and CDW as well as the evolution of magnetic structure in $NdSb_xTe_{2-x+\delta}$.

The reorientation of magnetic moments has been proposed to break various symmetries and consequently tune the topological states in AFM TSM [90–92]. For example, in a TSM candidate YbMnSb₂ [90], a *C*-type AFM ordering with out-of-plane or canted moments have been predicted to give rise to a gapped Dirac crossing or Weyl nodes, respectively. Similar modulation of topological states depending on moment orientation has also been predicted [91,92] and experimentally verified [91] in another TSM candidate FeSn. Substituting Co for Fe in FeSn reorient AFM moments from in-plane to out-of-plane direction which breaks the nonsymmorphic symmetry leading to a theoretically predicted gap at the Dirac point [91]. In *Ln*SbTe, the topological band structure can be controlled by Sb-Te substitution, providing access to an ideal Dirac state located near the Fermi level (E_F) for intermediate Sb compositions CeSb_{0.51}Te_{1.40} [62] and GdSb_{0.46}Te_{1.48} [61] where all trivial bands at the E_F are gapped out by CDW. However, as discussed earlier, both CeSb_xTe_{2-x-δ} [62] and GdSb_xTe_{2-x-δ} [33] exhibit similar spin orientation for entire Sb composition. Therefore, NdSb_xTe_{2-x+δ} studied in this work which displays switching of easy axis between in-plane and out-of-plane directions could be an ideal platform to investigate the interplay between moment reorientation and non-trivial band topology.

In addition to symmetry breaking induced by moment reorientation, tuning magnetic states by applying magnetic field has also been proposed to modify the topological phases in CeSbTe [28]. The AFM ground state in CeSbTe is found to exhibit field-driven metamagnetic transition and a subsequent ferromagnetic (FM)-like polarization [28], which provides an approach to switch on/off the time-reversal symmetry and is predicted to tune topological states [28]. Also a field driven moment polarization to FM state has been demonstrated to lead to a topological phase transition from AFM topological insulator to time reversal symmetry-breaking Weyl state in MnBi₂Te₄ [93]. Here, as seen in field-dependence of magnetization M(H) measurements at T = 2K for all NdSb_xTe_{2-x+ δ} samples (Fig. 4), the isothermal magnetization becomes sublinear at high field but lacks a clear saturation behavior seen in true ferromagnets and their values at $\mu_0 H = 9$ T are smaller than the saturation moment of 3.62 μ_B for a Nd³⁺ ion. Therefore, the high-field sublinear magnetization in NdSb_xTe_{2-x+ δ} might be attributed to a possible new canted AFM state with partial polarization of moments. We calculated the derivative of M(H) data $[dM/d(\mu_0 H)]$ to precisely determine the field-driven metamagnetic (MM) transition and partial spin polarization (PP), which was used in an earlier NdSbTe study where a sharp peak followed by a broad hump or shoulder at higher field are defined as metamagnetic transition ($H_{\rm MM}$) and partial spin polarization ($H_{\rm PP}$) fields,

respectively [65]. The field-dependence of $dM/d(\mu_0 H)$ data under H/ab and H/c fields for samples representing three different AFM regions [AFM_{ab}, cAFM, and AFM_c in Fig. 3(a)] showing H_{MM} and H_{PP} are presented in Fig. 5(a). Based on these results, we constructed a magnetic phase diagram at T = 2 K [Fig. 5(b)] which depicts the evolution of magnetic states with Sb content x. First, in AFM_{ab} region, the x = 1 sample exhibits AFM to partial moment polarization featuring sublinear magnetization at higher in-plane magnetic field (H//ab) while the x = 0.82 undergoes AFM to MM transition before partial polarization at H/ab only, consistent with their in-plane moments as discussed earlier. Decreasing Sb content below x < 0.82 systematically reduces both H_{MM} and H_{PP} in cAFM region, however these transitions occur under both in-plane and out-of-plane fields that is line with their canted moments. As mentioned above, further reducing Sb content to x = 0.10and 0 switches easy axis towards the out-of-plane direction and thus features transition from AFM to partial spin polarization (without low-field metamagnetic transition) for H//c field only. This demonstrates rich magnetic phases in $NdSb_{x}Te_{2-x+\delta}$ depending on both Sb composition and applied field, which provides a rare platform to explore coupling between magnetism and electronic band topology.

