First-principles study of magnetic states and the anomalous Hall conductivity of $M$-Nb$_2$S$_6$ ($M$=Co, Fe, Mn, and Ni)

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(Dated: February 10, 2022)

Inspired by the observation of the extremely large anomalous Hall effect in the absence of applied magnetic fields or uniform magnetization in CoNb$_2$S$_6$ [Nature Comm. 9, 3280 (2018); Phys. Rev. Research 2, 023051 (2020)], we perform a first-principles study of this and related compounds of the $M$-Nb$_2$S$_6$ type with different transition metal $M$ ions to determine their magnetic orders and the anomalous Hall conductivity (AHC). We find that non-coplanar antiferromagnetic ordering is favored relative to collinear or coplanar order in the case of $M$=Co, Fe and Ni, while ferromagnetic ordering is favored in Mn$_2$B$_6$ at low temperatures. The AHC in these materials with non-coplanar spin ordering can reach about $e^2/h$ per crystalline layer, while being negligible for coplanar and collinear cases. We also find that the AHC depends sensitively on doping and reaches a maximum for intermediate values of the local spin exchange potential between 0.3 and 0.8 eV. Our AHC results are consistent with the reported Hall measurements in CoNb$_2$S$_6$ and suggest a possibility of similarly large anomalous Hall effects in related compounds.

I. INTRODUCTION

Despite having a simple semiclassical explanation, the Hall effect [1] is a remarkable quantum phenomenon that is caused by a subtle influence of the electromagnetic vector potential on the quantum mechanical phase of electrons. In weak magnetic fields, the transverse Hall voltage, being determined by the Lorenz force, is proportional both to the electric current and the magnetic field. However, in the limit of strong magnetic fields, deviations from this simple dependence become prominent, culminating in two-dimensional systems as quantized values of Hall conductance [2], precise enough to serve as a universal standard [3].

Reaching quantized levels of the Hall conductance in the original experiments [2, 4] required a combination of low carrier density and high magnetic fields. The reason is that the quantum Hall effect occurs when the electronic density $n_{2d}$ and quantum flux density, $n_{q} = B/e\phi_0 = 2eB/h$ are comparable. Thus, reaching the quantized limit in conventional materials with their high carrier density would require extremely high magnetic fields, 10$^4$ T and above. This might seem to imply that bulk materials are destined to remain in the semiclassical – weak-field – limit of the Hall effect. This is however not true, and the reason is that electrons in solids experience additional interactions that can have a dramatic effect on their quantum mechanical phase. These are exchange interactions with magnetic moments of atoms and spin-orbit interactions. Both can influence the Bloch wave functions of electrons, resulting in nontrivial features in bands structures, leading to nonzero Hall effect even in the absence of external magnetic field [5-9]. A classic example is the anomalous Hall effect (AHE) in ferromagnetic (FM) iron [10, 11], where magnetic ordering with the help of spin-orbit interaction can deflect moving electrons in a way similar to the conventional orbital magnetic field. A more exotic possibility, which does not require explicit spin-orbit coupling, relies on the interaction between spins of itinerant electrons and magnetic moments of atoms ordered into non-coplanar states [12–15]. In this case, the Berry phase imprinted on electrons can reach values that are equivalent to having a flux quantum piercing the magnetic unit cell, which translates into an extremely large effective magnetic field. Notably, the magnetic ordering itself can be purely antiferromagnetic (AFM), that is, lacking any net magnetic moment. Moreover, the AHE may persist even if there is no static magnetic order, but only finite scalar chirality [15, 16].

In recent years, several materials have been experimentally discovered that show complex AFM ordering and a very large AHC in the absence of magnetic fields or uniform magnetization [17–24]. Despite these successes, observing a quantized AHC in these systems has remained elusive. Even if a system has topologically non-trivial bands that could lead to quantization [25], accidental doping or magnetic domain structure may preclude its experimental observation [26].

A system that is perhaps one of the most promising candidates to exhibit a quantized AHC is CoNb$_2$S$_6$. It has the layered hexagonal structure of NbS$_2$, intercalated by triangular layers of Co [27]. Based on early neutron scattering data, the magnetic structure was believed to be a simple collinear AFM [28] (see Fig. 1f,g). So it came as a big surprise when an extremely large AHE, with the Hall conductivity comparable to $e^2/h$ per structural layer, was recently discovered in this system [21, 26]. Indeed, collinear AFMs are invariant under a combination of lattice translation and time-reversal; however, since time-reversal flips the sign of Hall effect and translation leaves it unchanged, the AHE must vanish and, therefore, such AFM order cannot be responsible for the observed anomalous Hall response. More recently, neutron scattering experiments were performed [29] on the new samples of CoNb$_2$S$_6$ and the results did not exclude more complex magnetic orderings, in addition to the collinear AFM.
The early neutron scattering measurement by Parkin et al. [28] reported the primary peak positions of CoNb$_3$S$_6$ at the M points of the Brillouin zone and their results can be interpreted as either a multi-domain 1q spin ordering with all three possible directions of q present, or a single-domain 3q spin ordering. Distinguishing between these two possibilities require detailed analysis of higher-order scattering peaks that had not been done at the time. Although it is difficult to identify the magnetic order in these compounds experimentally and noting that a non-zero AHE could be possible even without any static magnetic orders, we focus here on the possibility of the AHE in CoNb$_3$S$_6$ and related compounds originating from static non-coplanar magnetic orders. Such orders can be studied using first-principles calculations. One possibility that was not ruled out by the experiment is a "tetrahedral" non-coplanar AFM state. In a given triangular Co plane, the ordering simultaneously involves three symmetry-related AFM wavevectors, in contrast to the simple collinear AFM state, which only has one. The tetrahedral order has four spins per magnetic unit cell, pointing towards corners of a regular tetrahedron (see Fig. 1j-k). This state was theoretically proposed as a non-coplanar AFM order. Distinguishing between these two possibilities require detailed analysis of higher-order scattering peaks that had not been done at the time. For energetics and band structure calculations of MNB$_3$S$_6$, we use DFT by adopting the VASP code. For the Berry curvature and AHC calculations in a dense $k$-mesh, we construct the spin-resolved real-space Hamiltonian by interpolating band structures obtained from first-principles:

