Quantum oscillations and magnetic reconstruction in the delafossite PdCrO₂

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We report quantum oscillation data on the metallic triangular antiferromagnet PdCrO₂. We find that, to very high accuracy, the observed frequencies of PdCrO₂ can be reproduced by reconstruction of the (nonmagnetic) PdCoO₂ Fermi surface into a reduced zone. The reduced zone corresponds to a magnetic cell containing six chromium sites, giving a \(\sqrt{3} \times \sqrt{3}\) in-plane reconstruction, and \(\times 2\) interplane reconstruction. The interplane ordering represents a reduction in lattice symmetry, possibly to monoclinic, and an associated lattice distortion is expected. In addition, we report a magnetic transition under an applied in-plane field that is probably equivalent to the spin-flop transition reported for CuCrO₂, and present data on its field-angle dependence. We also report measurements of the resistivity of PdCrO₂ up to 500 K.

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Magnetic ions coordinated on a triangular lattice often yield interesting magnetic properties. A prominent example is the insulating CrO₂ sheet, stabilized in materials such as LiCrO₂ and PdCrO₂. In these compounds, the Cr formal charge is +3, and its configuration is 3\(d^3\). The crystal field at the Cr sites is nearly octahedral, inducing a gap between the quasi-\(t_{2g}\) (\(d_{xy}, d_{xz},\) and \(d_{yz}\)) and quasi-\(e_g\) (\(d_{3z^2−r^2}\) and \(d_{xy}−zy\)) bands. The \(e_g\) bands lie above the Fermi level, and the \(t_{2g}\) bands are half-filled. The CrO₂ sheet is a Mott insulator; strong Hund’s rule coupling aligns the spins on each Cr site, giving a total spin on each Cr of nearly \(\frac{3}{2}\). The CrO₂ sheet is a Heisenberg system, that orders at low temperatures into the \(120°\) triangular Néel phase.

Of materials containing such a CrO₂ sheet, AgCrO₂, CuCrO₂, and PdCrO₂ have the delafossite crystal structure, while LiCrO₂, NaCrO₂, and KCrO₂ have the closely related ordered rock salt structure. All of these systems show \(120°\) order, with the Néel temperature depending strongly on the Cr-Cr spacing. The ordered rock-salt structure gives substantially smaller interplane spacings than the delafossite structure, and somewhat higher Néel temperatures [1].

The interlayer order is a more subtle problem than the intralayer order: in all of these systems the Cr sheets are stacked rhombohedrally, so the molecular fields from first- and second-neighboring layers cancel, and the ordering is determined by more subtle interactions susceptible to perturbation. The interlayer order has observable consequences: AgCrO₂ and CuCrO₂ show spin-driven ferroelectricity, while NaCrO₂ and LiCrO₂ do not [2]. AgCrO₂, and CuCrO₂ have the delafossite structure and NaCrO₂ and LiCrO₂ the ordered rock-salt structure, but the more essential difference appears to be the interlayer order: in AgCrO₂ [3] and CuCrO₂ [4,5], the interlayer ordering is ferroic, in that the vector chirality, the rotational sense of the spin helices that comprise the \(120°\) phase, is the same in all layers. In LiCrO₂, in contrast, neutron scattering data suggest that the vector chirality alternates from layer to layer [6], while the interplane order of NaCrO₂ is not clear [7].

This paper focuses on PdCrO₂. The Néel temperature of PdCrO₂ is \(T_N = 37.5\) K, and, like LiCrO₂, the vector chirality probably alternates from layer to layer [1,8], so this feature is not restricted to the ordered rock-salt structure.

Whereas the other compounds discussed above are insulators, PdCrO₂ is a metal [9], due to Pd 4\(d/5s\) conduction. The isostructural, nonmagnetic compound PdCoO₂ also has the Pd sheets, and its carrier mobility was found to exceed that of copper [10]. In PdCoO₂, the strongest \(d\) contribution appears to be from the 4\(d_{3z^2−r^2}\), however, the relative weight of this and Pd 5s at the Fermi level is not clear [11,12]. PdCrO₂, therefore, is an interesting system, comprised of highly conducting sheets interleaved with Mott insulating spacer layers. The resistivity anisotropy \((\rho_c/\rho_{ab})\) is about 180 at 295 K, and about 290 at 50 K [13]. It would be interesting to determine whether the metallic Pd conduction has any effect on the magnetic order.

