Magnetic order and single-ion anisotropy in Tb$_3$Ga$_5$O$_{12}$

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(Received 22 February 2019; published 26 September 2019)

Terbium gallium garnet (TGG), Tb$_3$Ga$_5$O$_{12}$, is well known for its applications in laser optics but also exhibits complex low-temperature magnetism that is not yet fully understood. Its low-temperature magnetic order is determined by means of time-of-flight neutron powder diffraction. It is found to be a multiaxial antiferromagnet with magnetic Tb$^{3+}$ ions forming six sublattices of magnetic moments aligned parallel and antiparallel to the (100) crystallographic directions of the cubic unit cell. The structure displays strong easy-axis anisotropy with respect to a twofold axis of symmetry in the local orthorhombic environment of the Tb$^{3+}$ sites. The crystal-field splitting within the single-ion ground-state manifold is investigated by inelastic neutron scattering on powder samples. A strong temperature dependence of the quasidoublet ground state is observed and revised parameters of the crystal-field Hamiltonian are given. The results of bulk magnetic susceptibility and magnetization measurements are in good agreement with values based on the crystal-field model down to 20 K, where the onset of magnetic correlations is observed.

DOI: 10.1103/PhysRevB.100.094442

I. INTRODUCTION

Many current studies of the garnet family focus on technological applications of Y$_3$Al$_5$O$_{12}$ and Y$_3$Fe$_5$O$_{12}$ (in optoelectronics [1] and magnonics [2–4], respectively), but the garnet structure also contains two interpenetrating half-garnet lattices (Fig. 1)—twisting spatial arrangements of corner-sharing triangles—which are of considerable interest for highly frustrated magnetism. An individual half-garnet or hyperkagome lattice, along with the pyrochlore lattice, are candidates for the construction of three-dimensional Coulomb phases [5]. There are two series of rare earth garnets, $R_3$Al$_5$O$_{12}$ and $R_3$Ga$_5$O$_{12}$ (where $R$ is a trivalent rare earth ion), and, as in rare earth pyrochlores [6], competition between exchange and dipolar interactions [7,8] and crystal field anisotropy produces contrasting properties throughout the series, often with highly anisotropic magnetic order at low temperature [9–15]. Gd$_3$Ga$_5$O$_{12}$ (GGG) [16] is the only fully frustrated garnet in which no long-range magnetic order has been observed to the lowest temperature [10,13,17], and recent investigations discovered cooperative magnetic multipolar degrees of freedom underlying the spin correlations of the low temperature phase of GGG [18].

In TGG, the non-Kramers ion Tb$^{3+}$ has a quasidoublet ground state composed of two closely spaced singlets as a result of the orthorhombic crystal electric field (CEF) [20]. Thus the magnetic properties of TGG are strongly anisotropic, and long range order is expected to be of the induced-moment type. Highly anisotropic magnetic properties, including step-like magnetization curves, demonstrate the importance of single-ion crystal field effects [20–22]. Rare earth garnets with noncollinear Ising anisotropy and dominant dipolar interactions are expected to order with a magnetic structure formed of interpenetrating antiferromagnetic chains running along the (100) directions [7,10,13]. TGG orders with this structure at $T_N \approx 0.25$ K [23], but it has been suggested that hyperfine interactions must be incorporated to explain the observed $T_N$ [14,15,23]. The existence of a correlated phase for $T > T_N$ has not been examined.

Moreover, at higher temperatures, in addition to its well-known large Verdet constant and transparency, which make it a suitable material for Faraday rotators and optical isolators [24–26], TGG also exhibits an acoustic Faraday effect [27,28], Cotton-Mouton [29], and a thermal Hall effect [30,31] (for which it is the prototypical system), suggestive of important spin-lattice interactions. Despite several attempts [32–34], the thermal Hall effect lacks an unequivocal microscopic description. Detailed knowledge of the magnetic interactions and crystal field wave functions in TGG plays a crucial role as a starting point required to understand any frustrated phase and the ordering mechanism that terminates it, as well as an eventual understanding of the magnetoelastic and magnetothermal effects.

Here, results of our studies on powder samples of TGG are presented. Neutron and x-ray diffraction were used to
determine details of the crystalline and magnetic structures (Sec. III A) that are fundamental for any further considerations. Inelastic neutron scattering (INS) was employed to observe crystal electric field transitions (Sec. III B) and this information was used to refine the parameters of the CEF Hamiltonian. The results of bulk magnetization measurements are compared to the calculated forms determined by the CEF model (Sec. III C). Discussion of the results is followed by remarks as to the outlook for future single crystal studies.

II. METHODS

A. Sample preparation

A single crystal in the form of a cylindrical rod (\( \varnothing = 6 \) mm and \( l \approx 50 \) mm) cut out of an ingot grown by the Czochralski method was acquired from FEE (Idar-Oberstein, Germany) and crushed to a fine powder in an agate pestle and mortar.

B. Synchrotron x-ray powder diffraction

For a precise determination of structural parameters, a high-resolution synchrotron x-ray powder diffraction (SXRPD) experiment was performed on the Materials Science Beamline at the Swiss Light Source (Paul Scherrer Institute, Villigen, Switzerland). The powder sample was loaded in a borosilicate glass capillary and a diffraction pattern was taken at room temperature (298 K). The \( \varnothing = 0.1 \) mm diameter of the capillary in combination with photon energy \( E = 22 \) keV (\( \lambda = 0.563 \) Å) were chosen to minimize the sample absorption.

C. Neutron time-of-flight powder diffraction

Neutron diffraction patterns were collected on the WISH time-of-flight diffractometer at ISIS neutron and muon facility (Rutherford-Appleton Laboratory, Harwell, UK). A dilution refrigerator insert was used to cool the sample below \( T_N \). For this procedure \( \sim 5.5 \) g of powder was compacted in a \( \varnothing = 5 \) mm copper can. In order to enable faster thermalization of the sample, the can was indium sealed and equipped with a capillary which allowed for the introduction of \(^4\)He exchange gas during the cooling process. This is particularly important given the low thermal conductivity of garnets at \( T < 1 \) K [35], the slow thermalization of their magnetic structures, and the low \( T_N \) of TGG. The sample was held at base temperature, \( T \approx 0.03 \) K for \( \sim 12 \) hours before diffraction measurements commenced. During initial runs, the solid methane moderator was above its intended operating temperature, which modifies the wavelength spectrum and reduces the neutron flux of the instrument, but subsequently diffraction patterns were recorded at base temperature \( (T_{\text{base}} \approx 0.03 \text{ K} ) \), 1.5, 5, and 15 K. The appearance of magnetic Bragg reflections indicated the thermalization of at least part of the sample below \( T_N \), but a significant decrease in the intensity of the magnetic peaks with concurrent development of diffuse scattering around the Bragg positions was observed during the collection of the base temperature diffraction patterns once the moderator reached full performance. Since the changes occurred only concurrently with the exposure of the sample to the neutron beam, they were attributed to beam heating. This implies that the measured low temperature data are an average over a range of temperatures, and the full low temperature ordered moment cannot be accurately determined. The collected patterns were analyzed by the Rietveld and the LeBail refinement methods with use of the FULLPROF suite [36].

