Impact of further-range exchange and cubic anisotropy on magnetic excitations in the fcc kagome antiferromagnet IrMn$_3$

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Abstract

Exchange interactions up to fourth nearest neighbor are shown within a classical local-moment Heisenberg approach to be important to model inelastic neutron scattering data on the fcc kagome antiferromagnet IrMn$_3$. Spin wave frequencies are calculated using the torque equation and the magnetic scattering function, $S(Q, \omega)$, is determined by a Green’s function method, as an extension of our previous work, LeBlanc et al. Phys. Rev. B 90, 144403 (2014). Results are compared with intensity contour data on powder samples of ordered IrMn$_3$, where magnetic Mn ions occupy lattice sites of ABC stacked kagome planes. Values of exchange parameters taken from DFT calculations used in our model provide good agreement with the experimental results only if further-neighbor exchange is included. Estimates of the observed energy gap support the existence of strong cubic anisotropy predicted by DFT calculations.

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I. INTRODUCTION

IrMn$_3$ provides an important example of a truly three dimensional (3D) kagome lattice giving rise to geometrical magnetic frustration from eight near-neighbor (NN) antiferromagnetic exchange interactions between Mn ions. ABC stacking of kagome planes of Mn ions in the cubic $\langle 111 \rangle$ directions gives an overall L1$_2$, AuCu$_3$-type, fcc structure with four NNs within each plane and two NN connecting each adjacent plane (see Fig. 1). Interest in the magnetic properties of the corresponding two-dimensional (2D) kagome NN Heisenberg antiferromagnet spans 25 years due to the macroscopic spin degeneracy of the basic 120° spin structure associated with corner-sharing triangles. The zero energy dispersionless (flat) spin wave mode at all wave vectors predicted from classical theory acquires dispersion in the presence of further-neighbor exchange interactions. Inelastic neutron scattering data on a system with weakly coupled kagome planes appear consistent with this scenario where the flat mode observed is gapped due to additional Dzyaloshinskii-Moriya interactions.

![ABC stacked kagome planes forming the fcc kagome lattice with magnetic ions occupying the cube face center sites. Illustrated are spin vectors forming the 120° $q=0$ [111] planar spin structure (zero anisotropy) and the exchange interactions $J_1$, $J_2$, $J_3$, and $J_4$. Also see Table I.](image)

Interest in IrMn$_x$ alloys over the past 15 years has mainly arisen due to applications in
spin-valve technology, where they have been widely used as the antiferromagnetic thin film of choice that pins the magnetic moments of an adjacent ferromagnet in the phenomenon known as exchange bias. Although there is no universally accepted microscopic mechanism for exchange bias, magnetic frustration is believed to be important and stoichiometric IrMn$_3$ appears to optimize the desired pinning.

Monte Carlo simulations of the NN Heisenberg fcc kagome lattice have shown that the basic co-planar $120^\circ$ q=0 magnetic structure observed in 2D persists in the 3D case, with the inter-spin angle being $120^\circ$ between all eight NNs (shown in Fig. 1). The spin degeneracy is reduced in 3D and exists in the form of sublattice magnetization switching in the stacked kagome planes. This persistent degeneracy is believed to be responsible for the first-order nature of the phase transition at $T_N$. Inspired by earlier electronic structure calculations, subsequent simulations of the 3D lattice with an effective local cubic anisotropy included provided evidence that anisotropy removes the basic kagome degeneracy and the structure becomes non co-planar with a finite magnetization (spin vectors are lifted out of the [111] plane). This release of frustration drives the transition to be continuous. This scenario was supported by spin wave calculations of the NN Heisenberg model with and without anisotropy which demonstrated that in the absence of anisotropy, the zero energy flat mode exists only in certain high symmetry directions in reciprocal space and that the addition of anisotropy induces a gap (Ref.14 hereafter referred to as I). Monte Carlo studies of [111] thin films confirmed that the q=0 magnetic order remains in these geometries and the impact of surface axial anisotropy was also considered.

Early neutron scattering experiments on sister compounds RhMn$_3$ and PtMn$_3$ revealed the $120^\circ$ magnetic order which was subsequently established also in single crystal neutron scattering studies on IrMn$_3$ by Tomeno et al and referred to as “T1” magnetic order, with no mention of the underlying kagome lattice structure of magnetic ions or any indication of a finite magnetization. More recently, single crystal [111] thin films of IrMn$_3$ also showed the same magnetic order as in the bulk where exchange bias was also studied. To our knowledge, there have been no neutron scattering experiments reporting on spin excitations in these magnetic fcc kagome systems. It is of interest to note another class of Mn-based compounds with the generic formula Mn$_3$AX also exhibits fcc kagome magnetism.