What is certainly true is that the Pd conduction can be used as a probe of the magnetic order. Quantum oscillation [14] and angle-resolved photoemission spectroscopy (ARPES) [15,16] studies have shown that the PdCrO₂ Fermi surfaces result from reconstruction of the PdCoO₂ Fermi surface into the magnetic zone. Observation of an unconventional anomalous Hall effect
in PdCrO$_2$ indicates further that the Cr spins are not coplanar [17]. In this paper, we present measurements of quantum oscillations in PdCrO$_2$, and confirm the overall frequencies reported in Ref. [14]. We add greater resolution and a careful comparison with the nonmagnetic PdCoO$_2$ Fermi surface: we show that reconstruction of the PdCoO$_2$ Fermi surface reproduces most of the observed oscillation frequencies to very high accuracy: the PdCrO$_2$ frequencies can be analyzed in detail without recourse to density functional theory calculations. We show that the magnetic coupling is $k_z$ dependent: it is much weaker at $k_z = \pm \pi$ than at $k_z = 0$. Also, the dominant magnetic scattering vectors are those corresponding to a 6-Cr magnetic unit cell giving a $\times 2$ interplane reconstruction in addition to the $\sqrt{3} \times \sqrt{3}$ in-plane reconstruction. The interplane order represents a reduction in lattice symmetry, from $R\bar{3}m$ (rhombohedral) to, for the highest-symmetry possibility, $C2/c$ (monoclinic). We also report a magnetic transition under applied field, at a similar field to a spin-flop transition reported in CuCrO$_2$, that was observed in the course of measurement of the oscillations. Finally, we present measurements of the resistivity of PdCrO$_2$ to high temperatures.

I. METHODS

Single crystals of PdCrO$_2$ were grown by a NaCl flux method, using PdCrO$_2$ powder synthesized via a solid-state reaction [18]. We measured magnetic oscillations in two samples of PdCrO$_2$, by torque magnetometry, using the same piezoresistive AFM cantilevers as in our previous study on PdCoO$_2$ [10,19]. The cantilevers were mounted on rotatable platforms with integrated field angle sensors. Some raw data for sample #1 are shown in Fig. 1.

Sample #1 was roughly $230 \times 300 \times 11$ $\mu$m, and sample #2 $160 \times 100 \times 15$ $\mu$m. For sample #1, the field was rotated about a (1000) axis (using hexagonal indexing), and for sample #2 a (1100) axis; these axes are illustrated at the bottom right of Fig. 2. Both samples showed strong low-frequency oscillations ($\sim 800$ T for field angle $\theta \to 0$) that dominate the data. In sample #1, at 0.7 K these oscillations were discernible at fields as low as 2 T. At high fields (above $\sim 10$ T), the oscillation amplitude was very large, leading to strong torque interaction: the oscillations had a triangular form, and strong sum and difference frequencies appeared in the Fourier transforms. While torque interaction complicates the analysis somewhat, it may be difficult to avoid if the samples are to be large enough for higher-frequency oscillations to be resolved.

The oscillation amplitudes were history dependent. Cooling the sample through $T_N$ in a 15-T field resulted in oscillation amplitudes for the $\alpha$, $\beta$, and $\gamma$ orbits (illustrated in Fig. 3) roughly four times as large as with zero-field cooling. These large amplitudes persisted as long as the field was maintained. Releasing the field and heating the sample to $\sim 1$ K caused the amplitudes to decrease, and reapplying fields up to 15 T did not recover the large amplitudes. The system therefore appears to be nonergodic. A possible origin of glassy behavior is domain-boundary Cr spins: the relatively large magnetic unit cell of PdCrO$_2$ means that in the absence of training a complex domain structure is likely.
The oscillation frequencies as a function of field angle are shown in Fig. 2. Four sets of intrinsic peaks can be identified, located, at $\theta = 0$, at around 0.8, 3.5, 10.5, and 27.4 kT. Following Ref. [14], we label these $\alpha$, $\beta$, $\gamma$, and $\delta$. All the frequencies scale broadly as $1/\cos(\theta)$, indicating that the Fermi surfaces are highly two dimensional.\(^3\)

It has been established that the PdCrO$_2$ orbits result from a $\sqrt{3} \times \sqrt{3}$ reconstruction of the PdCoO$_2$ Fermi surface, due to the 120° Néel order [14,16]. This reconstruction is illustrated in Fig. 3. The $\alpha$ and $\gamma$ orbits are fully reconstructed orbits, while $\beta$ and $\delta$ result from magnetic breakdown, i.e., the orbits cross gaps in $k$ space.

