Magnetic excitations in the XY-pyrochlore antiferromagnet Er$_2$Ti$_2$O$_7$

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The XY-pyrochlore antiferromagnet Er$_2$Ti$_2$O$_7$ is studied by heat capacity measurements and electron spin resonance spectroscopy performed on single crystal samples. The magnetic phase diagrams are established for two directions of applied field, $H \parallel [100]$ and $H \parallel [111]$. In the magnetically ordered phase observed below $T_N = 1.2$ K, the magnetic excitation spectrum consists of a Goldstone mode acquiring an isotropic gap in an applied field, and another mode with a gap softening in the vicinity of a field-induced phase transition. This second-order transition takes place at a critical field $H_c$ above which the magnetization process is accompanied by a canting of the magnetic moments off their local “easy-planes”. The specific heat curves for $H \parallel [100]$ ($H \gg H_c$) are well described by a model presuming a single dispersionless excitation mode with the energy gap obtained from the spectroscopic measurements.

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I. INTRODUCTION

Compounds with strong geometrical frustration of magnetic bonds have attracted much attention due to the unusual properties of their manifold ground states and their peculiar spin dynamics. For some systems the degeneracy of the ground state is infinite (in the nearest-neighbor exchange approximation), which leads to (i) delayed magnetic ordering and a wide temperature interval with a short-range correlated state, the so called cooperative paramagnet and (ii) an enhanced role for the weaker interactions in the eventual formation of a long-range order state at low temperature. Different mechanisms for lifting the macroscopic degeneracy and selecting a specific ground state is a matter of individual consideration for each system. In the case of pyrochlore magnets, (nearly) fully ordered states, cooperative paramagnetic states, spin-glass and spin-ice states have all been observed in different materials.

Experimental investigations of excitation spectra often provide the only viable method of determining a specific set of interactions to be taken into account for an individual magnetic compound. For example, in a Heisenberg pyrochlore magnet Gd$_2$Sn$_2$O$_7$, where the four-sublattice structure with $k = 0$ (a so-called plane cross or a Palmer-Chalker state) is found at low temperature, the spin-wave calculations performed for this type of ordering with known microscopic parameters of exchange, dipolar interactions and a single-ion anisotropy are in perfect agreement with the experimentally obtained picture, thus demonstrating the validity of this description.

Unlike the Gd-based Heisenberg pyrochlores, erbium titanate Er$_2$Ti$_2$O$_7$ has a strong local XY-type anisotropy of magnetic moments due to the large orbital moment of Er$^{3+}$ ions ($L = 6$) which is not quenched by the crystal field. The Curie-Weiss temperature in this system obtained from the susceptibility data points to an antiferromagnetic nearest-neighbor exchange interaction. Depending on the temperature interval of data fitting, the value of $\Theta_{CW}$ varies from -13 K in the higher temperature limit 80-300 K to -22 K if determined from data collected at temperatures below 50 K. Er$_2$Ti$_2$O$_7$ is known to undergo a second order transition to a long-range ordered state at about 1.2 K. A spherical neutron polarimetry experiment indicates that the magnetic structure is formed from six domains, all of which are equally occupied. In addition, recent high resolution neutron diffraction measurements detect the coexistence of short and long-range order. Low-lying excitation modes ungapped in zero field and near a critical field of about 15 kOe are also observed in this experiment. Muon spin-relaxation measurements show non-vanishing spin dynamics as the temperature approaches zero in contrast to the expectations for a conventional magnet.

The non-coplanar structure appearing below $T_N$ is presumed to be induced by fluctuations but this model predicts a strong first-order phase transition, while experimentally it is found to be continuous. A mean field model which includes single-ion anisotropy and anisotropic exchange interactions is capable of producing the required long-range order however thermal or quantum fluctuations are expected to be important in this material. Polarized neutron diffraction measurements suggest that in addition to crystal field parameters, an anisotropic molecular field tensor must be included in order to reproduce the experimental values of the local susceptibility tensor.