The high Néel temperature in IrMn$_3$, $T_N \approx 960$ K (with large values also reported in the sister compounds), attractive for device applications, can be associated with large exchange
Exchange Interactions in IrMn$_3$

| nth Near Neighbor | $J_1$ | $J_2$ | $J_3$ | $J_4$ |
|-------------------|------|------|------|------|
| Value from DFT (meV) | 40   | -5   | 10   | -5   |
| Neighbors in plane | 4    | 0    | 4    | 6    |
| Neighbors first adjacent planes | 4 | 6 | 8 | 0 |
| Neighbors second adjacent planes | 0 | 0 | 4 | 6 |
| Total | 8 | 6 | 16 | 12 |
| Vector | $[\frac{1}{2}, \frac{1}{2}, 0]$ | [1 0 0] | $[1, \frac{1}{2}, \frac{1}{2}]$ | [1 1 0] |
| Distance (a) | 0.707 | 1 | 1.225 | 1.414 |

TABLE I. Exchange parameters for the fcc kagome lattice IrMn$_3$ (see Fig. 1) where $J > 0$ implies antiferromagnetic coupling. DFT values are taken from Ref. [10]. Distances are relative to the lattice constant $a = 3.76$ Å.

interactions. For example, NN $J_1 \simeq 40$ meV (here, $J > 0$ implies an antiferromagnetic interaction) has been estimated for IrMn$_3$ from the density functional theory (DFT) calculations by Szunyogh et al. [10], where an effective spin $S = 2.2$ has been folded into the reported exchange constant values. This and related work also reported substantial longer range exchange interactions of an oscillatory nature, with $J_2 \simeq -5$ meV, $J_3 \simeq 10$ meV, $J_4 \simeq -5$ meV, as well as a large effective cubic anisotropy $K_{\text{eff}} \simeq 7.67$ meV (see Table I). The DFT results indicate that further exchange interactions beyond fourth-neighbor are negligibly small. Analysis shows that these longer range alternating antiferromagnetic and ferromagnetic exchange interactions are consistent with the 3D $q = 0$ spin structure and do not introduce additional frustration. A first-principles molecular spin dynamics study of PtMn$_3$ and RhMn$_3$ also reports enhanced second-neighbor exchange interactions.

In the present work, the earlier local-moment spin-wave and inelastic magnetic scattering intensity calculations reported in I are expanded to include the further-neighbor exchange $J_2$, $J_3$, and $J_4$. As in the 2D case, these additional interactions remove any flat modes and dispersion appears in all cases. The impact of cubic anisotropy, $K$, is again examined in the presence of the additional exchange interactions. Scattering intensity, $S(Q, \omega)$, contours are calculated for both single crystal and powder sample scenarios. Results are compared
with new inelastic neutron scattering data on powder samples of ordered phase IrMn₃. The
effects of further-order exchange are demonstrated to be important and an energy gap is
observed.

II. MODEL RESULTS

Modifications to our previous analysis in I to include further-neighbor exchange interac-
tions are described here. In that work, J represented the NN exchange in kagome planes
and J′ denoted inter-plane exchange. Here, we set J₁ = J = J′. The 120° q=0 spin structure
is characterized by a three magnetic sublattice magnetization vectors labeled as A, B and
C, and we consider the following Hamiltonian

$$H = \sum_{i<j} J(|r_i - r_j|)S_i \cdot S_j - K \sum_{\gamma} \sum_{l \subset \gamma} (S_l \cdot n_{\gamma})^2. \tag{1}$$

Note that $K = \frac{1}{2}K_{\text{eff}} \geq 0$ has a different easy direction for each of the three sublattices.
Here, $\gamma$ represents sublattice A, B and C and $l$ is summed over the $N_{\gamma}$ spins of sublattice
$\gamma$, $S_i$ are unit classical Heisenberg spin vectors at each site and $n_{\gamma}$ are unit vectors in the
cube axes directions, $n_A = \hat{x}$, $n_B = \hat{y}$, and $n_C = \hat{z}$. The lattice constant of ordered IrMn₃
has been determined at room temperature to be around $a=3.76$ Å. The Mn ions occupy
face center sites, with $\gamma = A$ at $[0, \frac{1}{2}, \frac{1}{2}]$, $\gamma = B$ at $[\frac{1}{2}, 0, \frac{1}{2}]$, and $\gamma = C$ at $[\frac{1}{2}, \frac{1}{2}, 0]$, separated
by a distance $a/\sqrt{2} = 2.67$ Å as depicted in Fig. 1. The model results presented below
include the exchange constants up to fourth near-neighbors, with values given in Table I, as
well as anisotropy. In order to demonstrate the impact of these further-neighbor exchange
interactions, as well as anisotropy, we also consider model results with some of the constants
set to zero, or anisotropy set to zero. We do not attempt to fit model parameters with the
data described below.