The cyclotron masses were determined for sample #1 by Lifshitz-Kosevich fits to the temperature dependence of the

1While measuring the angle dependence of the oscillations in sample #1, we performed a “de-Gauss” routine at the start of each run, in which the field was ramped to $-2$ T, then $+1$ T, then $-0.5$ T, and so on. The aim was to reduce the probability of the system evolving gradually into a more ordered state, with larger oscillation amplitudes, though it is not clear whether the procedure had much effect.

2Some additional peaks appear in the data, between the $\beta$ and $\gamma$ frequencies, and are plotted in the figure. It was not obvious in the raw data whether they were real peaks, interaction peaks, or spurious; they are probably spurious.

3For calculation of the $\alpha$ and $\gamma$ expected masses, we use the $\alpha_1$ and $\gamma_1$ orbits.
Fig. 4 reverses the vector chirality within each layer, and monoclinic): reflection about the glide plane indicated in translation along the interplane ordering vector restores the nonmagnetic first Brillouin zone (black), together with the PdCoO2 white to indicate the three spin directions of each layer. Right: the surface is reconstructed.

The observed orbit, which is at $k_z = 0$, has the $\sim 35\%$ mass enhancement noted earlier; but so does the observed $\beta$ orbit, which is located at $k_z = \pm \pi$.

Overall, then, a two-dimensional $\sqrt{3} \times \sqrt{3}$ reconstruction of the PdCoO2 frequencies, where the reconstruction is weak at $k_z = \pm \pi$, gives a good description of the PdCrO2 frequencies at $\theta = 0$. We now test the reconstruction at other field angles by performing a three-dimensional reconstruction.

We first need a three-dimensional magnetic cell. It must contain at least six Cr sites: three per layer to capture the 120° Néel order, and two layers to capture the alternating vector chirality. The cell we test is shown in Fig. 4. The Cr sites are stacked rhombohedrally, so at $T_N$ there are three equivalent choices for the interplane ordering vector. In choosing one of them, the lattice symmetry is reduced. The space group of the nonmagnetic lattice is $R\bar{3}m$. The highest-symmetry possible magnetic lattice has space group $C2/c$ (base-centered monoclinic): reflection about the glide plane indicated in Fig. 4 reverses the vector chirality within each layer, and translation along the interplane ordering vector restores the original lattice. Magnetostructural coupling should lead to a lattice distortion associated with the reduction in symmetry, and it is possible that more subtle features in the magnetic order reduce the symmetry further.

The reconstruction is performed as described in textbooks: sections of the nonmagnetic Fermi surface are translated by combinations of reciprocal lattice vectors of the reduced zone until all portions of the original surface are represented within the reduced zone. For our nonmagnetic surface, we take the parametrized model for the PdCoO2 Fermi surface that was determined in Ref. [10]. The topology of the reconstructed Fermi surfaces is determined by the orientations of the avoided crossings that would occur in the real system, but the calculation does not include finite avoided crossings. Further details of the calculation are given in the Appendix. Our results are shown in Fig. 5, together with the observed frequencies.

There are two $\gamma$ bands. The lower band $\gamma_1$ derives from reconstruction of the belly orbit, and $\gamma_2$ the neck orbit. $\gamma_2$ is not observed in the data because the magnetic coupling is weak for the neck orbit.

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4In Refs. [6] (LiCrO2) and [1] (PdCrO2), single-crystal neutron scattering data are analyzed using an 18-site magnetic cell. This is the smallest that can preserve $R\bar{3}m$ symmetry: to preserve $R\bar{3}m$ symmetry the interplane ordering vector must take on all three possibilities in an ABCABC order, so six layers are required to capture both this and the alternating vector chirality. The most-likely magnetic structure that is obtained in Ref. [1] can be reduced to a six-site magnetic cell.

5The unit cell shown in Fig. 4 is metrically triclinic (none of the angles are 90°) but if the glide symmetry is preserved, then a 12-site monoclinic supercell can be constructed.
Three α bands appear in the data. The lower and upper bands α₁ and α₃ are reproduced well by the calculation. α₁ is from the nonmagnetic belly orbit and α₃ the neck orbit, so α₁ has a much lower amplitude in the data. The middle band α₂ is not reproduced in the calculations. It could be a breakdown orbit. There are prominent breakdown orbits in the data (β and δ), and the × 2 interplane reconstruction could lead to breakdown orbits that mix segments from the original neck and belly orbits.