We present the results of an extensive electron spin resonance study of Er$_2$Ti$_2$O$_7$, which is used as a high-resolution probe of the $q = 0$ energy structure for different directions of an applied field. A Goldstone mode is observed and its linear increase with an applied magnetic field is traced. Additionally, we have detected a gapped mode softening in the vicinity of a critical field. The
origin of both types of oscillation is discussed. Specific heat measurements were also carried out for $H \parallel [100]$ and $H \parallel [111]$ to complement the previously reported results\textsuperscript{12} for $H \parallel [110]$. The calorimetry results are compared with the ESR and inelastic neutron scattering data\textsuperscript{12} and an overall agreement between the different experimental methods is demonstrated.

II. EXPERIMENTAL PROCEDURES

A single crystal sample of Er$_2$Ti$_2$O$_7$ was grown by the floating zone technique\textsuperscript{12}. Small thin plates of a characteristic size $1 \times 1 \times 0.2$ mm$^3$ (about 1 mg in mass) containing the (110)-plane were cut out of the original larger sample. The plates were mounted in such a way that the magnetic field was applied in the sample plane in order to minimize the demagnetization effect. Specific heat was measured using a Quantum Design Physical Property Measurement System calorimeter equipped with a $^3$He option and a 90 kOe cryomagnet.

ESR measurements were carried out using a homemade transmission type spectrometer built on a $^3$He cryostat with a base temperature of 0.45 K. The lowest eigen-frequency of the resonant cavity was 25 GHz. The highest frequency was limited to about 70 GHz by the so-called “size effect” i.e. the condition that the half-wavelength of the microwave radiation inside the sample is of the same order as the sample size. For the pyrochlore magnets, this limit is especially rigid because of their large magnetic permeability. It has been shown in case of another pyrochlore, Gd$_2$Ti$_2$O$_7$, that above this frequency limit a parasitic absorption (not directly related to magnetic excitation spectrum of the samples) can disguise its resonance pattern\textsuperscript{12}.

III. EXPERIMENTAL RESULTS

The temperature dependence of the specific heat in zero external magnetic field is shown in Fig. 1. It demonstrates a sharp anomaly at $T_N = 1.2$ K in close agreement with the previously obtained results. This peak corresponding to a second-order phase transition into a magnetically ordered state is followed by $C_p \propto T^3$ drop of specific heat on decreasing the temperature. The magnetic entropy of the system obtained by integrating the $C_p/T$ curve with the phonon contribution subtracted (see below for details) saturates at the value of $R \ln 2$. This points to the effective $S = 1/2$ pseudo-spin character of the magnetic Er$^{3+}$ ions originating from the double degeneracy of its ground state in a crystal field.

In an applied magnetic field the transition shifts to lower temperatures and becomes difficult to detect in the $C(T)$ curves. The transition is also traced by recording the field dependence of the specific heat at constant temperature. The $C(H)$ dependence measured for two different directions of the external magnetic field, $H \parallel [100]$ and $H \parallel [111]$, shows that the peak anomalies are easily detectable (see the inset to Fig. 1) and that their behaviors are quite anisotropic in nature with the transitions occurring at $H_{c1}^{[100]} = 16.5 \pm 0.5$ kOe and $H_{c1}^{[111]} = 13.5 \pm 0.5$ kOe.

The observed transformation of the $C(T)$ curves on increasing the external magnetic field up to and above the $H_c$ for these two field directions is similar to the behavior reported in Ref. 12 for $H \parallel [110]$. In higher applied fields, the sharp low-temperature anomaly seen in the $C(T)$ curves is replaced by a much broader feature developing in the higher-temperature part of the curves (see Fig. 2).