$$\hat{H} = \sum_{\alpha \beta \sigma, R, R'} t_{\alpha \beta}(R - R') \hat{c}^\dagger_{R \alpha \sigma} \hat{c}_{R' \beta \sigma} + \sum_{R, \alpha i} \hat{H}_U(r_i),$$

where $t_{\alpha \beta}(R - R')$ is the spin-independent hopping integral between the Wannier function $\alpha$ within the unit cell $R$ and the Wannier function $\beta$ within the unit cell at $R'$. The $t_{\alpha \beta}(R - R')$ parameters are calculated from the non-magnetic Kohn-Sham Hamiltonian in the basis of the maximally localized Wannier functions obtained using the Wannier90 code [34, 35].

For the spin-dependent part of the Hamiltonian, we adopt multi-orbital Hubbard interaction correction term $\hat{H}_U$ for $d$ orbitals of every $M$ ion located at site $r_i$:

$$\hat{H}_U(r_i) = \sum_{\alpha \beta \gamma \delta \sigma \sigma'} U_{\alpha \beta \gamma \delta} \hat{c}^\dagger_{\alpha \sigma} \hat{c}_{\beta \sigma'} \hat{c}_{\gamma \sigma} \hat{c}_{\delta \sigma'}$$

$$\approx \sum_{\alpha \beta} \left( \hat{U}_H \tilde{n}_\alpha^i \tilde{n}_\beta^i + \hat{U}_F \tilde{S}_\alpha^i \cdot \tilde{S}_\beta^i \right)$$

where $U_{\alpha \beta \gamma \delta}$ is the on-site Coulomb interaction matrix between orbitals $\alpha, \beta, \gamma, \delta$ centered at the magnetic site $r_i$, $\hat{U}_H$ is the direct Coulomb interaction parameter for the density-density type interaction, $\hat{U}_F$ is the Hund’s coupling parameter for the spin-spin interaction, $\tilde{n}_\alpha^i$ is the density operator for the orbital $\alpha$ of magnetic atom $i$, and $\tilde{S}_\alpha^i = \sum_{\sigma'} \hat{c}_{\alpha \sigma'}^i \hat{c}_{\alpha \sigma'} \hat{c}_{\alpha \sigma'}$ is the local spin operator. Both $\hat{U}_H$ and $\hat{U}_F$ are taken averaged over orbital indices at the local site. In the mean-field approximation, $\hat{H}_U$ can be written in terms of the one-particle Hamiltonian,

$$\hat{H}_U(r_i) \approx \sum_{\sigma} \left( V_{\sigma} \cdot \delta_{\sigma \sigma'} + J_{\sigma} \cdot \sigma_{\sigma'} \right) \hat{c}_{\sigma}^i \hat{c}_{\sigma'}^i$$
where $V^i \simeq \sum_\beta \tilde{U}_H (\tilde{n}_\beta^i)$ and $J^i \simeq \sum_\beta \tilde{U}_F (\tilde{S}_\beta^i)$. The local spin exchange potential $J^i$ induces a spin polarization at the atomic site $i$ of the $M$ ion; within mean field, it is proportional to the local ordered magnetic moment itself.

For the magnetic states that we consider, symmetry dictates that $V^i$ is site-independent ($V_i = V$), and $J^i$ is site-dependent, but has constant magnitude, i.e., $|J^i| = J$. Since both $V$ and $J$ may differ for different $M$ ions, we determine them by fitting the band structure of Eqs. (1-3) to the one obtained with the spin-resolved DFT using the PBE exchange-correlation functional without spin-orbit coupling (see Fig. 3), assuming the same magnetically ordered states. This allows us to construct a compact representation of band structure that accounts for different stable magnetic orderings obtained within DFT. This band structure is used to efficiently calculate the AHC. In addition to needing a quantitatively accurate and compact band structure, the $V$ and $J$ terms in the tight-binding Hamiltonian can parametrize the Coulomb interaction effects of spin-polarized DFT bands starting with non-spin-polarized DFT bands. We find that both interaction ($V$ and $J$) terms are necessary to capture the exact spin-polarized DFT bands although they can not treat orbital-dependent interactions due to the approximation used in Eq. 3. Although most calculations in this paper are performed using spin-polarized DFT without tuning interaction parameters, we check the correlation effect on the AHC calculation by tuning the $J$ parameter as shown in Fig. 6.