For this calculation, we supposed that the system chose the interplane ordering vector that aligned the plane of (possible) glide symmetry with the field rotation plane. The other two possibilities would lead to the glide and rotation planes being separated by 120°. We also calculated this possibility, with the result shown in the Appendix. The match to the data is reasonable but not as good, so it appears that either the glide and rotation planes were aligned by chance, or the fields applied during the measurement reoriented the magnetic reconstruction.

We also show in the Appendix results for a strictly two-dimensional reconstruction, that would preserve the R3m symmetry of the nonmagnetic lattice. The results do not match the data well, and we conclude that the dominant magnetic scattering vectors are those of the magnetic cell shown in Fig. 4.

III. SPIN-FLOP TRANSITION

In addition to quantum oscillations, a large-scale feature appeared during the torque magnetometry measurements on sample #1: a first-order magnetic transition. It occurs at a field of around 6.5 T for θ = 90° (i.e., the field applied in the plane). On the high-field side of the transition, the sample has a much larger c-axis magnetic moment than on the low-field side. Figure 6 shows the set of torque curves obtained in this study, divided by the applied field to yield M⊥, the magnetization perpendicular to the applied field, and smoothed so as to exclude the oscillations and show broad-scale features. The transition and its hysteresis are readily apparent, at θ ~ 90°. It appeared for sample #1 but not #2, so it is sensitive to the direction of the applied field.

Over a small angle range, quantum oscillations were visible both below and above the transition; the Fourier transforms are shown in the Appendix. At each angle within this range, the oscillations have a lower amplitude on the high-field side of the transition than on the low-field side, and the dominant peak shifts to a slightly higher frequency. That the transition affects the oscillations in a consistent manner shows that it is a bulk property.

This transition is at a similar field to a spin-flop transition reported for CuCrO₂ [21], 5.3 T for temperatures well below Tₘ, and probably has the same origin. In CuCrO₂, it has been found by single-crystal neutron diffraction that the spins lie in ⟨0001⟩-(1100) planes [22,23]. The transition occurs when the field is applied in a ⟨1100⟩ direction, but there is no transition for fields applied in a ⟨1000⟩ direction. It has been shown by symmetry arguments [21] and by direct observation [22] that the transition is a spin flop, where the spin plane rotates by 90°. The spin flop is illustrated in Fig. 7, and discussed in greater detail in Ref. [21].

⁹Ok et al. also identify three α bands [14]. However, the frequencies they label α₁ and α₂ are both part of α₃ in our labeling scheme, and Ok et al. do not resolve the frequency we label α₁.
For PdCrO$_2$, although observation of an unconventional anomalous Hall effect shows that the spins are not coplanar, neutron scattering data show that the spins lie approximately in a (0001)-(1100) plane, similar to CuCrO$_2$. Furthermore, and also as in CuCrO$_2$, the transition occurs when the field is applied in a (1100) direction (sample #1), but not in a (1000) direction (#2). Therefore, the transition is probably the same spin flop.

IV. RESISTIVITY

PdCoO$_2$ has been used as a nonmagnetic analog of PdCrO$_2$ in order to extract the magnetic contribution to the specific heat and electrical resistivity [8, 13]. We have shown here that it is a very good comparison: the PdCrO$_2$ Fermi surfaces are, to the room-temperature resistivity determined in Ref. [10]: 2

in order to extract the magnetic contribution to the specific heat

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samples. Sample A had two pairs of voltage contacts. The point at

ear between

the same feature appears in PdCrO$_2$. The quantum oscillation frequencies indicate that the dominant magnetic scattering vectors are those of a six-site magnetic cell with lower symmetry than the nonmagnetic lattice. An associated lattice distortion is expected to onset at $T_N$. It has been looked for by both neutron and x-ray diffraction, and not found [1], so any distortion must be small.