Following the results of an inelastic neutron scattering experiment for $H \parallel [110]$ in Ref. 12 which showed for the lower excitation mode the dispersion vanishing at $H \gg H_c$, one can attempt to approximate Er$_2$Ti$_2$O$_7$ in high fields as an ensemble of non-interacting two-levels systems. In this case, the molar specific heat can be expressed in the form:

$$C_p = \alpha R \left(\frac{\Delta}{kT}\right)^2 \frac{e^{-\Delta/kT}}{1 + e^{-\Delta/kT}} + \beta T^3,$$

(1)

where the first term represents the Schottky anomaly ($\Delta$ is the gap value, $\alpha \approx 1$ is a numerical coefficient), and the second term is the phonon contribution. Taking $\alpha$, $\beta$ and $\Delta$ as fitting parameters, one can perfectly well reproduce the specific heat curves in higher fields (50, 70 and 90 kOe) in Fig. 2 for $H \parallel [100]$ over the entire temperature range shown. For all scans we obtain $\alpha = 0.84 \pm 0.01$ and $\beta \approx (1.33 \pm 0.03) \times 10^{-3}$ J/(K$^4$ mole Er). The latter value agrees well with the specific heat of an isomorphous nonmagnetic compound Y$_2$Ti$_2$O$_7$ (see e.g. Ref. 19) rescaled to Er$_2$Ti$_2$O$_7$ by molar mass in the approximation that the Debye temperature $\theta \propto 1/\sqrt{M}$. 

![Graph](image)
The field dependence of the gap values is presented in the resulting phase diagram (Fig. 3) and discussed below. Only the high temperature part of the heat capacity measured at \( H = 30 \, \text{kOe} \) can be satisfactorily described by equation (1), as the low energy contributions to the entropy are not taken into account in this simple model. The observed behavior of the specific heat indicates that the high field excitation spectrum may indeed be modeled as a set of two-level systems with a field-dependent gap, while nearer to the critical field, a more important role is played by the dispersion of the excitation mode.

The specific heat reported by Ruff et al.\textsuperscript{12} for a magnetic field along the [110] direction exhibits the same properties. The high field \( C_p(T) \) curves shown in Fig. 1 of Ref. 12 can also be described in the manner outlined above. The corresponding gap value at \( H = 70 \, \text{kOe} \) is shown on Fig. 6 for reference.

Using the same model, however, we could not obtain satisfactory fits to our data obtained for \( H \parallel [111] \). The solid lines on the lower panel of Fig. 2 drawn with the same set of parameters as for \( H \parallel [100] \) illustrate the point that some of the magnetic entropy for this field direction is redistributed to lower temperatures. One could suggest that the energy of the long-wave excitations grows more slowly for a magnetic field applied along the [111] axis than for other field directions. The relationship between the heat capacity data and the results of the magnetic resonance measurements is discussed below.

The resonance absorption spectra recorded at the lowest experimentally available temperature of 0.45 K at constant frequency on increasing/decreasing field sweeps are presented in Fig. 3. Rather similar results have been obtained at \( H < H_c \) for fields along the [100], [110] and [111] axes. Two spectral lines of Lorentzian shape are observed below \( T_N \). The resonance field of line 1 grows linearly with the increasing frequency while the position of line 2 shifts to lower fields. Their relative intensities vary with field direction but the halfwidth of both lines lie in the interval \( \Delta H_{1,2} \approx 1 - 1.2 \, \text{kOe} \). The nearly frequency independent resonance field of the second line in the vicinity of the critical field points to the possible complete softening of this resonance mode, although our experimental frequency limit does not allow for spectral records below 25 GHz (corresponding to energies of about 0.1 meV). At higher fields, for \( H > H_c \), a third narrow line \( (\Delta H_3 \approx 0.2 \, \text{kOe}) \) appears in the spectra for \( H \parallel [100] \) and [110]-axes, while for \( H \parallel [111] \)-axis an absorption of this type is not seen. Instead, two much broader spectral lines with considerably smaller intensities (lines 3a, b on the lowest panel of Fig. 3) are observed. A quantitative analysis of the frequency-field diagram is performed below.

