Order by Disorder and Energetic Selection of the Ground State in the XY Pyrochlore Antiferromagnet \( \text{Er}_2\text{Ti}_2\text{O}_7 \). An Inelastic Neutron Scattering Study

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(Dated: October 24, 2014)

Examples of materials where an “order by disorder” mechanism is at play to select a particular ground state are scarce. It has recently been proposed, however, that the antiferromagnetic XY pyrochlore \( \text{Er}_2\text{Ti}_2\text{O}_7 \) reveals a most convincing case of this mechanism. Observation of a spin gap at zone centers has recently been interpreted as a corroboration of this physics. In this paper, we argue, however, that the anisotropy generated by the interaction-induced admixing between the crystal-field ground and excited levels provides for an alternative mechanism. It especially predicts the opening of a spin gap of about 15 \( \mu \text{eV} \), which is of the same order of magnitude as the one observed experimentally. We report new high resolution inelastic neutron scattering data which can be well understood within this scenario.

Geometrically frustrated magnetism is a forefront research topic within condensed matter physics, as testified by the wealth of exotic phenomena discovered over the past years\textsuperscript{2–6,9–13}. For instance, the problem of an XY antiferromagnet on the pyrochlore lattice (the celebrated lattice of corner sharing tetrahedra) has been considered with much interest since this model displays an extensive classical degeneracy\textsuperscript{3,14}, along with classical and quantum order by disorder (ObD) effects\textsuperscript{2,6,9,11,13}. The elegant concept of ObD\textsuperscript{11,115} is a cornerstone of ordering in frustrated condensed matter systems. ObD comes into play by selecting a ground state, either because fluctuations away from this particular configuration allow for a relative gain of entropy compared to other classically degenerate states, or because quantum mechanical zero point fluctuations define a minimum in the total energy.

Until now, the number of confirmed examples for ObD in real materials have remained scarce\textsuperscript{14,15}. For ObD to be an efficient selection mechanism, the classical ground state degeneracy must be extremely robust and the minimal theoretical model not openly subject to additional terms that would spoil the accidental emerging symmetry and lift the degeneracy. Recently, the XY pyrochlore antiferromagnet \( \text{Er}_2\text{Ti}_2\text{O}_7 \) has been proposed as a candidate that satisfies these conditions in a rather compelling way\textsuperscript{3,6,110}. Given the unique position of \( \text{Er}_2\text{Ti}_2\text{O}_7 \) among frustrated quantum magnets, it is of foremost importance to scrutinize the soundness of this proposal.

The crystal electric field (CEF) acting on the Kramers \( \text{Er}^{3+} \) ion is responsible for a strong XY-like anisotropy, with easy magnetic planes perpendicular to the local \( (111) \) ternary axes\textsuperscript{3,15}. Combined with antiferromagnetic interactions, an extensive classical degeneracy is expected\textsuperscript{6,113}. Despite this degeneracy, \( \text{Er}_2\text{Ti}_2\text{O}_7 \) undergoes a second order phase transition towards an antiferromagnetic non-collinear \( \mathbf{k} = 0 \) Néel phase at \( T_N = 1.2 \text{ K} \)\textsuperscript{113}. In this configuration, denoted \( \psi_2 \) and depicted in Fig. 1a, the magnetic moments are perpendicular to the \( (111) \) axes\textsuperscript{3,20} and make a zero net magnetic moment per tetrahedron.