The distortion, if it occurs, should resemble that observed in CuCrO$_2$ [26]; there is also evidence for a lattice distortion in AgCrO$_2$ [27]. In CuCrO$_2$, the vector chirality is the same in each layer and the magnetic cell contains three sites [2, 4, 5]. The magnetic transition is split into two [28], which is expected for easy-axis-type 120° antiferromagnetism: spins first order along the easy axis, then, at a lower temperature, along an in-plane axis. Ultrasound velocity measurements show that the lattice distortion starts at the upper transition, with no apparent anomaly at the lower transition [28]. The transition of PdCrO$_2$ may also be split, although if so the splitting is small and not generally observed in experiment [1]. Either way, the magnetic order implies two separate reductions in symmetry from $R3m$: the choice of interplane ordering vector at the (possible) upper transition, and the choice of spin plane at the lower transition (where, again, in PdCrO$_2$ the spins are only approximately coplanar). The two symmetry reductions are likely to couple. The ultrasound data on CuCrO$_2$ suggest that in that system it is the interplane ordering vector that more strongly drives the distortion.

The angle dependence of the spin-flop transition in PdCrO$_2$ is interesting: the transition field appears to vary smoothly from $\sim 15$ T on one side of the plane (at $\theta \sim +55^\circ$) to nearly zero on the other (at $\theta \sim -55^\circ$). At $B = 0$, however, the field angle is a meaningless parameter, so if there is in fact an end point near $B = 0$, then the high- and low-field states are adiabatically connected to each other. Whether this is true and how it might occur for a 90° spin flop requires further investigation.

In summary, PdCrO$_2$ is an interesting system comprised of alternating highly conductive sheets and Mott-insulating spacer layers. It provides a good model for study of the interaction between metallic conduction and antiferromagnetic

The form of the resistivity of PdCrO$_2$ is well established. There is a sharp cusp at $T_N$, and above $T_N$ the magnetic component of the resistivity $\rho_m$ remains well below its saturation value due to short-range correlation. The increase and eventual saturation of $\rho_m$ as the temperature increases leads to a convex temperature dependence of the resistivity.

The in-plane resistivity of PdCoO$_2$, in contrast to PdCrO$_2$ and PtCoO$_2$, is essentially linear from $\sim 200$ to at least 500 K. It may be that the temperature dependence of $\rho_m$ obscures an optical phonon contribution. $\rho_m$ is expected to saturate when the Cr spins become completely uncorrelated. If $\rho_m$ is estimated by subtracting the resistivity of PdCoO$_2$ from that of PdCrO$_2$, then our data indicate that $\rho_m$ continues to increase at temperatures well above room temperature, i.e., $\rho_m$ continues to evolve to temperatures an order of magnitude greater than $T_N$. The Weiss temperature of PdCrO$_2$ is $\sim -500$ K [8], so correlations between the Cr spins are expected to persist to temperatures up to $\sim 500$ K, and a high saturation temperature of $\rho_m$ may be expected.

V. DISCUSSION AND CONCLUSION

The transition of PdCrO$_2$ may also be split, although if so the splitting is small and not generally observed in experiment [1]. Either way, the magnetic order implies two separate reductions in symmetry from $R3m$: the choice of interplane ordering vector at the (possible) upper transition, and the choice of spin plane at the lower transition (where, again, in PdCrO$_2$ the spins are only approximately coplanar). The two symmetry reductions are likely to couple. The ultrasound data on CuCrO$_2$ suggest that in that system it is the interplane ordering vector that more strongly drives the distortion.

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In summary, PdCrO$_2$ is an interesting system comprised of alternating highly conductive sheets and Mott-insulating spacer layers. It provides a good model for study of the interaction between metallic conduction and antiferromagnetic

FIG. 8. The resistivity of PdCrO$_2$ against temperature, for two samples. Sample A had two pairs of voltage contacts. The point at 295 K, with error bars, indicates the room-temperature resistivity: 8.2 ± 1.0 $\mu\Omega$ cm. Data on a PdCoO$_2$ sample are also shown, scaled to the room-temperature resistivity determined in Ref. [10]: 2.6 ± 0.2 $\mu\Omega$ cm. Bottom panel: $d\rho/dT$ for sample A.
order: the conduction is well characterized, arising from a single, open Fermi surface, and much is known about the magnetic order. Also, there are related compounds that permit precise comparisons, such as the nonmagnetic PdCoO$_2$, and other CrO$_2$-based delafossite and ordered rock-salt materials. Several avenues of inquiry remain open.

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APPENDIX

In this Appendix we first present some supplementary data, and then give further details on the calculation of the three-dimensional reconstruction.

Figure 9 shows the Lifshitz-Kosevich fits to the oscillation amplitudes determined over the field range 8.5–9.5 T. Four such fits were done, over 1-T field ranges starting at 7.5, 8.5, 9.5, and 10.5 T. The masses reported in the main text are the averages of the masses obtained from these fits.