The AHC $\sigma_{xy}$ is computed using the Berry curvature obtained from band structure as follows:

$$\sigma_{xy} = -\frac{e^2}{h} \frac{1}{2\pi} \sum_n \int dk \ n_F(e_{nk}) \cdot \Omega^x_n(k) \tag{4}$$

where $\Omega^x_n(k) = -Im(\partial_x u_{nk}\partial_y u_{nk})$ is the Berry curvature of band $n$ with momentum $k$ and $n_F$ is the Fermi function. $u_{nk}$ is the periodic part of the Bloch wavefunction obtained by solving the spin-resolved Hamiltonian in Eq. (1). The calculation of Eq. (4) requires integration over a very fine $k$-mesh as an important large contribution of $\Omega_n^x$ can occur in a small region of the Brillouin zone (B.Z.). Here, we adopt the Wannier-berry package [36] using the recursive adaptive refinement method based on symmetries for the smooth convergence of the $k$-mesh integration. For the AHC calculations of $M$Nb$_3$S$_6$ compounds in the non-collinear spin configurations, we used the $10 \times 10 \times 8$ $k$-mesh with 10 recursive refinement iterations and temperature smoothing of the Fermi functions at 10 K which is well below the experimental Neel temperature of 29 K in the case of $M = \text{Co}$.

III. RESULTS AND DISCUSSIONS

A. Magnetic structures and moments in $M$Nb$_3$S$_6$

In $M$Nb$_3$S$_6$ compounds, transition metal $M$ ions are intercalated between NbS$_2$ layers, forming triangular lattices stacked along the $c$-axis at two distinct Nb sites locations in an alternating fashion (see Fig 1a,b). The issue of the magnetic ground-state in $M$Nb$_3$S$_6$ has not been settled yet experimentally, and we therefore consider a variety of collinear and non-collinear spin configurations of the $M$ ions (see Fig 1d-k). Some of them have the magnetic unit cell identical to the primitive unit cell (d-e); some double unit cell (f-g), and some – quadruple (h-k). The magnetic moment and the energy for each compound and spin structure are computed using DFT.

All experimentally relevant spin structures can be obtained from the following ansatz for the three-
The simplest collinear cases are c-FM in which all \( M \) spins are aligned ferromagnetically (see Fig. 1d) and c-AFM0 where \( M \) spin directions are FM in the \( a-b \) plane but anti-parallel between two crystallographically inequivalent layers (see Fig. 1e). In both cases, the magnetic unit cell coincides with the primitive unit cell – all \( q_i = 0 \). We note that in the crystal structure of \( \text{M} \text{Nb}_3\text{S}_6 \), the inversion symmetry is broken; this allows for nonzero Dzyaloshinskii-Moriya interaction that tends to favor in-plane alignment of spins [38].

In c-AFM1 and c-AFM2 cases, spins are ferromagnetically ordered along one of the in-plane Co-Co bond directions and modulated antiferromagnetically along the other in-plane Co-Co bond directions. In this so-called 1q structure, the ordering wavevector is half of the in-plane reciprocal vector (a high-symmetry point \( M \) in the primitive Brillouin zone), which is orthogonal to the ferromagnetically ordered direction. In the 1q structure, there are two distinct relative spin arrangements of \( M \) spins in the adjacent layers, one favors AF interlayer coupling (see Fig. 1f) and the other FM interlayer coupling (see Fig. 1g).

In the non-collinear cases, we consider the magnetic unit cell containing four distinct spin orientations per layer (see Fig. 1b dashed line) These orders involve two or three ordering wave vectors that correspond to \( M \) points in the primitive Brillouin zone. The coplanar ordering (cpl-) involves two ordering vectors ("2q" structure) and spins that lie in a single plane in the spin space; their sum is zero, so they can be arranged into a rhombus (see Fig. 1h-i).

The non-coplanar (ncpl-) AFM involves all three M-point wave-vectors ("3q" structure). The spin orientations correspond to the directions from the center toward vertices of a regular tetrahedron [14]. The 3q state on a triangular lattice is special as it corresponds to a scalar chirality, \( \chi_{ijk} = S_i \cdot [S_j \times S_k] \), that is constant for all elementary triangular plaquettes \( (i, j, k) \). For two dimensional systems this leads to Anomalous Hall effect even in the absence of spin-orbit interactions [14, 16]. In the bilayer structure of \( \text{M} \text{Nb}_3\text{S}_6 \), the two \( M \) layers can have either the same or the opposite scalar chirality. Only in the former case we anticipate AHC to be present, since in the latter case layer translation followed by time reversal is a symmetry that prohibits finite AHC. In our DFT calculations we compute the energies of both states (ncpl-AFM1, see Fig. 1j and ncpl-AFM2, see Fig. 1k).