A theory based on a Hamiltonian written in terms of interacting pseudospins \( 1/2 \), each describing the single-ion CEF ground doublet, along with four anisotropic nearest-neighbor exchange parameters \( J_{zz}, J_{z\perp}, J_{\perp\perp}, J_{\perp z} \), has been proposed for \( \text{Er}_2\text{Ti}_2\text{O}_7 \)\textsuperscript{3}. For the set of parameters determined by inelastic neutron scattering (INS) experiments in a large applied magnetic field\textsuperscript{21}, the theory\textsuperscript{6} predicts a quantum ObD selection of \( \psi_2 \), on the basis of a linear spin wave calculation\textsuperscript{21,31,32}, as well as thermal ObD at \( T_c \), also selecting \( \psi_2 \)\textsuperscript{21}. Another consequence of ObD in \( \text{Er}_2\text{Ti}_2\text{O}_7 \) is the opening of a spin gap, previously inferred from EPR experiments\textsuperscript{22}, as well as from deviation of the \( T^3 \) law in specific heat measurements\textsuperscript{23}, and very recently confirmed from INS measurements\textsuperscript{21}. However, while the spin gap is a necessary consequence, it is not a definitive proof of this scenario: whatever the mechanism, a spin gap is expected since the ordered \( \psi_2 \) ground state breaks a global discrete symmetry\textsuperscript{3}.

In this work, we follow a different route and consider an anisotropic bilinear exchange Hamiltonian written for the \( \text{Er}^{3+} \) moments along with the CEF contribution (henceforth referred to as Model A). As shown in Ref. \textsuperscript{14}, an energetic selection of the \( \psi_2 \) state is possible for a specific range of anisotropic exchange parameters, owing to magneto-crystalline effects described by the CEF (see also Ref. \textsuperscript{20}). We model the spin excitations...
CEF energetic selection mechanism (Model A) – This approach considers a mean-field anisotropic bilinear exchange Hamiltonian written for the $\mathrm{Er}^{3+}$ moments $\vec{J}_i$ at sites $i$ of the pyrochlore lattice (see Refs. [1] and [26]). It also contains explicitly the CEF contribution, $\mathcal{H}_\text{CEF}$:

$$\mathcal{H} = \mathcal{H}_\text{CEF} + \frac{1}{2} \sum_{i,j} \vec{J}_i \cdot \vec{K}_{i,j} \cdot \vec{J}_j$$

(1)

where $\mathcal{H}_\text{CEF} = \sum_{n,m} B_{nm} O_{nm}$ is written in terms of $O_{nm}$ Stevens operators$^{[12,13]}$. The $B_{nm}$ have been determined to fit a number of experiments, including the intensities and positions of the crystal field levels, as well as the susceptibility$^{[20,30,31]}$. In the following, those values are considered as fixed parameters. $\vec{K}_{i,j}$ denotes an anisotropic coupling tensor, defined in the $(\vec{a}, \vec{b}, \vec{c})$ frame attached to the $\mathrm{Er}^{3+} - \mathrm{Er}^{3+}$ bond.$^{[31]}$ It is described by 3 symmetric parameters, $K_{a,b,c}$, and an anti-symmetric exchange constant (Dzyaloshinskii-Moriya like), $\vec{K}$.$^{[31]}$

In this model, the molecular field induces an admixture between the ground and excited CEF levels, leading to an effective magnetic anisotropy. This point is best evinced by considering the problem of an $\mathrm{Er}^{3+}$ ion in a local magnetic field $\vec{h}$: $\mathcal{H} = \mathcal{H}_\text{CEF} + \mu_\text{B} \vec{h} \cdot \vec{J}_i$. Figure 1 (b) shows the ground state energy of $\mathcal{H}$ computed as a function of $\theta$ and $\phi$ (in the local basis) for a field $h = 1$ T, which is the actual order of magnitude of the molecular field in $\mathrm{Er}_2\mathrm{Ti}_2\mathrm{O}_7$ (see below). Minima along the 6-fold directions of the CEF, slightly tilted away from the XY plane perpendicular to the local [111] direction, are clearly observed. As shown in Fig. 1 (c), the tilt grows as $h^2$ but remains less that one degree for realistic values of $h$. The potential well in the vicinity of the minima can be approximated by a highly anisotropic harmonic potential whose curvature along $\theta$ (denoted by $C_\theta$) is far steeper than along $\phi$ (denoted by $C_\phi$). The average curvature, given by $\sqrt{C_\theta C_\phi}$, is approximately $3 \times 10^{-2}$ K at $h = 1$ T, a value about the same order of magnitude as the one emerging from zero point fluctuations (quantum ObD$^3$).