A typical transformation of the resonance spectrum on increasing the sample temperature above the Neel point is shown in Fig. 4. The records at \( \nu = 40.2 \, \text{GHz} \) reveal an intensity loss for all three spectral lines on heating. Lines 1 and 2 almost disappear at the transition temperature (about 1.0 K and 0.8 K for the corresponding resonance fields). Although line 3 exists in the higher fields and is therefore not directly related to any low-field antiferromagnetic order, it is observed only at low temperature, broadening and disappearing on warming above 1.5 K. As seen in Fig. 4 the absorption in the sample is accompanied by a broad non-resonant background gradually vanishing in the high field limit. This background signal grows in amplitude on approaching the Neel temperature and finally replaces the three lines of the low-temperature spectrum. Comparing the spectra recorded at various frequencies one finds that the background is excited only by the microwave field component directed along the external magnetic field. Therefore, the absorption is associated with the longitudinal oscillation of the magnetic moment and should be damped in the vicinity of the field at which the magnetization reaches its maximum value. A similar effect was observed in the Heisenberg pyrochlore system \( \text{Gd}_2\text{Ti}_2\text{O}_7 \) where the longitudinal susceptibility was supposed to result from a partial disorder of the magnetic ground state.\textsuperscript{5} As in the present case, this mode had very strong relaxation (of unknown origin) which makes it impossible to observe other than
in the form of a broad non-resonant background vanishing above the saturation field. Approximating the system as a set of "rigid" sublattices canting to the magnetic field, one obtains the slow response of the system to the microwave field to be $\chi \sim \cos^2 \phi$, where $\phi$ is an angle between the sublattice and the field. Since the total magnetization of the system is $M = M_c \sin \phi$ ($M_c$ is a maximum magnetization), the resulting contribution of such a degree of freedom to the dynamic susceptibility of the sample can be estimated as $\chi'' \propto 1 - (M/M_c)^2 = 1 - (H/H_c)^2$ ($H_c$ is a critical field). The above approach is qualitatively illustrated in the inset to Fig. 3. The imaginary part of the dynamic susceptibility extracted from the absorption spectrum record was fitted by a sum of three resonance lines of Lorentzian shape and a non-resonant background with $H_c$ taken as a fitting parameter. The fit shown by the solid line was achieved for $H_c = 23.6$ kOe which compares reasonably well to the value of the critical field, given the simplicity of the above approximation. One should also mention, that although this transition is associated with slowing down the magnetization growth, it is not a usual saturation field, as will be discussed below.

**IV. DISCUSSION**

For comparison with the experimentally observed spectra one should first determine the $g$-values of a single Er$^{3+}$ ion in a crystal field (CF) for the various directions of an external magnetic field. The total momentum in the ground state of Er$^{3+}$ ion (electron shell configuration 4f$^1$6s$^2$) is $J = 15/2$ ($L = 6$, $S = 3/2$). The CF of a trigonal $D_{3d}$ point symmetry at the erbium sites (the three-fold axes are parallel to local [111] axes) splits the 16-fold state into 8 doublets. A calculation of this splitting using the CF Hamiltonian parameters rescaled from another pyrochlore compound Ho$_2$Ti$_2$O$_7$ (given in Ref. [21]) describes well the observed gaps between the lowest and two excited CF levels and gives the following expression for the lowest Kramers doublet: $\psi_{1,2} = 0.543|\pm\frac{11}{2}\rangle \pm 0.238 |\pm\frac{5}{2}\rangle - 0.563 |\pm\frac{1}{2}\rangle \mp 0.388 |\pm\frac{3}{2}\rangle + 0.426 |\pm\frac{13}{2}\rangle$. Using these wave-functions, one can easily calculate the single ion energy splitting under magnetic field applied at an arbitrary angle $\alpha$ with respect to a local trigonal axis:

$$\Delta \varepsilon = \mu_B H \sqrt{g_\|^2 \cos^2 \alpha + g_\perp^2 \sin^2 \alpha},$$