D. Inelastic neutron scattering

Time-of-flight neutron spectroscopy experiments were conducted on the MARI (direct geometry) and IRIS (indirect geometry) time-of-flight spectrometers at ISIS (Rutherford-Appleton Laboratory, Harwell, UK). For MARI, 7.3 g of powder was wrapped in an aluminium foil sachet and placed in a \( \varnothing = 45 \) mm Al can. Spectra with \( E_i = 150, 75, 50, \) and \( 12 \) meV were recorded at \( T = 5 \) K and \( T = 100 \) K, using a Fermi chopper with a gadolinium coated slit package. The best energy resolution at energy transfer \( E \approx 5 \) meV was obtained with \( E_i = 12 \) meV, where \( \Delta E \approx 0.19 \) meV. For IRIS, the same amount of sample, also wrapped in an Al sachet, was placed in a \( \varnothing = 24 \) mm can. Datasets were collected with the PG(002) and PG(004) analyser setups. The former allows for measurement within an almost symmetric, narrow dynamical range \(( -0.55 < E < 0.57 \text{ meV}) \) and its excellent resolution of 17.5 \( \mu \text{eV} \) at the elastic line was used to investigate the components of the ground-state quasidoublet. The latter setup expands the dynamical window on the neutron energy loss side \(( \sim 2.1 < E < 15.5 \text{ meV}) \), providing a 54.5 \( \mu \text{eV} \) resolution at the elastic line, thus enabling observations of CEF excitations at higher energies. The measurements were conducted at temperatures in range \( 1.8 < T < 100 \) K.
TABLE I. Structural parameters obtained by Rietveld refinement of SXRPD pattern measured at room temperature. Tb2 stands for superstoichiometric Tb$^{3+}$ ions at the 16a gallium position.

| Atom | Wyckoff position | x    | y    | z    | Occupancy | $B_{iso}$ (Å$^2$) |
|------|------------------|------|------|------|-----------|-------------------|
| Ga1  | 16a              | 0    | 0    | 0    | 1.969(3)  | 0.217(5)          |
| Tb2* | 16a              | 0    | 0    | 0    | 0.031(3)  | 0.217(5)          |
| Tb   | 24c              | 1/8  | 0    | 1/4  | 3         | 0.239(1)          |
| Ga2  | 24d              | 3/8  | 0    | 1/4  | 3         | 0.283(3)          |
| O    | 96h              | 0.286(1) | 0.0566(1) | 0.6500(1) | 12         | 0.23(2)           |

E. Magnetization

Magnetic susceptibility data were collected on a small portion of the same powder sample ($m = 95.6$ mg), in an applied magnetic field of 0.01 T, using a Quantum Design MPMS XL-7 superconducting quantum interference device (SQUID) magnetometer. Using the same device, the magnetization was measured with a smaller sample ($m = 8.15$ mg) of fields up to 60 kOe and temperature range 1.8 $< T < 300$ K.

III. RESULTS

A. Crystalline and ordered magnetic structure

TGG crystallizes in the body-centred cubic $Ia\overline{3}d$ space group (No. 230). The Tb$^{3+}$ ions occupy the 24c Wyckoff positions (with point group $D_2$ or 222), forming two half-garnet sublattices (Fig. 1). The lattice constant of TGG at room temperature, determined from a Rietveld refinement of the x-ray diffraction pattern, is $a = 12.35225(1)$ Å. The presence of superstoichiometric Tb$^{3+}$ ions in Czochralski-grown TGG has been discussed in the literature [34]. Because of the large contrast between Ga$^{3+}$ and Tb$^{3+}$ in x-ray diffraction, Rietveld refinement enabled a precise determination of the stoichiometry of our sample, which was found to be Tb$_{3.031(3)}$Ga$_{4.969(3)}$O$_{12.0}$. (The small difference between the coherent neutron scattering lengths of Tb and Ga does not allow for cross-verification by neutron diffraction [37]). Superstoichiometric Tb$^{3+}$ ions were found exclusively at the octahedral Ga site 16a (with trigonal point group $C_3$ or 3).

Attempts to refine a population of terbium ions on other sites were found [39] (not shown, no unexpected changes of parameters or misfits from the x-ray experiment against the data measured at 15 K). All of the magnetic Bragg peaks can be indexed by the propagation vector $\mathbf{k} = (0, 0, 0)$.

Using the ISOTROPY package [38], eight irreducible representations of the space group and the propagation vector were found [39] ($2 \times 1$-dimensional: $m\Gamma_1^+, m\Gamma_3^{-}$; $2 \times 2$-dimensional: $m\Gamma_1^+, m\Gamma_3^{-}$; $4 \times 3$-dimensional: $m\Gamma_4^+, m\Gamma_5^{-}$, $m\Gamma_6^+ \text{ and } m\Gamma_7^{-}$; in Miller-Love notation). Their magnetic moment basis functions were projected out and combined with corresponding order parameter directions. These generated 14 possible sets of basis vector directions. 11 sets were immediately excluded as they are incompatible with the presence or absence of certain magnetic Bragg peaks in the experimental diffraction pattern, i.e., the diffraction data show the presence of $(1, 1, 0)$ antiferromagnetic and absence of $(2,0,0)$ ferromagnetic reflections (Fig. 3). Rietveld refinement of the remaining possibilities, and examination of the relative intensities of the observed reflections, allowed us to unambiguously assign the one-dimensional irreducible representation $m\Gamma_2^+$ to the observed pattern (Fig. 3). This structure has the magnetic space group $Ia\overline{3}d'$. The scale factor of the Rietveld refinements was set by refining the room temperature structural model obtained from the x-ray experiment against the data measured at 15 K (not shown, no unexpected changes of parameters or misfits from the x-ray experiment against the data measured at 15 K).