As described in I, in the absence of anisotropy spins are coplanar with zero net mag-
netization. Anisotropy serves to lift the spin vectors out of the plane and induce a finite
magnetization in a ⟨111⟩ direction. This effect is characterized by $\alpha$, the cosine of the angle
between each sublattice spin and its anisotropy axis ($\alpha = \cos(S_i \cdot n_i)$, $i = A, B, C$), and $\beta$,
the cosine of the angle with respect to the other two anisotropy axes ($\beta = \cos(S_i \cdot n_j)$,
$i \neq j$) where $\alpha^2 + 2\beta^2 = 1$. The modified ground state energy per spin that includes
further-neighbor exchange is given by (compare Eqs. 2 and 3 in I where $S=1$ was used for
\[ E/N = 4(J_1 + 2J_3)(\beta^2 - 2\alpha\beta) - K\alpha^2 + 3J_2 + 6J_4 \] (2)

and is minimized when \( \alpha \) has the value

\[ \alpha = \sqrt{\frac{1}{2} + \frac{1}{2\sqrt{1 - 1/[1 + (K + 1)^2/8]}}} \] (3)

where \( K = K/(2J_1 + 4J_3) \) and \( \beta = \sqrt{1 - \alpha^2} \) using the positive values of the square roots to give physical solutions. Note that there are eight degenerate ground states corresponding to the eight \( \langle 111 \rangle \) axes.\(^{13}\) The analysis of spin excitations in this section correspond to a single domain [111] crystal. Powder averaged results are discussed in the following section.

The basic structure of the linearized spin wave theory presented in I remains the same with further-neighbor exchange added. The \( 6 \times 6 \) matrix characterizing dynamic fluctuations of the transverse spin components in a local coordinate system is again given by

\[
-i\omega \begin{pmatrix}
\tilde{S}_A^x \\
\tilde{S}_B^x \\
\tilde{S}_C^x \\
\tilde{S}_A^y \\
\tilde{S}_B^y \\
\tilde{S}_C^y
\end{pmatrix} =
\begin{pmatrix}
0 & Y_{AB} & -Y_{AC} & X & Z_{AB} & Z_{AC} \\
-Y_{AB} & 0 & Y_{BC} & Z_{AB} & X & Z_{BC} \\
Y_{AC} & -Y_{BC} & 0 & Z_{AC} & Z_{BC} & X \\
W & T_{AB} & T_{AC} & 0 & Y_{AB} & -Y_{AC} \\
T_{AB} & W & T_{BC} & -Y_{AB} & 0 & Y_{BC} \\
T_{AC} & T_{BC} & W & Y_{AC} & -Y_{BC} & 0
\end{pmatrix}
\begin{pmatrix}
\tilde{S}_A^x \\
\tilde{S}_B^x \\
\tilde{S}_C^x \\
\tilde{S}_A^y \\
\tilde{S}_B^y \\
\tilde{S}_C^y
\end{pmatrix}
\] (4)

where \( Y, T, \) and \( Z \) are defined in I and with the following modifications:

\[ X = 8(J_1 + 2J_3)(\beta - 2\alpha)\beta - 2K\alpha^2 \]

\[ + 6J_2 - 2J_2(\cos Q_x a + \cos Q_y a + \cos Q_z a) \]

\[ + 12J_4 - 4J_4(\cos Q_x a \cos Q_y a + \cos Q_z a \cos Q_x a + \cos Q_y a \cos Q_z a)]/S \]

\[ W = 8(J_1 + 2J_3)(2\alpha - \beta)\beta + 2K(\alpha^2 - 2\beta^2) \]

\[ - 6J_2 + 2J_2(\cos Q_x a + \cos Q_y a + \cos Q_z a) \]

\[ - 12J_4 + 4J_4(\cos Q_x a \cos Q_y a + \cos Q_z a \cos Q_x a + \cos Q_y a \cos Q_z a)]/S \]

and

\[ \lambda_{AB} = 4S^{-1}[J_1 + 2J_3 \cos Q_x a] \cos (\frac{1}{2}Q_x a) \cos (\frac{1}{2}Q_y a) \]

\[ \lambda_{BC} = 4S^{-1}[J_1 + 2J_3 \cos Q_y a] \cos (\frac{1}{2}Q_x a) \cos (\frac{1}{2}Q_y a) \]

\[ \lambda_{AC} = 4S^{-1}[J_1 + 2J_3 \cos Q_z a] \cos (\frac{1}{2}Q_x a) \cos (\frac{1}{2}Q_y a) \] (6)
where coordinates are in terms of cube axes with lattice constant $a$. Note that for the spin-wave frequency, the bare exchange and anisotropy constants are divided by $S$.