where $g_\| = 0.24$ and $g_\perp = 7.6$ are the $g$-factors parallel and perpendicular to [111]. A more general calculation which involves the solution of a total atomic Hamiltonian and uses high temperature susceptibility data in the fitting procedure gives slightly different lowest doublet wave-functions but the same $g$-values. It should be noted...
that recent neutron scattering measurements\textsuperscript{15} have determined a much less pronounced XY-anisotropy of the local susceptibility tensor with the values of $g_{\parallel} = 2.6$ and $g_{\perp} = 6.8$. When the field is applied along [100]-axis all four magnetic moments in the unit cell are in equivalent positions with $\cos \alpha = 1/\sqrt{3}$. Field directions $H \parallel [110]$ and [111] create two nonequivalent positions with $\cos \alpha_1 = \sqrt{2}/3$, $\cos \alpha_2 = 0$ and $\cos \alpha_1 = 1$, $\cos \alpha_2 = 2\sqrt{2}/3$ respectively. The corresponding effective $g$-values are collected in a Table I.

Let us now analyze the resonance spectra measured at the temperature 0.45 K for the three different orientations of the external magnetic field. The frequency-field diagram is presented in Fig. 5. The resonance spectrum observed below the critical field $H_c$ consists of two branches. Branch 1 is increasing in field and can be approximated by a linear dependence $\nu = \gamma H$ with the value of $\gamma$ slightly dependent on the field direction: $\gamma_{[100]} = 3.8 \pm 0.1$ GHz/kOe, $\gamma_{[110]} = 3.6 \pm 0.1$ GHz/kOe, $\gamma_{[111]} = 4.1 \pm 0.1$ GHz/kOe, which correspond to effective $g$-values 2.7, 2.6 and 2.9 respectively. This linear in field gap signifying the existence of a Goldstone mode at zero magnetic field is quite unexpected for a strongly anisotropic magnetic system where all the acoustic modes should be influenced by spin-orbit coupling. The residual gap due to a possible (within the experimental accuracy) deviation from the linear extrapolation to zero field does not exceed a value $\sim 10$ GHz $\approx 0.04$ meV. One should also note that the power law dependence of the specific heat $C_p \propto T^3$ extending to very low temperatures\textsuperscript{8,9} confirms the existence of gapless magnetic excitations at zero field. Obviously, this mode cannot be interpreted in terms of a single ion splitting. The observed increase of the gap, which is linear in field, should be ascribed to a uniform oscillation of the magnetization resulting from the in-phase spin motion in the planes perpendicular to the local [111] axes.

Branch 2 has a gap that diminishes quickly in the vicinity of the critical field. Although the high frequency measurements were hindered by the “size effect” and the zero field gap of branch 2 is not determined from our experiment, one can relate this branch to the gapped mode with $\Delta_0 \approx 0.4$ meV observed by inelastic neutron scattering\textsuperscript{12} The field dependence of this branch in the vicinity of the second order transition $H = H_c$ is similar to the behavior of the optical branch near the spin-flip transition: $\Delta \sim \Delta_0 \sqrt{1-(H/H_c)^2}$. Using this formula one can satisfactorily approximate the field dependence of branch 2 for all three field orientations and therefore estimate the critical field values: $H_c^{[100]} = 16.5 \pm 0.5$ kOe; $H_c^{[110]} = 16.0 \pm 0.5$ kOe and $H_c^{[111]} = 15.0 \pm 0.5$ kOe. An optical branch with an analogous field dependence was observed in the spectrum of the Heisenberg pyrochlores Gd$_2$Ti$_2$O$_7$ and Gd$_2$Sn$_2$O$_7$. It corresponds to the out-of-phase oscillations of spins in local “easy” planes.