![FIG. 2. Rietveld refinement of synchrotron x-ray powder diffraction pattern. Green vertical lines mark the positions of Bragg reflections.](image)

![FIG. 3. Rietveld refinement of magnetic reflections in the neutron powder diffraction difference pattern, produced by the $m\Gamma_2^+$ irreducible representation of $Ia\overline{3}d'$, corresponding to $Ia\overline{3}d'$ magnetic space group. The observed reflections are indexed and can be seen to correspond to the propagation vector $\mathbf{k} = (0, 0, 0)$.](image)
FIG. 4. Ordered magnetic structure of TGG. (a) and (b), red arrows show the magnetic moments for a selection of the garnet sites from each of the two hyperkagome sublattices. For sites where magnetic moments are not shown, their anisotropy can be inferred by the ones shown on those triangles of the same sublattice that lie in parallel planes. [Such triangles also have the same color shading by the ones shown on those triangles of the same sublattice that lie sites from each of the two hyperkagome sublattices. For sites where arrows show the magnetic moments for a selection of the garnet lattice, making them individually antiferromagnetic. In the course of the magnetic intensity refinement, the resemblance to the calculated form factor less good in the high-\[\bar{Q}\] region of the data. Retrieving a similar dependence from the \(E \approx 35\) meV feature shown in Fig. 6(c) was not possible due to the presence of an optical phonon branch at \(E \approx 40\) meV, but excited state transitions to these levels that follow the magnetic form factor were observed in the high temperature spectra, confirming their CEF nature and energies.

To ensure that all of the excitations we consider in our fitting are indeed crystal field excitations, their \(\bar{Q}\) dependence was analyzed. The energy-integrated intensity of the quasidoublet and the first two groups above it (i.e., \(E \approx 5, 28\) meV) closely follow the dipole magnetic form factor of Tb\(^{3+}\) ion \([42]\), as shown in Fig. 7. Phonon contributions make the resemblance to the calculated form factor less good in the high-\[\bar{Q}\] region of the data. Retrieving a similar dependence from the \(E \approx 35\) meV feature shown in Fig. 6(c) was not possible due to the presence of an optical phonon branch at \(E \approx 40\) meV, but excited state transitions to these levels that follow the magnetic form factor were observed in the high temperature spectra, confirming their CEF nature and energies. The excitation that can be seen at \(E \approx 18\) meV in Fig. 6(c) was found to follow the \(\bar{Q}\)\(^2\) dependence expected of a phonon branch.

To refine the parameters of the crystal field Hamiltonian, we require the energies of the crystal field excitations.
FIG. 6. Powder neutron scattering spectrum of TGG at $T = 5$ K measured by means of (a) IRIS, (b) and (c) MARI spectrometers with (a) PG (002) analyzer, (b) $E_i = 12$ meV, and (c) $E_i = 50$ meV. Intensities of inelastic features in all panels were normalized to 1. Polygons drawn with a dashed line mark the integration ranges of data presented in Fig. 8.

Extracting these quantities is complicated when the separation of the excitations is close to the instrumental resolution, especially when the excitations are broadened beyond the resolution limit, and because of the close spacing of the two members of the quasidoublet. The energy difference of $\Delta = 0.22$ meV between the two states of the quasidoublet means both levels are similarly populated at $T = 5$ K ($F_1/F_0 = e^{\Delta/kT} = 0.57$), and contributions to the spectra by excitations from both levels could be comparable.

FIG. 7. $|\vec{Q}|$ dependence of the energy-integrated intensity of the observed features normalized to 1 at $|\vec{Q}| = 1$. Blue and cyan markers show the transitions from ground state measured at low temperature. Green markers show excited state transitions from levels at $E \approx 28$ meV that appear at 100 K (Fig. 9). Black dots mark the quasidoublet transition intensity (Fig. 10). The red dashed line shows the magnetic form factor of the Tb$^{3+}$ ion. The form factor is scaled and a small flat background is added to fit the low-$|\vec{Q}|$ data of low energy excitations (blue markers).

We integrated the data in ranges indicated in Fig. 6, using simple boxes for the levels at $E \approx 5, 28$ meV, and a parallelogram for the levels at $E \approx 35$ meV. The resulting spectra are shown in Fig. 8. For the feature at $E \approx 5$ meV [Fig. 8(a)], positions of single transition lines cannot be resolved using either spectrometer. Previous investigations of the TGG crystal field spectrum using Raman scattering [43] identified this feature as containing four excitations from the ground state. We therefore fitted four Gaussian peaks constrained to have a common width ($FWHM = 0.85(3)$ meV). A comparison between the reported transition energies [43] and the values deduced from the fits is made in Table II, and shows good agreement between both datasets. At $E \approx 28$ meV [Fig. 8(b)] two distinct peaks are observed, with a separation well exceeding the splitting of the ground-state singlets, suggesting they are separate levels. Both peaks were fitted with Gaussian peaks ($FWHM = 1.93(7)$ meV). The group at $E \approx 35$ meV [Fig. 8(c)] was fitted with four more Gaussians ($FWHM = 0.55(5)$ meV). We note that the widths of the measured features are considerably broader than the instrumental resolution, with the exception of the members of the group lying around 35 meV.

### TABLE II. CEF transition energies observed by Raman spectroscopy [43], calculated with parameters given in Ref. [20], determined by fits to INS data [Figs. 8(a), 8(b) and 8(c)] and calculated with the refined parameters (Table III). All values are given in meV.

| $E_{obs}$ Raman [43] | $E_{calc}$ [20] | $E_{obs}$ INS | $E_{calc}$ |
|----------------------|----------------|----------------|----------------|
| 0.4                  | 0.219(1)       | 0.22           |
| 4.2                  | 4.52(2)        | 4.50           |
| 5.3                  | 5.23(3)        | 5.25           |
| 6.2                  | 6.04(4)        | 6.01           |
| 6.6                  | 6.51(3)        | 6.54           |
| 36.6                 | 26.30(4)       | 26.60          |
| 38.6                 | 28.38(5)       | 28.27          |
| 44.6                 | 34.16(8)       | 34.26          |
| 46.7                 | 34.85(5)       | 34.73          |
| 49.9                 | 35.59(5)       | 35.63          |
| 50.7                 | 36.31(5)       | 36.38          |
| 55.4                 | 38.20          | |
Changes in the measured spectra between \( T = 5 \text{ K} \) and \( T = 100 \text{ K} \) are shown in Fig. 9(a) and 9(b). The most striking difference is the appearance of strong intensities at \( E \approx 21.29 \text{ meV} \), which follow the magnetic form factor of Tb\(^{3+} \) (Fig. 7), and can therefore be identified as transitions from thermally populated levels at \( E \approx 5 \text{ meV} \) [Fig. 9(c)], consistent with the assignment of excitations at \( E \approx 35 \text{ meV} \), whose form factor could not be verified, as a crystal field excitation.

2. Crystal electric field Hamiltonian

The crystal-field Hamiltonian for \( D_2 \) point-group symmetry is given by

\[
\hat{H}_{\text{CEF}} = \tilde{B}_0^0 \hat{O}_0^0 + \tilde{B}_2^0 \hat{O}_2^0 + \tilde{B}_4^0 \hat{O}_4^0 + \tilde{B}_2^2 \hat{O}_2^2 + \tilde{B}_4^2 \hat{O}_4^2 + \tilde{B}_2^4 \hat{O}_2^4 + \tilde{B}_4^4 \hat{O}_4^4,
\]

where \( \hat{O}_q \) are the (cosine) Stevens operators \( \hat{O}_q^\pm(\mathbf{J}) \equiv \hat{O}_q^\pm(\mathbf{J}_\parallel, \mathbf{J}_\perp) \). The crystal-field coefficients \( \tilde{B}_q^k \) can be found in standard literature about the Stevens’ equivalent method [44–46].