In conclusion, we have investigated the magnetic properties of NdSb_xTe_{2-x+ δ} over the entire composition range. This work reveals an interesting non-monotonic evolution of T_N accompanied by a systematic reorientation of moments from in-plane to out-of-plane direction with decreasing Sb content *x*. The rich magnetic phases in NdSb_xTe_{2-x+ δ} provide useful insights for the evolution of magnetism in *Ln*SbTe materials, offering a good platform for tunable topological states.

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Table I: Elemental compositions used in the source materials and final compositions in the grown

 crystals determined by EDS.

Source materials			EDS composition	$NdSb_{x}Te_{2-x+\delta}$	
Nd	Sb	Te		x	δ
1	0	2	NdTe ₂	0	0
1	1.2	1	NdSb _{0.10} Te _{1.93}	0.10	0.03
1	0.5	1.5	NdSb _{0.29} Te _{1.73}	0.29	0.02
1	0.2	0.8	NdSb _{0.45} Te _{1.57}	0.45	0.02
1	1.1	0.9	NdSb _{0.60} Te _{1.48}	0.60	0.08
1	0.3	0.7	NdSb _{0.82} Te _{1.22}	0.82	0.04
1	1.2	0.8	NdSb ₁ Te _{1.08}	1	0.08

Figure 1



FIG. 1. (a) Optical microscope images of $NdSb_xTe_{2-x+\delta}$ ($0 \le x \le 1$) crystals. (b) X-ray diffraction result for $NdSb_xTe_{2-x+\delta}$. (c) Evolution of lattice parameters *a*, *b* (Left vertical axis), and *c* (Right vertical axis) with varying Sb content *x*. The Blue, orange, and grey regions represent tetragonal (Tetr), orthorhombic (Orth), and tetragonal (Tetr) lattices, respectively.





FIG. 2. Temperature dependence of susceptibility for $NdSb_xTe_{2-x+\delta}$ samples under in-plane (H||ab, magenta) and out-of-plane (H//c, blue) magnetic fields of $\mu_0H = 0.1$ T. Inset: Low-temperature susceptibility under in-plane H||ab field to show magnetic transition clearly. The dashed lines are guide to eye. (b) Temperature dependence of derivative $d\chi/dT$ of $NdSb_xTe_{2-x+\delta}$ samples. The solid triangles denote T_N .

Figure 3



FIG. 3. (a) Evolution of T_N with varying Sb content x. Different colored regions denote distinct orientations for Nd moments. The in-plane, canted, and out-of-plane antiferromagnetic configurations are denoted as AFM_{ab}, cAFM, and AFM_c whereas PM represents a paramagnetic state. The moment orientations in each phase region are schematic to show the magnetic easy axis. Here, χ_{ab} and χ_c represent the susceptibility along the *ab*-plane and *c*-axis, respectively. (b) Evolution of effective magnetic moment μ_{eff} and Curie-Weiss temperature θ_{CW} with varying Sb content x. The dashed line represents the theoretical μ_{eff} . (c) Angular dependence of susceptibility for x = 0, 0.60, and 1 sample under magnetic fields of $\mu_0 H = 0.1$ T and T = 2 K.





FIG. 4. Field dependence of magnetization for $NdSb_xTe_{2-x+\delta}$ samples under in-plane (H||ab, magenta) and out-of-plane (H//c, blue) magnetic fields at T = 2 K.





FIG. 5. (a) Field dependence of derivative $dM/d(\mu_0 H)$ under in-plane (H||ab, magenta) and outof-plane (H//c, blue) magnetic fields at T = 2 K. The metamagnetic and partial spin polarization fields are represented as H_{MM} (Blue color) and H_{PP} (Red color), respectively. (b) Magnetic phase diagram constructed from the field dependence of magnetization measurements presented in Fig. 3. The in-plane, canted, and out-of-plane antiferromagnetic configurations are denoted as AFM_{ab}, cAFM, and AFM_c. The metamagnetic transition for in-plane and canted antiferromagnetic states are represented as MM_{ab} and MM_{cAFM}, respectively. The partial spin polarization is denoted as PP.