In general, numerical analysis is required to obtain the spin wave frequencies as a function of wave vector but some special cases mentioned in I can again be determined analytically. For all of the numerical results shown below, values from the DFT calculation given in the Introduction were used.

**A. Zero Anisotropy**

For the case $K = 0$, the eigenvalue problem involves the $3 \times 3$ symmetric matrix Eq. (7) in I with elements now given by

\[
\begin{align*}
A_1 &= X^2 - (\lambda_{AB}^2 + \lambda_{AC}^2)/2 \\
A_2 &= X^2 - (\lambda_{AB}^2 + \lambda_{BC}^2)/2 \\
A_3 &= X^2 - (\lambda_{AC}^2 + \lambda_{BC}^2)/2 \\
B_1 &= -X\lambda_{AB}/2 - \lambda_{AC}\lambda_{BC}/2 \\
B_2 &= -X\lambda_{AC}/2 - \lambda_{AB}\lambda_{BC}/2 \\
B_3 &= -X\lambda_{BC}/2 - \lambda_{AB}\lambda_{AC}/2
\end{align*}
\]

(7)

The special case $Q_x = Q_y = Q_z$ again yields eigenvalues of the general form given by Eq. (11) in I but further reduction in terms of the $J_i$ is not illuminating. However, for the case where the wave vector is directed along a cube axis, $Q_y = Q_z = 0$ for example, it can be shown that one of the three modes takes the form

\[
\omega_1^2 = S^{-2}(2J_2 + 8J_4 - 8J_3)[(6J_1 + 12J_3)(\cos Q_x a - 1) \\
+ (2J_2 + 8J_4 + 4J_3)(\cos Q_x a - 1)^2]
\]

(8)

This yields the zero energy mode Eq. (13) of I in the absence of further-neighbor exchange. It is also zero if $2J_2 + 8J_4 = 8J_3$, which is not possible in the case of IrMn$_3$ with $J_3 > 0$ and $J_2, J_4 < 0$. Thus, as in the 2D case, further-neighbor exchange removes the flat mode.
FIG. 2. Spin wave modes along the ΓX, ΓM and ΓR directions illustrating the impact of anisotropy and further-neighbor exchange interactions. In the case of NN exchange only with $K=0$ (panel (c)), there is a zero energy mode along ΓX (black line) and the remaining two modes (red and green lines) are degenerate. Values from the DFT calculation given in the Introduction were used.

B. Effects of Anisotropy

Fig. 2 shows the spin wave frequencies $\omega$ along the ΓX(100), ΓM(110) and ΓR(111) directions with only NN exchange $J_1$ included and also with further-neighbor $J_2$, $J_3$, and $J_4$ added, with and without cubic anisotropy $K$ (also see I). The effect of further-neighbor interactions is to lift the degeneracy that gives rise to the flat zero energy mode (black line)
that occurs along ΓX with only NN interactions (Figs. 2a and c). The resulting dispersion of this mode is now nearly the same as the other two branches. Note as well that the degeneracy of the other two modes (red and green lines) occurring around the X point in the case of only NN exchange is split with longer-range interactions included. As before, the impact of anisotropy $K > 0$ is to introduce a gap at the zone center (Figs. 2b and d). Also note that two of the modes along the Γ - M line are degenerate with further-neighbor exchange added (green and black lines in Figs. 2a and b) that were well separated with only NN interactions (Figs. 2c and d). In contrast, the near degeneracy of these modes around the R point is little impacted by including further-neighbor exchange.

At the zone center $Q = 0$, the small $K$ dependence of the modes can be calculated and are given by

$$
\omega_1 \approx \omega_2 \approx 2S^{-1}\sqrt{(J_1 + 2J_3)K}
$$

$$
\omega_3 \approx 4S^{-1}\sqrt{(J_1 + 2J_3)K}
$$

yielding the prediction of small $Q$ gaps of about 13 meV and 26 meV, respectively, for IrMn$_3$.