We can summarize the general expressions of the CEF eigenstates as

\[
\begin{align*}
|\Gamma_1\rangle &= \sum_{M_J = 2, 4, 6} b_M (|M_J\rangle + |-M_J\rangle) + b_0 |0\rangle, \\
|\Gamma_2\rangle &= \sum_{M_J = 1, 3, 5} a_M (|M_J\rangle - |-M_J\rangle), \\
|\Gamma_3\rangle &= \sum_{M_J = 1, 3, 5} a'_M (|M_J\rangle + |-M_J\rangle), \\
|\Gamma_4\rangle &= \sum_{M_J = 2, 4, 6} b'_M (|M_J\rangle - |-M_J\rangle),
\end{align*}
\]

where all coefficients \( a_M, b_M, a'_M, b'_M \) are real, and the \( |M_J\rangle \equiv |J = 6, M_J\rangle \) are eigenstates of the \( \hat{J}_z \) operator. The generic states \( |\Gamma_n\rangle \) are labeled according to their irreducible representation \( \Gamma_n \), which we deduce by comparison with character tables.

Previous analyses of isostructural Tb\(_3\)Al\(_5\)O\(_{12}\) (TAG) have attempted to simplify the CEF scheme by retaining cubic or tetragonal symmetries (reducing the number of operators to four or five, respectively), with the orthorhombic terms expected to produce only a small modification of the parameters. We find that such approaches cannot describe our experimental data, and so we use the orthorhombic Hamiltonian throughout the following. This is in contrast to other RE garnets (see, for example, Refs. [47,48]) where the use of higher symmetries has been successful.

Refinement of the nine crystal-field parameters \( \tilde{B}_q^k \) of Eq. (1) was attempted with the use of the SPECTRE software [49]. The refined quantity was the transition energies (and not the intensities), retrieved from fitting to the INS spectra. Refinement of values previously published for TGG [20], from the results of Raman spectroscopy measurements of a small number of levels (see Table II) and single crystal magnetization, did not lead to satisfactory results. Several other sets of initial parameters were tested (Dy [50], Er [51], Eu [52], and Nd [53] gallium garnets scaled accordingly to the differences between radial momenta of RE\(^{3+}\) and Tb\(^{3+}\) ions, and parameters for Tb\(_n\) in YAG [54]). We have made attempts of refinement with sets of basis functions limited to the \( ^7F_0 \) ground-state multiplet (LS coupling), \( ^7F_N (N \in [0, 6]) \) states lying between 253 and 698 meV [55] (intermediate coupling), and including all higher \( ^{2S+1}L_J \) terms. Although good agreement between calculated and observed energies was found in the two latter cases (twofold reduction of goodness of fit parameter when going beyond LS coupling and without any further improvement when expanding the basis more), a proper convergence of this multiparameter refinement was not achieved. The previously listed initial sets of parameters together with some of the best final sets refined with SPECTRE were introduced into a custom least-squares fitting routine refining the eigenvalues of the Stevens Hamiltonian [Eq. (1)] (i.e., restricted to LS coupling). The algorithm reached convergence and the results ranged from good to excellent agreement with the measured values. The optimal solution was chosen by investigating the neutron spectra calculated using the refined parameters and comparing them with INS data (Fig. 9) and the set giving the closest distribution of calculated intensities with respect to the observed one was chosen (based on the result

![FIG. 9. (a) \(|\tilde{Q}|\)-integrated (\(|\tilde{Q}|\)) = 2.25 ± 1.25 Å\(^{-1}\)) excitation spectra measured on MARI with \( E_i = 12 \text{ meV} \). (b) \(|\tilde{Q}|\)-integrated (\(|\tilde{Q}|\)) = 4 ± 2 Å\(^{-1}\)) excitation spectra measured at MARI with \( E_i = 50 \text{ meV} \). At \( T = 100 \text{ K} \) two features appear with \( |\tilde{Q}|\) dependence following the magnetic form factor (Fig. 7). Their energy correspond to the transitions, represented by cyan and magenta arrows, in the schematic CEF excitation spectrum shown in (c). Dashed lines present the neutron scattering intensities calculated within dipole approximation convoluted with Gaussian peaks. For illustrative purposes we use the FWHM retrieved from resolution calculations for MARI with proper chopper setup. Vertical lines mark the energies of expected transitions from thermally populated states having nonzero intensities.](https://doi.org/10.1103/PhysRevB.100.094442)
of the initial EuGG refinement with SPECTRE). This result was also characterized by the best value of the goodness of fit parameter $\chi^2 = 2.1 \times 10^{-3}$. The contributions to the wave functions in the $|M_J\rangle$ basis in Table V is given in Appendix B.

The neutron scattering intensities calculated within the dipole approximation using refined CEF parameters for $T = 5$ K and $T = 100$ K are plotted along with experimental results in Fig. 9(a) and 9(b). The calculated positions of the excitations are in good agreement with measured intensities. The shapes of the features at higher energies are reasonably reproduced by the calculations. In addition, the almost nonexistent scattering intensity calculated for the transition between the ground-state and the highest level of the lowest multiplet is consistent with it not being observed in the experimental data.

When compared with previous determinations of the crystal field parameters, i.e., Ref. [20] (see Table III), we find reasonable agreement between most parts of the Stevens parameters, but there are differences in the size and sign, particularly for the highest order parameters. As can be seen in Table II, rather few of the levels were previously determined experimentally, with the results that the parameters are not very accurate. Indeed, when the full level scheme is calculated using the parameters of Ref. [20], we can see that the higher levels are predicted at much higher energies than we observe them (see Table II for comparison).