Returning to the model A Eq. (1), this effective anisotropy combined with appropriate exchange parameters stabilizes the $\psi_2$ state (note that other phases, namely a canted ferromagnet as well as the antiferromagnetic Palmer and Chalker state can also be stabilized depending on the values of $K_{a,b,c}$). Here, the moment direction at each of the 4 sites of a primitive tetrahedron basis is given by $(x, x, y), (-x, -x, y), (-x, x, -y), (x, -x, -y)$ with $y \approx 2x$ (in the cubic frame). Note that this is allowed by symmetry for the $\psi_2$ state, but not for the other component ($\psi_3$) of the $\Gamma_5$ two-dimensional representation$^{[31]}$. The dynamical structure factor $S(Q, \omega)$ that exposes the spin dynamics is modeled by a random phase approximation (RPA) calculation (see Refs. [13,15] and Supplemental Material$^{[31]}$). For the relevant set of $K_{a,b,c}$ parameters (see below), numerical calculations show the opening of a spin gap $\Delta_{\text{RPA}} \approx 15$ meV at Brillouin zone centers. To emphasize explicitly the influence of the CEF
levels in causing this gap, calculations have been performed for $B_{nm}$ parameters multiplied by a renormalization coefficient $\eta$. This has the effect of rescaling all CEF energy gaps by $\eta$. In these calculations, the exchange coupling remains fixed, so that the molecular field and the Néel temperature are essentially unchanged, but the effective magnetic anisotropy inherited by admixing of the ground CEF doublet with excited CEF states decreases with increasing $\eta$. $S(Q, \omega)$ is also globally unchanged, yet the spin gap gradually decreases and tends to zero for large $\eta$. Numerical calculations show that $C_\theta \sim h$ while $C_\phi \sim h^5/\eta^4$. Ultimately, as $\eta \to \infty$, the $U(1)$ classical ground state degeneracy within the Hilbert space strictly composed of of a direct product of single-ion CEF ground doublets is recovered (see Fig. 1f). In that limit, quantum and thermal ObD would become the sole mechanism able to lift the accidental degeneracy.

Comparison with experiments – To determine the $K_{a,b,c,4}$ couplings, the INS data were fitted to the calculated $S(Q, \omega)$ within the RPA. The neutron measurements were performed on a large $\text{Er}_2\text{Ti}_2\text{O}_7$ single crystal grown with the floating zone technique. The crystal was inserted in a copper sample holder and attached on the cold finger of a dilution fridge, allowing one to cool the sample down to 50 mK. Data were collected on the IN5 time-of-flight instrument (ILL) which combines high flux with position sensitive detectors allowing for single crystal spectroscopy. Measurements were carried out with an incident neutron wavelength of 6 Å in zero field and under an applied magnetic field of 1.5 and 2.5 T along [1,−1,0]. The spin excitation spectra measured along the high symmetry directions of the cubic unit cell at 50 mK are shown in Fig. 2. These results compare well with prior measurements (see Ref. [19] and the supplemental material of Ref. [2]). Because of the magnetic $\psi_2$ domains, the identification of the expected four different spin wave branches is not straightforward. This means that the inelastic peaks in Fig. 2 contain several modes within the experimental resolution. This is evidenced in the high resolution set-up, using a wavelength of 8.5 Å. The highly Gaussian (nearly triangular) profile of the resolution line inherent to the counter-rotating disk choppers instrument, which yields an energy resolution of about 20 $\mu$eV, reveals two acoustic-like modes coming from different magnetic domains (see Supplemental Material). Using the same high resolution set-up, the
zero field data confirm the opening of a spin gap at zone centers: as shown in Fig. 2(c), the energy resolution permits to discriminate between the inelastic scattering and the strong Bragg intensity at the \( Q = (111) \) position. Above the elastic line, the neutron intensity first shows a dip and then a peak, a behavior that is typical of a spin gap. Fitting the data through a Lorentzian profile convoluted with the resolution function (see blue line in Fig. 2) yields \( \Delta_{G}^{\text{Sp}} \approx 43 \mu \text{eV} \). This value compares very well with previous estimates\(^{21-23} \).