We finally note that all the AF states that we consider can be smoothly distorted into each other: 3q state can be “flattened” into a 2q state by decreasing, e.g., the \( z \) component of spin (coefficient \( C \) in Eq. (5)); further, the 2q structure can be distorted into 1q by reducing one of the remaining spin components, e.g. \( y \) (coefficient \( B \)), to zero. In this work we do not exhaust all the possible ordered states. Instead, we compare the most symmetric states, where nonzero amplitudes \( A, B, C \) are all equal in magnitude.

| Mag. mom. & energy \( M=\text{Mn} \) | \( M=\text{Fe} \) | \( M=\text{Co} \) | \( M=\text{Ni} \) |
|---|---|---|---|
| \( m/m_0 \) [\( \mu_B \)] | 3.9/5.0 | 3.1/4.0 | 1.5/3.0 | 0.7/2.0 |
| c-FM [eV] | -74.900 | -73.412 | -72.211 | -69.734 |
| c-AFM0 [eV] | -74.869 | -73.423 | -72.221 | -69.743 |
| c-AFM1 [eV] | -74.814 | -73.425 | -72.242 | -70.576 |
| c-AFM2 [eV] | -74.811 | -73.404 | -72.222 | -70.588 |
| cpl-AFM1 [eV] | -74.816 | -73.423 | -72.244 | -70.576 |
| cpl-AFM2 [eV] | -74.811 | -73.410 | -72.232 | -70.589 |
| ncpl-AFM1 [eV] | -74.816 | -73.426 | -72.245 | -70.576 |
| ncpl-AFM2 [eV] | -74.811 | -73.408 | -72.224 | -70.589 |

Table I shows spin magnetic moments and total energies per formula unit of the \( M \text{Nb}_3\text{S}_6 \) comparing collinear (c-) FM & AFM states and non-collinear coplanar (cpl-) & non-coplanar (ncpl-) spin states computed using DFT. The ordered magnetic moments are almost the same for all magnetic states; the spin magnetic moments (\( m_0 \)) for the free \( M^{2+} \) ions are also given. Bolded values correspond to the lowest energy states.
spin-orbit coupling in these compounds by comparing the collinear magnetic energies along various ordered spin directions. Regardless of the spin directions, energies of all compounds are almost the same (within 1meV) except FeNb$_3$S$_6$ for which the energy of spin direction along the easy plane was lower than one along the $z$-axis by $\sim$18meV per formula unit. Given the small difference of $\sim$1meV between collinear and non-collinear spin states, it is important to include the spin-orbit effect for the energy calculation in FeNb$_3$S$_6$. The spin-orbit effect is much smaller for other compounds, which is typically expected in these transition metal ions with 3$d$ orbitals.

All MNB$_3$S$_6$ compounds studied here favor the ncl-AFM structures energetically except the $M = \text{Mn}$ case (see Table I). We find that the variant with Mn orders ferromagnetically. Qualitatively, this is consistent with the Mn magnetic moment being the largest, and thus most strongly coupled to itinerant electrons; this favors ferromagnetic ordering via the conventional double-exchange mechanism. Indeed, the magnetic ground state exhibits metallic behavior with large spins of the intercalated Mn ions strongly hybridized with itinerant Nb $d_z^2$ orbitals in the NbS$_2$ layers. Due to strong hybridization, any misalignment of ordered moments will frustrate electronic kinetic energy, and therefore is energetically penalized. The next energy state is c-AFM0, with ferromagnetic Mn planes stacked antiferromagnetically along the $c$ axis. This is consistent with having dominant ferromagnetic coupling between Mn in individual layers, and a weaker ferromagnetic coupling between them.

On the other hand, for smaller moments and thus weaker exchange fields, the physics is expected to be controlled by Fermi surface instabilities. Given the hexagonal symmetry of the crystal, having a magnetic order with multiple ordering wave vectors then becomes favorable as it allows us to gap larger sections of the Fermi surface. The 3$q$ ncl-AFM state is particularly attractive: At a commensurate carrier density, such weak-to-intermediate coupling instability can fully gap the Fermi surface, producing an insulator with quantized AHC [14, 39]. Indeed, the lowest energy spin configuration that we were able to obtain for MNB$_3$S$_6$ for $M = \text{Fe, Co, and Ni}$ correspond to the 3$q$ state; it is ncl-AFM1 for $M = \text{Fe, Co and ncl-AFM2 for } M = \text{Ni}$. The difference between these noncoplanar states is that one has the same sign of scalar spin chirality in both $M$ layers within the unit cell (ncl-AFM1), while the other (ncl-AFM2) has opposite in sign chiralities in two layers. Because of the scalar chirality structure, we expect non-zero net AHC for Fe and Co cases, and zero AHC for Ni case. This is indeed confirmed numerically in the later subsection (see Fig. 5).