Nonetheless, these previously determined parameters and eigenfunctions could be used successfully to reproduce a step-like magnetization curve measured on a single crystal sample at 4.2 K [20]. However, all of the parameter sets obtained in the course of our refinements give a similar structure of the wave functions. They also describe the magnetization step, which is a rather generic feature present for similar compositions of the quasidoublet eigenstates, and one of the levels at $E_a \approx 5$ meV. The significant contribution from high-$M_J$ states to this higher level causes, due to the Zeeman term, its rapid descent in field towards the energies of the quasidoublet states leading to increase in its thermal population and contribution to the overall magnetization. By comparison of the quasidoublet wave functions obtained from our parameter set to the one obtained from the parameter set in Ref. [20], we find that the symmetry character $|\Gamma_3\rangle$ and $|\Gamma_1\rangle$, of the ground singlet, $|p\rangle \equiv |\psi_0\rangle$, and the first-excited one, $|q\rangle \equiv |\psi_1\rangle$, are swapped. More explicitly, $|p\rangle \equiv |\psi_0\rangle \approx -0.09|\psi_0\rangle + 0.03|\psi_1\rangle + 0.71|\psi_2\rangle$ and $|q\rangle \equiv |\psi_1\rangle \approx 0.18|\psi_0\rangle + 0.12|\psi_1\rangle - 0.33|\psi_2\rangle - 0.83(0)$ in this work (Table V in Appendix B), whereas in Ref. [20] we have $|p\rangle \approx -0.05|\psi_0\rangle + 0.17|\psi_1\rangle - 0.16|\psi_2\rangle - 0.94(0)$ and $|q\rangle \approx -0.06|\psi_0\rangle - 0.15|\psi_1\rangle + 0.69(1)$. The order of the $|\Gamma_3\rangle/|\Gamma_1\rangle$ states was found to play an important role in obtaining a good reproduction of the transition intensities in INS.

On the other hand, at $E \approx 5$ meV, there is a strong discrepancy between the calculated and observed line widths, which are much larger than the instrumental resolution. Furthermore, although a single transition between the members of the quasidoublet can be seen at $E = 0.22$ meV at $T = 5$ K (though the peak is again much broader than the instrumental resolution), the lowest energy part of the spectrum displays a very strong temperature dependence. At the lowest measured temperature $T = 1.8$ K the $S(|\bar{Q}|, \omega)$ map of this part of the spectrum [Fig. 10(a)] suggests that the single excitation is replaced by a band of dispersive excitons. A clearly visible contribution of two or more peaks to the first transition has developed, as shown in Fig. 10. On the high energy tail of the main peak is another weak peak at $E \approx 0.45$ meV that develops with the same temperature dependence as the multi-peak structure of the main excitation. A shoulder can be distinguished in the quasidoublet gap at $T < 3.5$ K, which suggests another peak. This observation, along with the large widths of other excitations (i.e., $|\psi_2\rangle$ with $E_a \approx 5$ meV), suggests interactions between Tb$^{3+}$ ions, or coupling with lattice excitations modifies the excitation spectrum beyond a single-ion picture.

**TABLE III.** Set of refined crystal-field parameters giving the best value of standard goodness of fit parameter, $\chi^2 = 2.1 \times 10^{-3}$. Parameters are given in Stevens notation.

| Parameter | Ref. [20] (meV) | This work (meV) |
|-----------|----------------|-----------------|
| $E^2_{1g}$ | $5.05 \times 10^{-2}$ | $2.66 \times 10^{-2}$ |
| $E^2_{1u}$ | $-2.60 \times 10^{-1}$ | $-3.24 \times 10^{-1}$ |
| $E^2_{2g}$ | $-4.10 \times 10^{-3}$ | $-2.47 \times 10^{-3}$ |
| $E^2_{2u}$ | $3.00 \times 10^{-3}$ | $3.70 \times 10^{-3}$ |
| $E^2_{4u}$ | $1.50 \times 10^{-2}$ | $8.51 \times 10^{-3}$ |
| $E^2_{6g}$ | $-5.39 \times 10^{-6}$ | $1.12 \times 10^{-5}$ |
| $E^2_{6u}$ | $1.36 \times 10^{-5}$ | $-1.21 \times 10^{-5}$ |
| $E^2_{8u}$ | $-1.09 \times 10^{-4}$ | $-4.73 \times 10^{-5}$ |
| $E^2_{8u}$ | $5.32 \times 10^{-7}$ | $8.96 \times 10^{-5}$ |

**FIG. 10.** (a) $S(|\bar{Q}|, \omega)$ of the low-lying CEF singlet measured on IRIS at $T = 1.8$ K. Hints of dispersive character of the measured excitation can be observed. Intensity of inelastic feature normalized to 1 on logarithmic scale. (b) Temperature dependence of $|\bar{Q}|$-integrated ($|\bar{Q}| = 0.4-1.8$ Å$^{-1}$) cuts through the ground-state quasidoublet.
the calculated results deviate strongly from the experimental data. Similarly, the collapse of the $M(H/T)$ curves that is expected for paramagnetic spins can be observed down to 50 K. For $T < 50$ K deviation from this scaling, coinciding with the departure of the susceptibility from the Curie-Weiss law, becomes increasingly pronounced [inset of Fig. 11(a)]. These behaviors are consistent with the onset of a correlated cooperative paramagnetic phase. Our measured susceptibility and magnetization agree well with another recent report [56].

IV. DISCUSSION

We have investigated the crystal and magnetic structures of TGG, together with the crystal field Hamiltonian for its single-ion physics. These are all relevant for its low temperature physics and magnetothermal effects.

Firstly, the crystal structure of our samples is close to the ideal garnet structure, but there are superstoichiometric terbium ions (Tb2). These Tb$^{3+}$ ions in excess act as defects which enhance the scattering of phonons and cause the lowering of the thermal conductivity [31]. For this reason, in Czochralski-grown crystals, where their abundance goes up to $\approx 1\%$, they are considered responsible for the manifestation of the thermal Hall effect [35]. Such concurrence—Czochralski-grown crystals have the highest percentage of Tb2 defects and lowest diagonal thermal conductivity—motivated the theoretical work of Ref. [34], in which Mori et al. posit that the resonant skew scattering of phonons from the CEF states of superstoichiometric Tb$^{3+}$ ions is the origin of the thermal Hall effect. In this theory two assumptions are crucial: (i) the Tb2 ions are somewhat correlated in space, (ii) the CEF splitting of their lowest levels (labelled as $|a\rangle$, $|b\rangle$ in Ref. [34]) is comparable with $\Delta$, the CEF gap for the quasidoublet states, $|\rho\rangle$, $|q\rangle$, of the regular Tb sites.

Our study provides insights to this theory, from both structural and single-ion perspectives. We detect superstoichiometric terbium ions only on the 16a (Ga1) Wyckoff position, where their concentration is 1.6%. The importance of this

C. Magnetization

The magnetic susceptibility is presented in Fig. 11(a). Above $T = 50$ K it can be fitted by a Curie-Weiss law with $\theta_{\text{CW}} = -7.98$ K (hence the frustration parameter $f = |\theta_{\text{CW}}|/T_{\text{N}} \approx 32$). By means of $\mu_{\text{eff}} = \sqrt{\frac{3\mu_B^2}{N_A k_B}} \chi T$, we extracted $\mu_{\text{eff}} = 9.913(3) \mu_B$ as paramagnetic moment per Tb$^{3+}$ ion over the range $260 < T < 300$ K. Such value differs by only 2% from $\mu_{\text{qTb}^3} = 9.72 \mu_B$, the theoretical paramagnetic moment per Tb$^{3+}$ ion in the $^7F_6$ ground multiplet [46]. Calculation of the susceptibility using the refined crystal-field parameters (Table III) gives excellent agreement with the experimental data.