Figure 10 shows the Fourier transforms of the oscillations above and below the putative spin-flop transition, for the narrow angle range over which oscillations were observed on both sides of the transition. On the high-field side of the transition, the oscillations have lower amplitudes and the main peak a higher frequency. (The reduction in amplitude may not be immediately apparent in the figure, but normally quantum oscillation amplitudes grow rapidly as the field is increased, and the reduction in amplitude is very clear in the raw data.)

We now give further details on the reconstruction. The first step is to determine the misalignment, if any, between the sample and the field angle sensor mounted on the rotator platform. This misalignment is determined, for sample #1, by comparison of the observed $\delta$ frequencies and the scaled PdCoO$_2$ frequencies, shown in Fig. 2. The best match is obtained when a $0.3 \pm 0.2^\circ$ offset is added to the measured field angle. This is our misalignment, and correction for it is incorporated into all plots of sample #1 frequencies presented in this paper.

We take the nonmagnetic Fermi surface to be the parametrized PdCoO$_2$ Fermi surface that was determined in Ref. [10]. The parameters specifying the Fermi surface are the radius of a circular, cylindrical base Fermi surface $k_{00}$, and then the amplitudes of the corrugations on this base. $k_{60}$, for example, is the amplitude of the hexagonal distortion. $k_{01}$ sets the difference between the radii of the neck and belly orbits. A thorough description of this system of parametrization is given in Ref. [29].

We allow three parameters to be adjusted, within small ranges that are consistent with experiment, to match the overall levels of the observed PdCrO$_2$ frequencies. We emphasize that this adjustment only tunes the overall levels, and does not substantially alter any of the substructure. These parameters are as follows: (1) the overall size of the Fermi surface ($k_{00}$), (2) the magnitude of the hexagonal distortion ($k_{60}$), and (3) the in-plane lattice constant $a$. The $k_z$-dependent Fermi surface corrugations ($k_{01}$, $k_{02}$, and $k_{31}$) are left fixed at the amplitudes found for PdCoO$_2$. When $k_{60}$ is adjusted, the higher-order

![FIG. 9. Lifshitz-Kosevich fits to the amplitudes determined over the field range 8.5–9.5 T.](image)

![FIG. 10. Fourier transforms of the oscillations below (thin lines) and above (thick lines) the magnetic transition, which at the angles in this figure occurs at $B \sim 10$ T.](image)

![FIG. 11. (Color online) Oscillation frequency against field angle for two further possibilities for the magnetic reconstruction, discussed in the text.](image)
amplitudes $k_{12,0}$ and $k_{13,0}$ are adjusted by hand in response, to smooth the faces and sharpen the corners of the Fermi surface so as to match the Fermi surfaces observed by ARPES \cite{15,16}.

The best match is found with $k_{00} = 0.9219 \, \AA^{-1}$, $k_{40} = 0.036 \, \AA^{-1}$, and $\alpha = 2.927 \, \AA$. Further refinement (for example, by allowing a low-temperature lattice distortion, or modification of the $k_z$-dependent corrugations, or finite avoided crossings) might yield slightly different results: the obtained lattice parameter, for example, is not precisely in line with expectations, as it has been measured to be 2.923 \, \AA at room temperature and some thermal contraction is expected \cite{1}. However, these fitting parameters are within bounds permitted by experiment and are adequate to model the angle dependence.

In addition to the reconstruction shown in the main text, we tested two further possibilities. The first is the same reconstruction but with the other possible orientation relative to the field rotation plane, that the glide plane is rotated from the field rotation plane by 120°. The second tests a hypothetical magnetic zone with vertical side walls, that would preserve the $R\overline{3}m$ symmetry of the nonmagnetic lattice. The results are shown in Fig. 11.

The rotated reconstruction, shown in the left-hand panels in Fig. 11, gives a reasonable match to the data, but not as good as with the glide and rotation planes aligned: it yields the wrong sign on the slope of $\alpha_3$ against $\theta$, and too low an amplitude on the substructure of the $\gamma_1$ frequencies. The $R\overline{3}m$ reconstruction gives a worse match: although it reproduces the $\alpha_1$ and $\alpha_3$ frequencies reasonably well, it does not reproduce the structure on $\gamma_1$ at all. Therefore, we conclude that the magnetic reconstruction illustrated in Figs. 4 and 5 is the correct one.

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