To determine the exchange parameters, we calculate \( S(Q,\omega) \) assuming an equal population of the six \( \psi_2 \) magnetic domains. On the basis of exhaustive calculations as a function of the parameters \( K_{a,b,c,4} \) in zero and applied magnetic field, the INS data were fitted by matching the location of the maximum INS intensity in several directions. A good agreement is found for the following values:

\[
K_a \sim 0.003 \pm 0.005 \text{ K} \quad K_b \sim 0.075 \pm 0.005 \text{ K} \\
K_c \sim 0.034 \pm 0.005 \text{ K} \quad K_4 \sim 0 \pm 0.005 \text{ K.} (2)
\]

Many others sets that capture independently the magnetization or the excitation spectrum can be found but the present determination provides values that capture all these experimental data\(^{21} \). With these exchange parameters, the spin gap is evaluated at \( \Delta_{G}^{\text{RPA}} \approx 15 \mu \text{eV} \), a value smaller than \( \Delta_{G}^{\text{Sp}} \), but of the correct order of magnitude.

**Discussion** — It is instructive to compare our results (Eq. 2) with those found using the pseudospin 1/2 approach (Model B\(^{2} \)). The corresponding anisotropic exchange Hamiltonian is described in detail in Ref. \(^{2} \) and is based on the anisotropic couplings \( (J_{+\pm}, J_{\pm}, J_{z\pm}, J_{zz}) \) acting between pseudospin 1/2 components written in their local basis. The calculation of the dynamical structure factor for this model was performed within the Holstein-Primakov approximation, using the Spinwave software developed at the LLB\(^{25} \). Following the same fitting procedure as above, a set of parameters is obtained which largely confirm the results of Ref. \(^{2} \) (see Table I\(^{25} \)). The most striking point is that Models A and B lead to very similar \( S(Q,\omega) \). This is due to the fact that both models adopt a predominant effective Hamiltonian with bilinear couplings in terms of pseudospin 1/2 operators when projected in the CEF ground doublet.\(^{11} \)

Specifically, there is a relationship\(^{20} \) between the two sets of anisotropic exchange couplings based on the \( g_\perp \) and \( g_z \) Landé factors deduced from the ground state doublet wavefunction\(^{11} \). Table I\(^{11} \) which allows to compare \( K_{a,b,c,4} \) transformed in the \( (J_{+\pm}, J_{\pm}, J_{z\pm}, J_{zz}) \) language with the values determined from model B and from Ref. \(^{2} \), shows that the sets of values are similar whether determined from either model. Further, owing to the ObD mechanism, Model B leads to a spin gap \( \Delta_{G} = 21 \mu \text{eV} \) a value of the same order of magnitude as the one \( \Delta_{G}^{\text{RPA}} = 15 \mu \text{eV} \) obtained in Model A.

While the maps in Fig. 2 demonstrate an overall agreement with experiment, some discrepancies are observed nonetheless, which equally affect Models A and B. The most important difference concerns the acoustical like mode stemming from \((0,0,2)\), which seems to disperse continuously up to 0.45 meV in the neutron data. Within the experimental uncertainty, there is no gap opening when this branch crosses the optical one (see the arrows in Fig. 2(a), left and right columns of the \( H=0 \) panel). Such a gap opening occurs in the calculations, separating the acoustic branch from a higher energy optical branch. Furthermore, both models predict two well-separated modes at the zone centers \( Q = (1,1,1), (2,2,0) \) and \((0,0,2)\) at about 0.45 and 0.5 meV, whereas a single one is observed in experiment (middle column of the \( H=0 \) panel in Fig. 2(a)). More elaborate models are probably necessary to explain these features, taking into account the long-range part of the dipolar interaction or more complex coupling terms than bilinear ones.