The magnetization per Tb$^{3+}$ ion as a function of applied field $H$ for different temperatures, and in terms of the ratio $H/T$ is plotted in Fig. 12(a) and 12(b), respectively. Solid lines in panel (a) show the result of calculations using the refined CEF parameters (Table III), which are in good agreement with the data down to 20 K. Below this temperature,

![Image](https://example.com/image.png)
observation is that the 16$^{a}$ position has a $C_{3v}$ symmetry, which is higher than the $D_2$ and $S_3$ symmetries, respectively, for the 24$c$ (Tb) and 24$d$ (Ga2) sites (see Table I). For $J = 6$, five singlets and four doublets are expected in $C_{3v}$ symmetry, and the assumption that the crystal field scheme at this site will be generally similar to that at the main terbium site is to be examined carefully. Results from point charge calculations (which include the six nearest oxygen ions) suggest a similar structure of the quasidoublet ground state with four symmetry-allowed doublets lying at energies higher than 90 meV. However, one has to keep in mind the limited reliability of this method. Regarding the spatial correlations of Tb2 ions and their crystal field scheme, direct experimental probing is hindered by the low percentage of superstoichiometric abundance in TGG.

The low-temperature magnetic structure of TGG that we have observed is a multiaxial antiferromagnet composed of magnetic moments that have noncollinear local anisotropy axes. Any singlet $|\psi_n\rangle$, from the CEF spectrum of the Tb$^{3+}$ ion in TGG, is characterized by $\langle \psi_n | J | \psi_n \rangle = 0$ [where $J = (J_x, J_y, J_z)$ are the angular momentum operators], a consequence of the complete removal of the degeneracy of the free ion states by the crystal field [45,57]. The magnetic moments found in the ordered phase of TGG must therefore be induced by perturbations which couple or mix the CEF ground singlet with higher CEF eigenstates, i.e., an induced moment system. Usually this mixing is achieved by the exchange interactions, but a curious feature of the magnetic structure is that it nullifies the nearest neighbor Heisenberg terms $S_i \cdot S_j$, since the spin expectation values of induced moments $S_i$ are expected to be orthogonal [see below Eq. (4)]. Our measurements of the magnetic susceptibility and magnetization suggest nonetheless that antiferromagnetic correlations build up below $T \approx 20$ K, far above the magnetic ordering temperature. One may well ask how this combination of anisotropy, frustration, and magnetic order arises.

An insightful general argument about anisotropies in non-Kramers systems was given by Griffith in Refs. [57], and it concerns instances where an even number of low-lying singlets is well isolated from higher excited states. In the simplest case of two low-lying singlets it was shown that the axial anisotropy of the magnetic moment is determined by their time-reversal (TR) behavior, i.e. the local Ising axis of a non-Kramers ion is independent of the direction of the applied field (provided this is only of perturbative order [58]).

As already pointed out in Ref. [59], this is of interest for the quasidoublet states ($|p\rangle, |q\rangle$) of the CEF in similar garnets and, in order to understand and possibly predict which of the three twofold axes of the $D_2$ CEF symmetry is the anisotropy axis for the magnetic moment, we follow Ref. [57] and investigate the matrix elements $\langle q|J|p\rangle$ and their behavior under TR transformation.

The action of the time-reversal operator $\Theta$ on angular momentum states $|\psi\rangle = \sum_{J,M_J} C_{J,M_J} |J,M_J\rangle$ is given by $|\psi\rangle^K = \Theta|\psi\rangle = \sum_{J,M_J} C_{J,M_J}^* (-1)^{J-M_J} |J,-M_J\rangle$ [45]. Then the time reversal of the CEF states in Eq. (2) reads

$$
|\Gamma_1\rangle^K = |\Gamma_1\rangle,
|\Gamma_2\rangle^K = |\Gamma_2\rangle,
|\Gamma_3\rangle^K = -|\Gamma_3\rangle,
|\Gamma_4\rangle^K = -|\Gamma_4\rangle.
$$

Calculating the matrix elements $\langle \Gamma_{\alpha}\mid J \mid \Gamma_{\beta}\rangle$, by means of the general expansions in Eq. (2), reveals the correspondence

$$
\mathbf{m}(\Gamma_3, \Gamma_4) \parallel \mathbf{m}(\Gamma_2, \Gamma_1) \parallel \mathbf{x},
\mathbf{m}(\Gamma_3, \Gamma_4) \parallel \mathbf{m}(\Gamma_2, \Gamma_1) \parallel \mathbf{y},
\mathbf{m}(\Gamma_3, \Gamma_4) \parallel \mathbf{m}(\Gamma_1, \Gamma_4) \parallel \mathbf{z},
$$

between the symmetry of the states and the direction of $\mathbf{m}(\Gamma_{\alpha}, \Gamma_{\beta})$ along $\mathbf{x}$, if $|\Gamma_{\alpha}\rangle, |\Gamma_{\alpha \neq \beta}\rangle$ have different basis decomposition (one with $M_{J_{\parallel}}$-even and the other with $M_{J_{\parallel}}$-odd) and different TR behavior (one even the other odd), along $\mathbf{y}$, if they have different basis decomposition but the same TR transformation, and along $\mathbf{z}$, if they have the same basis decomposition but opposite TR transformation.

Diagonalization of the crystal-field Hamiltonian in Eq. (1) with the parameters listed in Table III leads to a spectrum where the lowest singlet $|p\rangle$ and the first excited one $|q\rangle$ are, respectively, of the $|\Gamma_3\rangle$ and $|\Gamma_1\rangle$ type in Eq. (2) (see also Table V), $\mathbf{m}(p, q) \parallel \mathbf{x}$, according to Eq. (4), and as observed in the magnetic structure. We find indeed that $\langle p|J_z|q\rangle = \langle p|\hat{J}_z|q\rangle = 0$, identically, and so the low-temperature induced magnetic moment will be proportional to $\mathbf{m}(p, q) = -g_{\mu_B} \langle p|\hat{J}_z|q\rangle \mathbf{x}$ (see Appendix A for the different $\mathbf{x}$ in TGG).

The magnetic structure of TGG that we have determined is identical to that found previously [14], and to that found in various other rare earth garnets, including Dy$_3$Al$_5$O$_{12}$ (DAG). In the case of DAG, Wolf and collaborators showed that this magnetic structure minimizes the dipolar interactions for magnetic moments with $|100|$ Ising anisotropy and overall constraint of cubic symmetry [7], and this was subsequently confirmed by the application of the Luttinger-Tisza method [8]. In analogy with these RE garnets, it is typical also in TGG to ascribe the ordered structure predominantly to dipolar interactions.