C. Dynamic structure factor

The Green’s function method used in I was applied here to calculate the part of the dynamic structure factor that contributes to the inelastic neutron scattering cross section

$$
S(Q, \omega) = \sum_{m,n=x,y,z} S^{mn}(Q, \omega)(\delta_{mn} - \hat{Q}_m \hat{Q}_n)
$$

where $S^{mn}(Q, \omega)$ is the double Fourier transform of the correlation function $<S^m_i(0)S^n_j(t)>$, to provide an indication of the inelastic neutron scattering response for IrMn$_3$ with all four NN exchange interactions and anisotropy included.

Fig. 3 shows $S(Q, \omega)$ assuming a single magnetic (111) domain for $Q$ along the three principal cube directions with further-neighbor exchange as well as anisotropy included. Corresponding results with only NN exchange and $K$ included for $Q$ along [100] may be found in I. Of particular note is that for the cases with $Q$ along [100] and [110], the intensity is expected to be relatively small in the first Brillouin zone but is substantially larger in the second zone. This is not the case for $Q$ along [111] but the overall intensity is much weaker for the wave vector in this direction.
FIG. 3. Dynamic structure factor, $S(Q, \omega)$, for IrMn$_3$ assuming a single magnetic (111) domain along the three principal cubic directions. Indicated wave vectors show the symmetric points in the dispersion. Exchange interactions $J_1$, $J_2$, $J_3$, and $J_4$, as well as anisotropy are included. Values from the DFT calculation given in the Introduction were used.

Fig. 3 can be compared with Fig. 2b, illustrating the removal of the low frequency mode along $\Gamma X$ with the addition of extra neighbors and the splitting of the degeneracy of the higher frequency modes. Noticeably, along the [111] direction there is a mode not seen in Fig. 3c that is present in Fig. 2b. This is not an added degeneracy, but is rather due to a perfect cancellation in intensity when applying Eq. 10. A similar effect is observed in the elastic scattering results of Fig. 3 of I where there is no peak at [111] unless anisotropy is added.

III. INELASTIC NEUTRON SCATTERING

A. Model for scattering cross section

To provide a more meaningful comparison with the experimental data, the relevant parts of the scattering cross section for inelastic magnetic scattering are also calculated

$$\left( \frac{d^2\sigma}{d\Omega dE'} \right)_{\text{inel}} \propto \frac{k'}{k} |F(Q)|^2 S(Q, \omega), \quad (11)$$
where $S(Q, \omega)$ is given by Eq. 10 and $F(Q)$ is the magnetic form factor. Additional kinematic factors in the cross section, e.g., $k'/k$ (where $k$ and $k'$ are the initial and final scattering wavevector magnitudes, respectively) are accounted for in the reduction of the experimental data. This form factor is given by, in the usual dipole, or spherical, approximation,

$$F(s) = Ae^{-as^2} + Be^{-bs^2} + Ce^{-cs^2} + D,$$

where $s = Q/4\pi$. This function approaches zero at $s > \sim 1\text{Å}^{-1}$ and is negligible for $Q > \sim 4\pi\text{Å}^{-1}$.

In principle the values of the dimensionless constants depend on the oxidation state of Mn. However, little variation is observed so the tabulated values for Mn0 were used.\textsuperscript{22} The wavevector is calculated through $Q = \frac{2\pi}{a} \sqrt{h^2 + k^2 + l^2}$, using the low temperature lattice constant $a = 3.76(1)$.\textsuperscript{17}

\textbf{B. Calculations for comparison with neutron scattering data}

For comparison of the calculated cross section with the data from the polycrystalline sample used in the SEQUOIA experiment, $S(Q, \omega)$ is averaged over all crystallographic directions, yielding

$$S(Q, \omega) = \int_{\phi=0}^{2\pi} \int_{\theta=0}^{\pi} S(Q, \omega) \sin \theta \, d\theta \, d\phi,$$

where $\theta$ and $\phi$ are the azimuthal and polar angles describing the orientation of $Q$. This integral can be approximated through Monte Carlo integration using the following expression:

$$S(Q, \omega) = \frac{1}{n} \sum_{i=1}^{n} S(Q, \arccos(2a_i - 1), 2\pi b_i, \omega).$$

In the equation above, $a$ and $b$ represent two random numbers from 0 to 1 and $n$ is the number of iterations. Typically, 1000 random directions are chosen for each $Q$. For the model results presented below, the scattering cross section Eq. 11 was then calculated.

Figure 4 shows the calculated powder-averaged spectra, for comparison to the inelastic scattering results acquired with $E_i = 100$ meV, for four cases of setting selected further neighbor exchange constants equal to zero. All of the results include anisotropy. The energy gap appears to be about 13 meV from these model results.