Reference \(^{21} \) reports the evolution of the gap as a function of temperature, and ascertains that it varies as the square of the \( \psi_2 \) order parameter. The gap calculated in the framework of Model A shows instead a linear evolution with the order parameter (see Fig. 2(d)). The difference between the linear and squared order variations is most pronounced in a narrow temperature range spanning \( T_N \) to 0.6 \( \times T_N \). Unfortunately, in this temperature range, we believe that the experimental uncertainty in Ref. \(^{21} \) is too large to allow one to discriminate between the two behaviors. Further experiments are planned to shed light on this issue.

To conclude, the present study shows that the molecular field induced admixture between CEF levels generates an effective magnetic anisotropy as a plausible mechanism for an energetic selection of the \( \psi_2 \) state. The proposed model captures a number of key features of the inelastic neutron scattering data, including the opening of a spin gap. Its order of magnitude shows that the proposed mechanism appears as efficient as the ObD scenario, questioning the completeness of the projected pseudospin 1/2 model\(^{23-25,10} \) as a minimal model of \( \text{Er}_2\text{Ti}_2\text{O}_7 \). Our study raises the question whether the competing quantum\(^{21,10} \) and thermal\(^{10} \) order by disorder is the sole or even the principal mechanism for the selection of \( \psi_2 \) in this material, and whether its advocacy as a rare example of ObD\(^{21,10,12} \) will stand the test of time. On a more positive note, it seems plausible that quantum fluctuations and anisotropy induced by CEF admixing cooperate to select \( \psi_2 \) in \( \text{Er}_2\text{Ti}_2\text{O}_7 \). Conversely, one might ask whether their competition might be responsible in part for some of the perplexing properties of \( \text{Er}_2\text{Sn}_2\text{O}_7 \).
Acknowledgments

We acknowledge E. Lhotel, L. Jaubert and P. McClarty for fruitful discussions, as well as F. Damay for a careful reading of the manuscript. We also thank S. Turc (cryogeny group at ILL) for his technical help while setting up the magnet and the dilution fridge. Research at the Perimeter Institute for Theoretical Physics is supported by the Government of Canada through Industry Canada and by the Province of Ontario through the Ministry of Economic Development & Innovation.

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Supplemental Material

I. MEAN FIELD MODEL

Our mean field study follows the approach of Ref.$^{24}$ it is based on the following Hamiltonian for rare earth (R) moments $\vec{J}_i$ at site $i$ of the pyrochlore lattice:

$$\mathcal{H} = \mathcal{H}_{CEF} + \frac{1}{2} \sum_{i,j} \vec{J}_i \cdot \mathcal{K}_{i,j} \cdot \vec{J}_j$$

(3)
In this expression, $H_{CEF} = \sum_{nm} B_{nm} O_{nm}$ is written in terms of $O_{nm}$ Stevens operators with $B_{20} = 616$ K, $B_{40} = 2850$ K, $B_{43} = 795$ K, $B_{60} = 858$ K, $B_{43} = -494$ K, $B_{66} = 980$ K, in Weylbourne conventions. These parameters lead to CEF excitations at 6.2, 7.5, 13.8, 44.9, 47.7, 52.6 and 69.6 meV which compare well with the first levels determined experimentally through neutron scattering at 6.4, 7.3 and 15.4 meV.