It is useful to also examine the DFT results in Table I for other states besides the lowest energy ones. From the magnetic structure, Fig. 1, the (c.pl.ncl)-AFM1 are favored by the antiferromagnetic nearest neighbor interlayer coupling, while (c.pl.ncl)-AFM2 are by the ferromagnetic one. The pattern of energies is consistent with this coupling determining the type of the ordered states: For a given material, all x-AFM1 states are consistently above or below the corresponding x-AFM2. We also notice that coplanar and non-coplanar states are extremely close in energy, so it is conceivable that in the real materials the energy balance could be tipped in the opposite direction.

### B. Electronic structure of MNB$_3$S$_6$

The DFT calculations described above provide us with the candidate magnetic ground states, and we anticipate that ncl-AF1 is likely to have significant AHC. In order to compute the Hall conductivity we need both the wave functions and energies of electrons with high momentum resolution. This is very computationally expensive within DFT, particularly so if the magnetic unit cell is enlarged and magnetization is noncollinear.

To reduce the computational cost, we construct effective tight-binding models for the materials of interest by fitting the relevant bands using Wannier interpolation technique [34]. We do this in two steps. First, we construct the real-space Hamiltonian (Eq. (1)) with the hopping parameters ($t_{\alpha\beta}$) between Wannier orbitals in the paramagnetic state obtained with the Wannier90 code [35]. As can be seen from the upper panels of Fig. 3, the matching between the Wannier bands and the nonmagnetic DFT bands is almost perfect. We then treat the local direct interaction ($V$) and the spin exchange ($J$) terms on the transition metal $M$ as free parameter to fit the band structure in magnetically ordered states. We use the c-AFM0 band structure computed without SOC since it is the simplest AFM structure that has the same unit cell as the crystal itself.

![FIG. 2. Density of states in CoNb$_3$S$_6$ projected to Co, Nb, and S ions computed using non-magnetic DFT.](image)

Fig. 2 shows the density of states for CoNb$_3$S$_6$ projected to different ions. The Co $d$ orbitals are concentrated near the Fermi energy in the range between $-2$ eV and 1 eV, and they are also strongly hybridized with
the itinerant Nb $d_{z^2}$ orbital. Other Nb $d$ orbitals are located at higher energies, over 2 eV above the Fermi level. The hybridization with the S ions occurs at much lower energies, below $-2$ eV. For other transition metals $M$, the orbital arrangement and energy scales are similar. Therefore, we use Nb $d_{z^2}$ and $M$ ion’s $d$ Wannier orbitals for constructing the non-magnetic Hamiltonian (Eq. (1)); $\hat{H}_{\text{DFT}} = 0$).

Fig. 3 top panel compares the obtained Wannier bands (red dashed lines) to non-magnetic DFT band results (green solid lines) along the high-symmetry directions in the B.Z. of the primitive cell (see Fig. 1c black line B.Z.). The Wannier bands obtained from the non-magnetic Hamiltonian in Eq. (1) reproduce the DFT bands almost exactly for all $M$Nb$_3$S$_6$ compounds. As the $M$ ion changes from Mn to Ni, the occupancy of $d$ orbitals increases from $d^8$ to $d^{10}$ and bands with $M$ ion character shift down in energy. This change is particularly noticeable by following the band evolution near the $k_z = \pi$ plane in the momentum space ($A-L-H-A$ line in Fig. 3).

To be able to construct the band structure of the magnetically ordered states, we introduce $V$ and $J$ parameters in Eq. (3). The $J$ field is an effective spin-exchange potential, which is proportional to the magnetic moments of the $M$ site; it is therefore site-dependent due to the varying spin directions in the AFM state. The $V$ parameter is the local effective potential for the density-density interaction and it determines the relative position of $M$ ion bands compared to the Nb $d_{z^2}$ band; it is site-independent. Figure 3 bottom panels compare the spin-polarized band structure obtained from the Wannier Hamiltonian (dashed lines) to spin-resolved DFT (solid lines) calculations for the c-AFM0 state (collinear state with FM planes of $M$, stacked antiferromagnetically along the c axis). As seen in Fig. 3, including $J$ changes the band structure noticeably relative to the non-magnetic bands except for $M = Ni$. Focusing on the states nearest to the Fermi level, we find the following parameters that allow an accurate fit of Wannier bands to DFT ($J = |J|$, which is the same on all magnetic sites in the states that we consider): $J = 1.3$ eV and $V = -0.4$ eV for Mn, $J = 0.8$ eV and $V = -0.5$ eV for Fe, $J = 0.5$ eV and $V = -0.1$ eV for Co, and $J = 0.2$ eV and $V = 0$ eV for Ni. Since the spin-exchange potential $J$ is the product of the Hund’s coupling $\tilde{U}$ and the magnitude of the ordered moment, and since the Hund’s coupling does not vary much with ion $M$, the correlation between the value of $J$ and the size of the ordered moments is expected. Indeed, Mn has the largest moment, and, predictably, has the largest value of $J$.