The strongest intensity features predicted by the model that includes further-neighbor exchange up to at least $J_3$ (Figs. 4a and b) at $Q$=1.67, 2.37, 3.75 and 4.10 Å$^{-1}$ are observed. This is not surprising as they are above the (100), (110), (210), and (211) magnetic Bragg
FIG. 4. Calculated, powder-averaged inelastic neutron scattering spectrum for $E_i = 100$ meV with (a) all four further-neighbor exchange $J_1$, $J_2$, $J_3$, and $J_4$; (b) with $J_1$, $J_2$, and $J_3$ only; (c) with $J_1$ and $J_2$ only; and (d) with nearest neighbor exchange $J_1$ only. Anisotropy is included in all cases and these model results suggest a gap of about 13 meV. Values from the DFT calculation given in the Introduction were used.

peak positions. Figure 4 shows that these excitations are less distinct from each other indicating that $J_3$ is important to provide the overall bandwidth of the excitation spectrum. A significant difference between the NN only model (Fig. 4a) and the others with further-neighbor exchange is that the inelastic feature above $Q=2.37$ Å$^{-1}$ (110 peak position) is much more intense and well-defined with the inclusion of $J_3$ and $J_4$. The other regions of well defined but lower intensity resulting from the further-neighbor model at larger wave vectors also do not occur within the NN only model.

As shown below, a comparison of the experimental data with theses figures show that the inclusion of all four exchange interactions yield the best agreement. In Figure 4 the kinematic
FIG. 5. The powder averaged calculation showing the full range of the dispersion for the model including anisotropy and all of the neighbor interactions. Values from the DFT calculation given in the Introduction were used.

constraints of the neutrons scattering experiment were applied to allow for comparison. In Figure 5 that constraint is removed to be able to compare to multiple $E_i$ values. In other words, it shows the full powder averaged dispersion with the contributions from all of the neighbors. The data in this figure can then be used as input in the Monte Carlo ray tracing simulations, that account for instrumental effects such as resolution and are described below, for closer comparison to the measurements.

C. Sample preparation

Mn and Ir powders in the molar ratio of 3:1 were mixed well and pelletized. The pellets were then sealed inside of a quartz tube under approximately 1/3 atmosphere of argon gas. The sealed ampoule was heated to 1050°C in 10 hours and kept at this temperature for 48 hours before cooling to 600°C in 6 hours. After dwelling at 600°C for 7 days, the ampoule was quenched into iced water. X-ray diffraction confirmed that the resulting pellet was nearly single phase IrMn$_3$ (ordered Mn and Ir, $Pm3m$ space group), with a small amount of MnO.
detected on the surface. No preferred crystal orientation was detected.

D. Inelastic neutron scattering: experimental details

Inelastic neutron scattering experiments were performed on the direct-geometry time-of-flight (TOF) chopper spectrometer SEQUOIA\textsuperscript{23,24} at Oak Ridge National Laboratory’s Spallation Neutron Source. The polycrystalline sample was cut into thin slabs to minimize neutron absorption in the INS experiment. The cell was then cooled down to 5 K in a closed cycle refrigerator. Spectra were collected with incident energies $E_i = 50$, 100, 300, and 500 meV in coarse energy resolution mode (elastic resolution of $\sim 4 \% E_i$) to investigate the spin wave excitations in the magnetically-ordered phase. An empty Al cell was measured in identical experimental conditions for the $E_i = 50$, 100, and 500 meV cases and the resulting spectra were subtracted from the corresponding sample spectra. Additional $E_i = 300$ meV data was also collected, but since this data was only used to investigate magnetic excitations above the cutoff for the Al phonon density-of-states the empty Al cell was not measured in this case. Coarse energy resolution mode was achieved using Fermi Chopper 2 operating at 240 Hz ($E_i = 50$ or 100 meV), 480 Hz ($E_i = 300$ meV) or 600 Hz ($E_i = 500$ meV), and the background from the prompt pulse was removed with a $T_o$ chopper operating at 60 Hz ($E_i = 50$ or 100 meV), 120 Hz ($E_i = 300$ meV) or 150 Hz ($E_i = 500$ meV).\textsuperscript{24}