$K = J + D$ is the sum of the anisotropic exchange tensor $J$ and of the $D$ dipolar interaction, limited to the contribution of the nearest-neighbors. Various conventions have been used to define $K_{i,j}$ Here, $K_{i,j}$ is defined in the $(\hat{a}, \hat{b}, \hat{c})$ frame linked with a R-R bond.

$$J_i \cdot K_{i,j} \cdot J_j = \sum_{\mu, \nu=x,y,z} J^\mu_i (K_{i,j}^\mu a_{i,j}^\nu + K_{i,j}^\nu b_{i,j}^\nu) + K_{i,j}^\mu \sqrt{2} \vec{b}_{i,j} \times (\vec{J}_i \times \vec{J}_j)$$

Considering for instance the pair of Er$^{3+}$ ions at $r_1 = (1/4, 3/4, 0)$ and $r_2 = (0, 1/2, 0)$, we define the local bond frame as: $\vec{a}_{12} = (0, 0, -1)$, $\vec{b}_{12} = 1/\sqrt{2}(1, -1, 0)$ and $\vec{c}_{12} = 1/\sqrt{2}(-1, -1, 0)$. This Hamiltonian, written in terms of bond-exchange constants, has the advantage to provide a direct physical interpretation of the different parameters. Note that $K_4$ is an anti-symmetric exchange constant (Dzyaloshinskii-Moriya like), while $K_{a,b,c}$ are symmetric terms. Owing to the form of the dipolar interaction, we have:

$$D_{i,j} = D_{nn} \left( a_{i,j} a_{i,j} + b_{i,j} b_{i,j} - 2 c_{i,j} c_{i,j} \right)$$

with $D_{nn} = \sum_{nn} (g_{i,j}^{nn} a_{i,j}^{nn})^2$ and where $r_{nn}$ is the nearest neighbour distance in the pyrochlore lattice. If we combine them with the anisotropic exchange constants $(J_a, J_b, J_c)$, we obtain:

$$K_a = J_a + D_{nn}$$
$$K_b = J_b + D_{nn}$$
$$K_c = J_c - 2 D_{nn}$$

As is usual in mean-field approximations, a self-consistent treatment is carried out to solve the problem: starting from a random configuration for the $\{\vec{J}_j\}$, the contribution to $H$ at site $i$ is diagonalized in the Hilbert space of the Er$^{3+}$ magnetic moment defined by the $\{|J_i\rangle\}$, $J_z = -15/2, ..., 15/2$ basis vectors. This yields the energies $E_{i,\mu}$ and the wave functions $|\phi_{i,\mu}\rangle$. The updated magnetic moments, $\langle \vec{J}_i \rangle'$, at each step of the iteration procedure, are given by:

$$\langle \vec{J}_i \rangle' = \sum_{\mu} e^{-E_{i,\mu}/k_B T} Z \langle \phi_{i,\mu} \rangle \langle \vec{J}_i \rangle |\phi_{i,\mu}\rangle$$

with

$$Z = \sum_{\mu} \exp -E_{i,\mu}/k_B T$$

These are then used to proceed to site $j$, and this is repeated until convergence.

II. RELATION WITH QUANTUM PSEUDO-SPIN HALF MODELS

This anisotropic exchange Hamiltonian can be rewritten in terms of couplings between the spin components of a pseudospin $1/2$ defined in subspace spanned by the ground CEF doublet states.

$$H = \sum_{i,j} J_{i,j} S_i^z S_j^z - J_{\pm} (S_i^+ S_j^- + S_i^- S_j^+) + J_{\pm}\left[ S_i^z (\gamma_{ij} S_j^z + \gamma_{ij} S_j^-) + \gamma_{ij} S_i^- S_j^\pm + i \leftrightarrow j \right]$$

$(J_{\pm, \pm, \pm, \pm})$ is the set of effective exchange parameters. Note that the “sanscrit” notation refers to local bases. The states of this pseudospin $1/2$ span this ground CEF wavefunctions doublet, using the relation:

$$g_j \vec{J} = g_\vec{S} \quad \text{or} \quad \vec{J} = \lambda \vec{S}$$

For the pyrochlores, of Fd3m space group, $\lambda = \frac{A}{g_j}$ matrix is diagonal and takes the form:

$$\lambda = \begin{pmatrix} \lambda_\perp & \lambda & \lambda_\perp \\ \lambda & \lambda_\perp & \lambda \end{pmatrix}$$