C. Results of AHC calculations

In this subsection, we compute the AHC of $M$Nb$_3$S$_6$ with $M = Mn$, Co, Fe, and Ni. In most of the states that we consider, there are symmetry reasons that ensure that AHC must be zero. In particular, c-AFM, copl-AFM, and ncpl-AFM2 magnetic structures in $M$Nb$_3$S$_6$ remain unchanged under application of the time-reversal symmetry followed by the spatial translation of the lattice. For instance, in the cases of collinear and coplanar antiferromagnetic states, the relevant spatial translation is the one that connects two nearest $M$ sites with the opposite spin orientations. On the other hand, noncoplanar
AFM states may or may not break this symmetry, depending on the relative sign of scalar spin chirality in the two magnetic layers [13, 14].

Figure 5 displays the AHC results as a function of the Fermi energy $E_F$ computed for $M$Nb$_3$S$_6$ compounds with the non-coplanar AFM spin texture. For that, we assume rigid bands and vary their occupancy. This allows us to see how sensitive our results are to the precise band alignment, which may not be perfectly captured by DFT, but also possible sensitivity to chemical or electrostatic gate doping. Consistently with our expectations, out of non-coplanar states ncpl-AFM1 state has finite and large AHC, while ncpl-AFM2 does not.

In the four compounds we studied, the Co and Fe variants show the largest AHC when the Fermi energy is equal to zero, which is the nominal value for charge-neutral systems. Still, the AHC magnitude depends rather sensitively on the Fermi energy, i.e., the electron filling. We also compute AHC in ncpl-AF1 and ncpl-AF2 states, even though from DFT calculations (Table I) we expect the Mn compound to be FM. Also, in the case of Ni the ground state is predicted to be ncpl-AF2, which should have no AHC. The differences of DFT energies are rather small, however, so we cannot rule out that the energy ordering of candidate states can deviate from the DFT predictions.

The sensitivity of the AHC values to the Fermi energy implies the importance of band crossings near the Fermi level. In CoNb$_3$S$_6$, the AHC value is $\sim 1.2 e^2/h$ without dopings and remains almost flat under electron doping (increasing the Fermi energy). This can be attributed to the major contribution to the Berry curvature $\Omega(k)$ in CoNb$_3$S$_6$ coming from the vicinity of the high-symmetry point $K$; there are no low-energy bands above the Fermi energy at $K$ point, which makes that contribution insensitive to the Fermi energy. On the other hand, the hole doping leads to visible reduction of the AHC values.

Typically, in calculations of the Berry curvature $\Omega$, the major contributions to $\Omega$ originate from a small momentum space region. To identify this region, we compute $\Omega^z(k)$ summed over occupied bands along the high-symmetry directions in Fig. 4. The sharp peak and valley structures indeed occur in small regions, but these regions vary depending on the specific transition metal ion $M$. We also note that AHC is mostly coming from the vicinity of $k_z = 0$ plane rather than the $k_z = \pi$ plane. This can be understood from band structure since the spin-polarized bands are still nearly degenerate along the path in the $k_z = \pi$ plane while the effect of symmetry breaking occurs more prominently near the $k_z = 0$ plane with avoided crossings along the high-symmetry $k$–path.

Finally, we verify that our AHC results are not very sensitive to the values of $J$ that we used to fit the magnetic band structure. In Fig. 6, we computed the AHC as a function of $J$ in the chiral ncpl-AFM1 state (recall that based on DFT, only Co and Fe compounds are expected to have this ground state). In all cases, sizable AHC values are obtained when $J$ is in the intermediate range $0.3 \text{eV} < J < 0.8 \text{eV}$. The zero-temperature values of $J$ for Co and Fe indeed fall within this range, while it is rather smaller in the case of Ni ($J = 0.2 \text{eV}$). On the other hand, Mn has very large exchange coupling ($J = 1.3 \text{eV}$). We recall, however, that the quoted values of $J$ include in themselves the ordered magnetic moment, and therefore the magnitude of $J$ is generally expected to decrease with increasing temperature. This opens an interesting possibility, in the case of the Mn-intercalated compound in particular, that despite the low-temperature state being
In this work, we searched among collinear and non-collinear magnetic configurations for the ground states of $M$Nb$_3$S$_6$ compounds with $M$ = Mn, Fe, Co, and Ni. In these materials, the magnetic ions $M$ are far apart and their interactions are predominantly mediated by itinerant electrons. This leads to long-range frustrated exchange interactions, as well as higher order – multi-spin – interactions. Even weak higher order interactions generated this way can lift the massive degeneracies common in Heisenberg models, leading sometimes to exotic non-coplanar states instead of the more common coplanar helical states [40, 41]. Indeed, we found that for FeNb$_3$S$_6$ and CoNb$_3$S$_6$, a non-coplanar AFM structure with uniform scalar spin chirality has the lowest energy among the plausible candidate states that we considered. In the case of Ni, the lowest energy state is also non-coplanar AFM, but with staggered scalar spin chirality. In contrast, the collinear FM was found to be the ground-state in MnNb$_3$S$_6$. We also performed the anomalous Hall conductivity calculations using the band structure obtained from the Wannier Hamiltonian fitted to magnetically ordered states without the spin-orbit coupling.