E. Inelastic neutron scattering: results

The inelastic neutron results are reduced from TOF and pixel position to $Q$ and $E$ using Mantid\textsuperscript{25} and the $S(Q,E)$ slices and cuts were generated from this reduced data using DAVE.\textsuperscript{27} The color contour plots in Fig. 6 summarize the main results from the SEQUOIA experiment. Panels (a), and (b) depict the IrMn\textsubscript{3} spectra collected with $E_i$= 50 and 100 meV, respectively. As $E_i$ is increased, the energy resolution coarsens, and the kinematic range broadens. Several prominent features are shown in the figure. First and most interesting are the three nearly-vertical columns of scattering indicated by the arrows in Fig. 6b. These columns are centered above $Q = 1.67$, 2.36, 3.75, and 4.10 Å$^{-1}$, which correspond to the (100), (110), and the combined (210) and (211) magnetic Bragg peak positions for IrMn\textsubscript{3}. A more detailed examination of these excitations show that they match the further-neighbor
FIG. 6. (a) Color contour plots of spectra measured on SEQUOIA. (a) For $E_i = 50$ meV we observe a magnon band due to a small MnO impurity, a nearly-vertical column of magnetic scattering centered about $Q = 1.67 \text{ Å}^{-1}$ corresponding to the magnon band for IrMn$_3$, and some phonon modes. (b) For $E_i = 100$ meV we observe four nearly-vertical columns of scattering correspond to IrMn$_3$ spin wave excitations.

model calculations as will be discussed in detail below. Second, there is a strong phonon background between $\sim 10 - 35$ meV. Third, the weakly-dispersive magnetic mode, between 10 and 18 meV in a $Q$ range of 0.7-2.5 Å$^{-1}$, can be attributed to a magnon band arising from 2 ± 1% of MnO impurities.

F. Comparison between SEQUOIA data and spin wave calculations

To perform a quantitative comparison to the theoretical calculations, the results were fed into a McVine Monte Carlo ray tracing simulation of the SEQUOIA instrument. The simulation provides a computer model of each component of the instrument including the sample. Simulated neutrons are then propagated through the model. Once a sufficient number are propagated, the calculated result is analyzed in the same way as the data. The
end result is a directly comparable calculation with the instrumental resolution implicitly included. McVine provides a straightforward way to model multiple contributions like impurity phases and multiple scattering. Both the excitations from the IrMn$_3$ and the MnO were modeled. However, the unknown phonon contribution was not included in the simulation. A few simulations with multiple scattering were tried but did not improve the agreement with the measured data so the given results show single scattering events only. For the simulations, $1 \times 10^{10}$ neutron probability packets were propagated from the source and through the various instrument components.

Figure 7 shows a constant-$Q$ cut of the experimental data, with an integration range of $Q = [1.5, 1.9]$ Å$^{-1}$. This $Q$ range is centered around the (100) Bragg peak which has both structural and magnetic components due to the known $q = 0$ magnetic order. The mode appears to be strongly-gapped, as the spectral weight decreases significantly at energy transfers below the MnO magnon band. These measurements estimate the value of the gap to be $12 \pm 2$ meV, a value in good agreement with the model results (based on DFT calculations of the cubic anisotropy) and the McVine simulations. Further refinement of the gap value is hampered by the phonon background.

As mentioned earlier, for the $E_i = 100$ meV measurements, the $Q=2.36$ Å$^{-1}$ peak is clearly observed in Fig. 6b. However for the calculations, it is not clearly observed in the NN model (Fig. 4c) and only becomes prominent when further-neighbor interactions are included (Fig. 4b). It can thus be concluded that a model with NN exchange only does
not explain the measured data. Further examination of Fig. 6b shows that the phonon background is greatly reduced above 40 meV, which ensures that there is a large region of Q and E space with magnetic contributions only. This provides a region in Q and E space that is cleanly magnetic and facilitates a straightforward and direct comparison with the models. To this end, a constant E-cut with an integration range of 40 - 70 meV was considered. The result is shown as the black circles in Fig. 8. The same cut was taken from the \( E_{\text{c}} = 100 \) meV McVine simulations and is shown as the red line superimposed on that data. Note that the simulation includes a small linear background contribution. The simulation accounts for the data quite nicely and quantitatively shows that the theoretical model is appropriate. If the calculated dispersion curves were significantly different it would show up in the widths of the peaks.

For a broader comparison, data was acquired at \( E_{\text{i}} = 300 \) meV. This setting allowed the data to be acquired with sufficient range and resolution to compare to the model over the 70 - 140 meV range. For this comparison 10 meV wide cuts were taken in 10 meV steps. The data are shown as points in Figure 9 and the simulations are superimposed on the data with lines of the same color. As mentioned above, a linear background was added to the simulation and an overall scale factor was also incorporated to allow for a direct comparison to the SEQUOIA data. There were no other free parameters in the fit. Note the excellent agreement between the model and the data confirming that the additional neighbor model accounts for the magnetic excitations observed in the measurements. A similar process of fitting the simulation with a scale factor and a linear background was tried for a case where \( J_4 = 0 \). The results are shown as the dashed curves in Figure 9. Note that at low energies this model does not reproduce the data and furthermore results in a narrower bandwidth of excitations yielding a flat line at 110 meV.