The analysis of the relative intensities of the magnetic Bragg peaks\textsuperscript{10} shows that the field-induced transformation of the magnetic structure can roughly be described as a rotation of magnetic moments in their local “easy-planes”. The critical field is determined by the maximum value of the magnetic moment which can be achieved without canting from these planes. This geometric restriction was analyzed theoretically and the corresponding critical fields were calculated\textsuperscript{11,12}. The ratio \(H_c^{[100]} / H_c^{[110]} = 2\sqrt{2}/(\sqrt{3} + 1) \approx 1.04\) is in agreement with our results within the experimental accuracy while the theoretical value for \(H_c^{[100]} / H_c^{[111]} = 7\sqrt{2}/(3\sqrt{3} + \sqrt{6}) \approx 1.3\) appears to be considerably larger than the value 1.1 obtained from our ESR experiments. In contrast, the corresponding ratio of critical fields measured by specific heat, 16.5 : 13.5 \approx 1.2 demonstrates better agreement with the model. The resulting phase diagram obtained on the basis of both measurements at $H \parallel [100]$ and [111] are shown in the upper panel of Fig. 4.

At $H > H_c$, for two directions of the magnetic field $H \parallel [100]$ and [110], there exists a single gapped re-
FIG. 6: Upper panel: The phase diagram of Er$_2$Ti$_2$O$_7$ for $H \parallel [100]$ (□, ■) and $H \parallel [111]$ (Ο, ⋄) obtained from specific heat and magnetic resonance data respectively. Lower panel: gap values determined from fitting the $C_p(T)$ curves for $H \parallel [100]$ by equation (1); $\nu(H)$ dependence of ESR branch 3 is shown by crosses, solid line is a linear extrapolation of this branch to high fields. The schematic transformation of the excitation spectrum at high fields (based on experimental data from Ref. [12]) is given in the inset.

One can also establish a relationship between the increase of the gap as observed directly by the ESR spectroscopy and the evolution of the specific heat curves under field. The gap values obtained from the specific heat in the approximation of a single dispersionless gapped mode (two-level Schottky anomalies) and the field dependence of branch 3 at $H \parallel [100]$ are presented in the lower panel of Fig. 6. The difference between them at fields just above the transition results from the effect of dispersion on the specific heat which is not taken into account in our fitting. The convergence of the Schottky anomaly gaps to the high field linear extrapolation of branch 3 is evidence of a decrease in the dispersion of this mode in the high field limit. A similar effect, shown schematically in the inset of Fig. 6, was directly observed in inelastic neutron scattering experiments for $H \parallel [110]$. The agreement between ESR and specific heat data (single ESR branch and satisfactory modeling by a two-level system) is also established for $H \parallel [110]$. In contrast, the failure of such a model for $H \parallel [111]$ (see Fig. 2) can be considered as a bulk analog of the absence of a well defined excitation branch in the ESR measurements.

V. SUMMARY

In conclusion, the specific heat and the low temperature magnetic resonance spectra were studied in the XY pyrochlore antiferromagnet Er$_2$Ti$_2$O$_7$. A Goldstone mode with a gap linearly dependent on the applied field is observed in the ordered state. Another mode has a zero field gap consistent with the inelastic neutron scattering results which softens in the vicinity of the second order transition driven by magnetic field. Presumably, these collective modes correspond to in-phase and out-of-phase oscillations of spins in local “easy” planes. The critical field values are determined in three principal field orientations both from ESR and specific heat measurements. This transition is associated with the maximum possible spin reorientation without canting from their planes. In the high field phase the excitation spectrum for two field directions $H \parallel [100]$ and $H \parallel [110]$ consists of a single branch. The linear increase of the gap of this branch in magnetic field is somewhat related to the $g$-factor values characteristic of a splitting of a single ion lowest doublet. The specific heat curves observed at high fields demonstrate a Schottky-type anomaly with a gap that also increases with field. No well defined excitation mode was observed above $H_c$ at $H \parallel [111]$. Accordingly, the specific heat in strong magnetic fields does not demonstrate a two-level splitting with a single gap. A large difference in $g$-factors for the two nonequivalent positions of the magnetic ions with respect to the field is suggested to account for the absence of a collective excitation mode. Our measurements provide a framework with which to calculate the excitation spectrum in Er$_2$Ti$_2$O$_7$ using an effective pseudo-spin-1/2 model.
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