In the theoretical description of the ordering transition in TGG of Ref. [23], Hammann and Manneville also concluded that the dipolar interaction is of primary importance, but found that the observed ordering temperature was too high to be accounted for only by induced moment ordering mediated by dipolar interactions [23]. They therefore showed that a molecular field theory taking into account also the role of hyperfine coupling, together with the dipolar interactions, would provide a reasonable estimate of $T_N^{\text{theo}} \approx 0.345$ K with respect to $T_N^{\text{exp}} \approx 0.25$ K, and implying the possibility of cooperative electronic and nuclear spin order. This theory was constrained by various experimental parameters of the time, which can now be examined more accurately.

Our study confirms directly the value of the (ground state) quasidoublet gap, and it suggests a time-reversal analysis of the direction of the local easy axes (among the three equivalent ones permitted by the crystalline point symmetry). The

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TABLE IV. Estimation of the key parameters for the mean-field theory in Ref. [23]. The quantities are given for Tb$^{3+}$ ions, where the Landé factor is $g = 3/2$ and the hyperfine coupling constant is $a_{f} \approx 2.2 \times 10^{-3}$ meV [23] (see Table 5.5 in Abragam and Bleaney [45]).

| Ref. [23] | This work |
|-----------|-----------|
| $|w|/\mu_{B}$ | 6.7 | 7.9 |
| $|\alpha|$ | 0.08 | 0.11 |

The aforementioned “off-diagonal” matrix element $ \langle p|J_{x}|q\rangle \approx 5.3 $, is a crucial quantity which relates to the coefficient $w$ in Ref. [23] via $ |w| = |m(p, q)| $, and therefore controls the key parameters for the mean-field theory presented therein (see Table IV). It is worth emphasising that our estimation ($w \approx 7.9 \mu_{B}$) is given by the wave functions, $ |p\rangle, |q\rangle $, of the ground-state quasidoublet obtained via exact diagonalization of the Hamiltonian in Eq. (1), i.e., crystal-field analysis of the neutron scattering data. In contrast, the estimation by Hammann and Manneville ($w \approx 6.7 \mu_{B}$) relies on a self-consistent fit to magnetization curves measured at different temperatures on a single crystal sample with magnetic field applied along the $\langle 111 \rangle$ direction [60].

Our direct estimation of $w$ opens up a possible route to understand coherently the behavior of the magnetic moment across different temperature regimes. In a forthcoming theoretical work we will study how the local states ($|p\rangle, |q\rangle$) of the Tb$^{3+}$ ions are the starting point for the definition of the new magnetic basis ($|p\rangle, |q\rangle$) which carries information about the admixture induced by the presence of the (molecular) field on one hand and of the thermal population of states on the other. (A similar reasoning was considered in Refs. [23,59], but actual local symmetries of the RE$^{3+}$ ions were disregarded as an element of interest.) This route can resolve the puzzle reported in Ref. [14], where neutron powder diffraction showed that full saturation of the magnetic moment was still not achieved down to 0.2 $T/T_{N}$ (at such temperatures $|m|/w \approx 0.6$ where $w$ is treated as the saturated magnetic moment per ion). In addition, this is expected to be the natural starting point to investigate which mechanism among others—to mention a few, competing interactions, fragmentation, fluctuations (outside of the neutron spectroscopy window)—act as leading term in the reduction of the magnetic moment.

Following the perspective of Ref. [23], the key parameters in Table IV also comprehend implications about the role of hyperfine fields in the magnetic ordering of TGG. The coefficient $|\alpha| = 2 |w| |\alpha_{f}| / \Delta g_{J} \mu_{B}$ is the dimensionless strength of the hyperfine coupling in the pseudospin-1/2 effective Hamiltonian obtained by projecting on the quasidoublet subspace—here $\Delta$ is the CEF ground state splitting (see Sec. III B) and $\alpha_{f}$ comes from the hyperfine Hamiltonian $H_{HF} = a_{f} J \cdot I$, which, together with the Zeeman term $H_{B} = -g_{J} \mu_{B} J \cdot B$, act as a perturbation to $H_{CEF}$ in Eq. (1).

In our experiment we cannot directly probe the behavior of the nuclear spins, and we cannot verify definitively the relevance of our findings with respect to the hyperfine aspects. If we were to strictly apply the criteria of Hammann and Manneville, then we would deduce that the molecular field is mainly of electronic nature and that the cooperative electro-nuclear polarization of the magnetic moment of the Tb$^{3+}$ ions will be relevant only at the lowest temperatures (our results in Table IV imply $x \gtrsim 1.5$ in Fig. 3 of Ref. [23], which is to say that significant nuclear polarization is only expected at $T \sim 83$ mK or $T/T_{N} \sim 0.33$). However, one ought to conduct a dedicated study of instances of hyperfine driven quantum criticality at a level which is beyond the scope of this manuscript.

The broadening and structure of the first excitation appearing at the lowest temperatures (Fig. 10) suggests that the crystal field level is developing into dispersive excitons, as expected in a singlet-singlet system [40], and it would be interesting to examine the extent of any mode-softening at the transition [61].

The crystal field states are of importance in the acoustic Faraday effect (AFE) [27,28] and acoustic Cotton-Mouton effect [29], and in quantitative explanations of the elastic constants [29] and may also be of importance in the thermal Hall effect. A detailed explanation of the acoustic Faraday effect exists [27] and depends in part on the character of the crystal field states of the main terbium site and their coupling with phonons. It is noted in Ref. [27] that the crystal field states were not precisely known, and a simplified scheme of cubic Tb$^{3+}$ site symmetry was adopted using a $\Gamma_{1}$ doublet and $\Gamma_{4}$ triplet, with specific field dependent components and matrix elements amongst them. A quantitative explanation of the temperature and field dependence of the elastic constants [29,62] has been attempted using the parameters of Ref. [20], but, as described above, a more accurate description should be expected using the parameters presented here, particularly since the intersections and crossings of some levels are modified by the use of our parameters.

V. CONCLUSIONS

The low-temperature magnetic structure of TGG is of the multiaxial antiferromagnet type. As closer examination has shown, the large anisotropy of the Tb$^{3+}$ magnetic moments and its direction can be explained by the time-reversal properties of the crystal-field states of this non-Kramers ion. The singlet ground state of the Tb$^{3+}$ ion in TGG indicates the presence of perturbations allowing this induced-moment type of system to develop long-range order. Determination of the nature of the ordering transition still requires additional experimental investigation.

Two striking anomalies can be recognised in the crystal-field spectra measured by neutron scattering. These are: Substantial differences between observed and expected linewidths of low-lying excitations and the pronounced temperature-dependent structure of the quasidoublet ground state [see Fig. 12(b)]. To address such effects, elucidating superexchange mechanisms between Tb$^{3+}$ ions, as well as coupling between magnetic and lattice degrees of freedom, should be a priority for further studies. These investigations indeed introduce a collective character by perturbing the microscopic states from those derived within a (semiclassical) single-ion approach.