The obtained results show the AHC on the scale of $\sim e^2/h$ per NbS$_2$ layer, which is comparable with the experimentally measured AHC values in CoNb$_3$S$_6$. The calculated AHC depends rather sensitively on the chemical potential, indicating that doping may shift the bands near the Fermi energy and significantly affect the AHC values. We also studied the effect of a local spin exchange field $J$ on the AHC calculation. We found that the intermediate values of $|J|$ produce the largest AHC. This is indeed the case for both Co and Fe ions. The $|J|$ value obtained from DFT for Ni ion is smaller and thus likely to produce weaker AHC even if ordered magnetically with uniform scalar chirality.

Given the expected small energy differences between coplanar and non-coplanar states, it is possible that their energies can be relatively rearranged in experiment by, e.g., quantum confinement (exfoliation), applying various strain fields, or by doping. Moreover, application of an external magnetic field, via coupling to orbital magnetization [42], can both align chiral domains in the ncpl-AFM1 state (predicted for $M$ = Co and Fe), and even induce transitions from ncpl-AFM2 (Ni case) to ncpl-AFM1, with a concomitant jump in AHC.

**IV. CONCLUSION**

In this work, we searched among collinear and non-collinear magnetic configurations for the ground states of $M$Nb$_3$S$_6$ compounds with $M$ = Mn, Fe, Co, and Ni. In these materials, the magnetic ions $M$ are far apart and their interactions are predominantly mediated by itinerant electrons. This leads to long-range frustrated exchange interactions, as well as higher order – multi-spin – interactions. Even weak higher order interactions generated this way can lift the massive degeneracies common in Heisenberg models, leading sometimes to exotic non-coplanar states instead of the more common coplanar helical states [40, 41]. Indeed, we found that for FeNb$_3$S$_6$ and CoNb$_3$S$_6$, a non-coplanar AFM structure with uniform scalar spin chirality has the lowest energy among the plausible candidate states that we considered. In the case of Ni, the lowest energy state is also non-coplanar AFM, but with staggered scalar spin chirality. In contrast, the collinear FM was found to be the ground-state in MnNb$_3$S$_6$. We also performed the anomalous Hall conductivity calculations using the band structure obtained from the Wannier Hamiltonian fitted to magnetically ordered states without the spin-orbit coupling.

The obtained results show the AHC on the scale of $\sim e^2/h$ per NbS$_2$ layer, which is comparable with the experimentally measured AHC values in CoNb$_3$S$_6$. The calculated AHC depends rather sensitively on the chemical potential, indicating that doping may shift the bands near the Fermi energy and significantly affect the AHC values. We also studied the effect of a local spin exchange field $J$ on the AHC calculation. We found that the intermediate values of $|J|$ produce the largest AHC. This is indeed the case for both Co and Fe ions. The $|J|$ value obtained from DFT for Ni ion is smaller and thus likely to produce weaker AHC even if ordered magnetically with uniform scalar chirality.

Given the expected small energy differences between coplanar and non-coplanar states, it is possible that their energies can be relatively rearranged in experiment by, e.g., quantum confinement (exfoliation), applying various strain fields, or by doping. Moreover, application of an external magnetic field, via coupling to orbital magnetization [42], can both align chiral domains in the ncpl-AFM1 state (predicted for $M$ = Co and Fe), and even induce transitions from ncpl-AFM2 (Ni case) to ncpl-AFM1, with a concomitant jump in AHC.

**ACKNOWLEDGMENTS**

We thank M. Norman for helpful and insightful discussions. HP, OH, and IM acknowledge funding from the US Department of Energy, Office of Science, Basic Energy Sciences, Division of Materials Sciences and Engineering. We gratefully acknowledge the computing resources provided on Bebop, high-performance computing clusters operated by the Laboratory Computing Resource Center.
at Argonne National Laboratory.