Finally, measurements with an \( E_{\text{i}} \) of 500 meV were used to capture the top of the magnon band. Figure 10 shows the results of the spin wave model and the corresponding simulation. The measurements reveal magnetic scattering higher in E than is described by the simulation. There are two possibilities to explain this excess scattering. First, the exchange parameters as predicted by the DFT calculations could be too low. Second, there could be another excitation above the magnetic excitations described above. Close examination of spin-dependent DFT calculations in Chen et al.[28] indicates there might be some nested orbitals around the M point which could be consistent with the observed scattering. Nev-
FIG. 8. A constant $E$ cut of the $E_i=100$ meV data shown in 6b is indicated by the black circles. The integration range is between 40 and 70 meV. The same analysis procedure was carried out on results from McVine Monte Carlo ray tracing simulations that included the theoretical model for the sample. The results from the simulation are given by the red line. The error bars for the simulations arise from statistical uncertainty in the Monte Carlo method and are significantly less than those of the measurement.

Nevertheless, further DFT calculations and measurements with single crystal samples will be needed to definitively distinguish these two possibilities.

IV. SUMMARY AND CONCLUSIONS

The geometry of intersecting \{111\} kagome planes describing the fcc kagome lattice offers a rare example to study a truly 3D frustrated kagome antiferromagnet, as exemplified by IrMn$_3$. The so-called $q=0$ triangular spin structure is preserved in the 3D system along with some of the degeneracies of the NN Hesienberg model which stimulated previous work in the 2D case. A focus of the present results is to quantify the role of further-neighbor exchange interactions within a local-moment model of spin wave excitations in IrMn$_3$, extending our
FIG. 9. The points are series of cuts from the $E_i = 300$ meV data. The solid curves are simulations using the described dispersion with all four exchange interactions included. Broken lines are from a model with $J_4 = 0$. There is a curve, and its corresponding simulation, for each incident energy which are offset for clarity. The same scale factor is applied to all of the curves and an individual sloping background is fit to account for varying backgrounds. Only three curves of the $J_4 = 0$ case are shown as they are sufficient to illustrate that inclusion of $J_4$ results in superior agreement with the data.

earlier work that involved only NN exchange.\textsuperscript{13} Earlier DFT calculations\textsuperscript{10,12} indicate that exchange up to fourth nearest neighbors are substantial in this transition-metal ion conductor that has been widely used as the exchange pinning antiferromagnetic thin film which is integral to spin valve technology. The model Hamiltonian also includes a cubic anisotropy term, which is predicted to be strong from the DFT theory, and induces a gap in the model spectrum.
FIG. 10. Data (a) and simulation (b) of the $E_i = 500$ meV configuration. The results indicate that the measurement shows additional scattering above the top of the band as compared to the model. Also, the high-energy magnetic scattering seems to decrease with $Q$ faster in the experimental data as compared to the simulation.

New inelastic neutron scattering data on powder samples is also presented and compared with corresponding results from the model. The comparison is facilitated through the use of ray tracing simulations that integrate model predictions with instrumental resolution as well as additional features of the sample, which in this case includes a small amount of magnetic scattering from a MnO impurity. There is also a substantial amount of IrMn$_3$ phonon scattering in our data that was not included in the simulations.

Model calculations show that the presence of a strong inelastic feature above the (110) magnetic peak, which is observed in the SEQUOIA data, is only consistent with a model including further neighbor interactions. This fact provides strong support for the conclusion that further-neighbor interactions are required to successfully model the spin wave spectrum of IrMn$_3$. Good agreement between the model and data in the case of constant energy cuts supports the conclusion that the DFT estimates of the exchange interactions are reliable.

The present work provides the first experimental evidence for the existence of the DFT
prediction of strong cubic anisotropy in IrMn$_3$ through the good agreement between the
spin-wave model and a gap observed in the inelastic spectrum. This is an important result
as our earlier work demonstrated that anisotropy induces a uniform magnetic moment along
[111] directions. This likely indicates an important mechanism for exchange coupling to
adjacent ferromagnetic thin films. Such a moment could also be employed to induce a single
magnetic domain in field cooled single-crystal samples.

Evidence also exists in the data for broad high energy (at and above 200 meV) magnetic
scattering that does not appear in the local-moment model. This feature deserves further
investigation in view of the possibility it is associated with the calculated band structure
supporting the anomalous hall effect in such non-collinear antiferromagnets.

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