In addition, the results of bulk measurements we have shown departures from a picture of noninteracting spins when cooling below 50 K. This is consistent with a suspected onset of the effects of interactions that may lead to a spin liquid
phase of so far unknown character. The presence of a disordered but correlated regime in TGG is not surprising given its moderate frustration \((f \sim 32)\). Nonetheless its appearance would be expected more in the range of temperatures \(|\theta_{\text{CW}}| > T > T_N\).

Further investigations with single crystal samples that could provide the detailed \(|Q|-dependence of the observed features are needed to determine precisely the perturbation affecting single-ion anisotropy and the exact character of the intermediate short-range order below 50 K in TGG. We believe that these would furthermore allow us to establish a consistent microscopic theory for the thermal Hall behavior. By enriching the picture of the low temperature physics in TGG, the results we present here constitute the basis for a full microscopic description of the observed effect.

ACKNOWLEDGMENTS

The authors would like to express their gratitude to Victoria García Sakai for her assistance during the IRIS experiment, and to O. Cépas, M. Gingras, M. Kenzelmann, M. Mori, and M. Zhitomirsky for stimulating discussions and insightful suggestions. Neutron scattering experiments were carried out at the spallation neutron source ISIS at the Rutherford Appleton Laboratory, UK. The presented graphical representations of the crystallographic structure were prepared with VESTA software [19].

APPENDIX A: SYSTEM OF COORDINATES

In the rare earth garnets, each magnetic ion sits on one of the vertices joining the triangles of the two hyperkagome sublattices [63], as represented in Fig. 1. From a local point of view, the orthorhombic point symmetry characterizes the symmetry of the coordination environment or crystal field of a given \(R^{3+}\) ion (not shown in Fig. 1—details in Sec. III B). From a global point of view (i.e., with respect to the crystal axes), the orientation of this local coordination environment varies from one site to the other. Throughout the whole garnet lattice, according to the space group \(I\bar{a}3d\), there are six orientations of the local axes with respect to the global cubic axes. In this work the local coordinates \(x_i, y_i, z_i\) are chosen to be parallel to the 2-fold rotation axes of the \(D_2\) point group, so that with respect to the global \(X, Y, Z\) coordinates, they are:

\[
x_1 = [0, 0, 1], \quad y_1 = \frac{1}{\sqrt{2}}[1, \bar{1}, 0], \quad z_1 = \frac{1}{\sqrt{2}}[1, 1, 0]; \quad (A1a)
\]

\[
x_2 = [0, 1, 0], \quad y_2 = \frac{1}{\sqrt{2}}[\bar{1}, 0, 1], \quad z_2 = \frac{1}{\sqrt{2}}[1, 0, 1]; \quad (A1b)
\]

\[
x_3 = [1, 0, 0], \quad y_3 = \frac{1}{\sqrt{2}}[0, 1, \bar{1}], \quad z_3 = \frac{1}{\sqrt{2}}[0, 1, 1]; \quad (A1c)
\]

\[
x_4 = [0, 0, \bar{1}], \quad y_4 = \frac{1}{\sqrt{2}}[\bar{1}, \bar{1}, 0], \quad z_4 = \frac{1}{\sqrt{2}}[\bar{1}, 1, 0]; \quad (A1d)
\]

\[
x_5 = [0, \bar{1}, 0], \quad y_5 = \frac{1}{\sqrt{2}}[\bar{1}, 0, \bar{1}], \quad z_5 = \frac{1}{\sqrt{2}}[1, 0, \bar{1}]; \quad (A1e)
\]

\[
x_6 = [\bar{1}, 0, 0], \quad y_6 = \frac{1}{\sqrt{2}}[0, \bar{1}, \bar{1}], \quad z_6 = \frac{1}{\sqrt{2}}[0, \bar{1}, 1]. \quad (A1f)
\]

APPENDIX B: EIGENVECTOR STRUCTURE OF CRYSTAL-FIELD STATES

See Table V.

| \(E\) [meV] | \(|\psi_{\text{G}}\rangle = \sum c_{i,MJ} |J, M_J\rangle\), expressed as expansions of the \(|M_J\rangle \equiv |J, M_J\rangle\) eigenstates of \(J_z\), in the Russell-Saunders coupling scheme (\(J = 6, L = 3, S = 3\)) for the \(^7\text{Fe}\) ground multiplet of \(\text{Tb}^{3+}\) ion. A blank entry means a zero coefficient. |
|---|---|---|---|---|---|---|---|---|---|---|---|
| \(\psi_0\) | \(\Gamma_1\) | 0.0 | -0.091 | -0.026 | 0.701 | 0.701 | -0.026 | -0.091 |
| \(\psi_1\) | \(\Gamma_1\) | 0.22 | 0.182 | 0.122 | -0.325 | -0.832 | -0.325 | 0.122 | 0.182 |
| \(\psi_2\) | \(\Gamma_2\) | 4.50 | -0.594 | 0.096 | 0.371 | -0.371 | -0.096 | 0.594 |
| \(\psi_3\) | \(\Gamma_1\) | 5.25 | 0.683 | 0.061 | -0.085 | -0.215 | -0.085 | 0.061 | -0.683 |
| \(\psi_4\) | \(\Gamma_1\) | 6.01 | 0.383 | 0.115 | 0.583 | -0.583 | -0.115 | -0.383 |
| \(\psi_5\) | \(\Gamma_2\) | 6.54 | -0.005 | 0.502 | 0.498 | -0.498 | -0.502 | 0.005 |
| \(\psi_6\) | \(\Gamma_3\) | 26.40 | 0.032 | 0.706 | 0.030 | 0.030 | 0.706 | 0.032 |
| \(\psi_7\) | \(\Gamma_1\) | 28.27 | 0.016 | 0.415 | 0.535 | -0.290 | 0.535 | 0.415 | 0.016 |
| \(\psi_8\) | \(\Gamma_2\) | 34.26 | 0.566 | 0.302 | -0.298 | 0.298 | -0.302 | -0.566 |
| \(\psi_9\) | \(\Gamma_3\) | 34.73 | -0.019 | -0.691 | 0.148 | -0.148 | 0.691 | 0.019 |
| \(\psi_{10}\) | \(\Gamma_4\) | 35.63 | -0.701 | 0.035 | -0.090 | -0.090 | 0.035 | -0.701 |
| \(\psi_{11}\) | \(\Gamma_2\) | 36.38 | 0.424 | -0.397 | 0.403 | -0.403 | 0.397 | -0.424 |
| \(\psi_{12}\) | \(\Gamma_1\) | 38.20 | 0.023 | 0.556 | -0.318 | 0.422 | -0.318 | 0.556 | 0.023 |
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