[1] E. H. Hall, American Journal of Mathematics 2, 287 (1879).
[2] K. v. Klitzing, G. Dorda, and M. Pepper, Phys. Rev. Lett. 45, 494 (1980).
[3] B. Jeckelmann and B. Jeanneret, Reports on Progress in Physics 64, 1603 (2001).
[4] M. Paalanen, D. Tsui, and A. Gossard, Physical Review B 25, 5566 (1982).
[5] M.-C. Chang and Q. Niu, Physical review letters 75, 1348 (1995).
[6] M.-C. Chang and Q. Niu, Physical Review B 53, 7010 (1996).
[7] G. Sundaram and Q. Niu, Physical Review B 59, 14915 (1999).
[8] N. Nagaosa, J. Sinova, S. Onoda, A. H. MacDonald, and N. P. Ong, Rev. Mod. Phys. 82, 1539 (2010).
[9] D. Xiao, M.-C. Chang, and Q. Niu, Rev. Mod. Phys. 82, 1959 (2010).
[10] J. Kondo, Progress of Theoretical Physics 27, 772 (1962).
[11] Y. Yao, L. Kleinman, A. H. MacDonald, Jin Sino, T. Jungwirth, D.-s. Wang, E. Wang, and Q. Niu, Phys. Rev. Lett. 92, 037204 (2004).
[12] Y. Taguchi, Y. Oohara, H. Yoshizawa, N. Nagaosa, and Y. Tokura, Science 291, 2573 (2001).
[13] R. Shindou and N. Nagaosa, Phys. Rev. Lett. 87, 116801 (2001).
[14] I. Martin and C. D. Batista, Phys. Rev. Lett. 101, 156402 (2008).
[15] Y. Machida, S. Nakatsuji, S. Onoda, T. Tayama, and T. Sakakibara, Nature 463, 210 (2010).
[16] Y. Kato, I. Martin, and C. D. Batista, Phys. Rev. Lett. 105, 266405 (2010).
[17] S. Nakatsuji, N. Kiyohara, and T. Higo, Nature 527, 212 (2015).
[18] A. K. Nayak, J. E. Fischer, Y. Sun, B. Yan, J. Karel, A. C. Komarek, C. Shekhar, N. Kumar, W. Schnelle, J. Kübler, C. Felser, and S. S. P. Parkin, Science Advances 2, e1501870 (2016).
[19] Y. Zhang, Y. Sun, H. Yang, J. Zelezny, S. P. P. Parkin, C. Felser, and B. Yan, Phys. Rev. B 95, 075128 (2017).
[20] M. Ihklaas, T. Tomita, T. Koretsune, M.-T. Suzuki, D. Nishio-Hamane, R. Arita, Y. Otani, and S. Nakatsuji, Nature Phys 13, 1085 (2017).
[21] N. J. Ghimire, A. S. Botana, J. S. Jiang, J. Zhang, Y.-S. Chen, and J. F. Mitchell, Nat Commun 9, 3280 (2018).
[22] M. Kimata, H. Chen, K. Kondou, S. Sugimoto, P. K. Mochuli, M. Ihklaas, Y. Omori, T. Tomita, A. H. MacDonald, S. Nakatsuji, and Y. Otani, Nature 565, 627 (2019).
[23] N. J. Ghimire, L. R. Dally, L. Poudel, D. C. Jones, D. Michel, N. T. Magar, M. Bleuel, M. A. McGuire, J. S. Jiang, J. F. Mitchell, J. W. Lynn, and I. I. Mazin, Sci. Adv. 6, eabe2680 (2020).
[24] T. Asaba, S. M. Thomas, M. Curtis, J. D. Thompson, E. D. Bauer, and F. Ronning, Phys. Rev. B 101, 174415 (2020).
[25] D. J. Thouless, M. Kohmoto, M. P. Nightingale, and M. den Nijs, Phys. Rev. Lett. 49, 405 (1982).
[26] G. Tenasini, E. Martin, N. Ubrig, N. J. Ghimire, H. Berger, O. Zaharko, F. Wu, J. F. Mitchell, I. Martin, L. Forró, and A. F. Morpurgo, Phys. Rev. Research 2, 023051 (2020).
[27] K. Anzenhofer, J. van den Berg, P. Cossee, and J. Helle, J. Phys. Chem. Solids 31, 1057 (1970).
[28] S. S. P. Parkin, E. A. Marseglia, and P. J. Brown, J. Phys. C: Solid State Phys. 16, 2765 (1983).
[29] O. Zaharko, Unpublished (2021).
[30] G. Kresse and J. Furthmüller, Phys. Rev. B 54, 11169 (1996).
[31] G. Kresse and D. Joubert, Phys. Rev. B 59, 1758 (1999).
[32] P. E. Blöchl, Phys. Rev. B 50, 17953 (1994).
[33] K. Burke, J. P. Perdew, and M. Ernzerhof, International Journal of Quantum Chemistry 61, 287 (1997).
[34] N. Marzari, A. A. Mostofi, J. R. Yates, I. Souza, and D. Vanderbilt, Rev. Mod. Phys. 84, 1419 (2012).
[35] G. Pizzi, V. Vitale, R. Arita, S. Blügel, F. Freimuth, G. Géranton, M. Gibertini, D. Gresch, C. Johnson, T. Koretsune, J. Ibañez-Azpiroz, H. Lee, J.-M. Lihm, D. Marchand, A. Marrazzo, Y. Mokrousov, J. I. Mustafa, Y. Nohara, Y. Nomura, L. Paulatto, S. Poncé, T. Ponweiser, J. Qiao, F. Thöle, S. S. Tsirkin, M. Wierzbowska, N. Marzari, D. Vanderbilt, I. Souza, A. A. Mostofi, and J. R. Yates, Journal of Physics: Condensed Matter 32, 165002 (2020).
[36] S. S. Tsirkin, npj Computational Materials 7, 33 (2021).
[37] K. Momma and F. Izumi, Journal of Applied Crystallography 44, 1272 (2011).
[38] O. Heinonen, R. A. Heinonen, and H. Park, arXiv:2109.13837 (2021).
[39] Y. Akagi and Y. Motome, Journal of the Physical Society of Japan 79, 083711 (2010).
[40] D. Solenov, D. Mozyrsky, and I. Martin, Phys. Rev. Lett. 108, 096403 (2012).
[41] S. Hayami, R. Ozawa, and Y. Motome, Phys. Rev. B 95, 224424 (2017).
[42] J. Shi, G. Vignale, D. Xiao, and Q. Niu, Phys. Rev. Lett. 99, 197202 (2007).