We finally obtain the following relations:

$$J_{zz} = \frac{\lambda_\perp^2 K_a - 2 K_c - 4 K_4}{3}$$
$$J_{\pm} = -\frac{\lambda_\perp^2}{12} \left[ 2 K_a - 3 K_b - K_c + 4 K_4 \right]$$
$$J_{\pm, \pm, \pm, \pm} = \frac{\lambda_\perp}{3\sqrt{2}} \left[ -2 K_a + 3 K_b - K_c + 4 K_4 \right]$$

and, conversely:

$$K_a = \frac{4}{3} J_{zz, \pm} + J_{\pm, \pm} + 4 \sqrt{2} J_{zz, \pm, \pm} + \frac{1}{3} J_{zz}$$
$$K_b = \frac{2}{3} J_{zz, \pm} + J_{\pm, \pm} + 4 \sqrt{2} J_{zz, \pm, \pm} + \frac{2}{3} J_{zz}$$
$$K_c = \frac{2}{3} J_{zz, \pm} + J_{\pm, \pm} + 4 \sqrt{2} J_{zz, \pm, \pm} + \frac{2}{3} J_{zz}$$

III. DETAILS ABOUT THE MAGNETIC STRUCTURE

The description of the possible magnetic structures in Er$_2$Ti$_2$O$_7$ is based on the symmetry analysis performed.
in the Fd3m space group for a \( \mathbf{k} = 0 \) propagation vector. Different structures are possible, belonging to different irreducible representations. As explained in the main text, \( \text{Er}_2\text{Ti}_2\text{O}_7 \) undergoes a transition towards an antiferromagnetic Néel phase below \( T_N \approx 1.2 \text{K} \). This ordered phase corresponds to the \( \psi_2 \) basis vector of the \( \Gamma_5 \) irreducible representation. The magnetic moment at site \( i \) is defined in a local frame \( (\hat{a}_i, \hat{b}_i, \hat{e}_i) \) given in Table II. Owing to symmetry, the moments in the \( \psi_2 \) configuration are of the form \((x, x, y), (-x, -x, y), (x, -x, y), (x, -x, -y)\). The experiment favors \( y = 2x \), hence a non-collinear structure, in which the magnetic moments are perpendicular to the local \((111)\) axes. In the general case, the projection of the moment along the CEF axis is:

\[ m_z = \frac{2x - y}{\sqrt{3}} \]

while the angle formed with the \( XY \) plane is:

\[ \cos \theta = \frac{1}{\sqrt{3}} \frac{(2x - y)}{2x^2 + y^2} \]

Assuming classical magnetic moments of equal amplitude \( m = \sqrt{2x^2 + y^2} \) on the different sites, and calculating the coupling matrices \( \mathbf{K} \), it is possible to calculate analytically the classical energy per spin in the \( \psi_2 \) state:

\[ E_c = \frac{1}{2} \left( 2K_a + 4K_a - 3K_b - 5K_c \right) + 3(2K_a + K_b + K_c) \cos 2\theta \]

In the \( XY \) case, \( y = 2x \):

\[ E_c = (2K_a - 3K_b - K_c)m^2 \]

| Site | 1 | 2 | 3 | 4 |
|------|---|---|---|---|
| CEF axis \( \hat{e}_i \) | (1,1,-1) | (-1,-1,1) | (1,-1,1) | (1,1,1) |
| Position \( (\hat{a}_i, \hat{b}_i, \hat{e}_i) \) | \( (\frac{1}{2}, \frac{1}{2}, 0) \) | \( (0, \frac{1}{2}, 0) \) | \( (0, \frac{1}{2}, \frac{1}{2}) \) | \( (\frac{1}{2}, \frac{1}{2}, \frac{1}{2}) \) |

TABLE II: \( (\hat{a}_i, \hat{b}_i, \hat{e}_i) \) frame for the different sites of a tetrahedron.

## IV. SPIN DYNAMICS IN THE RPA

The spin dynamics is calculated using the random phase approximation (RPA). Spin excitations are constructed on the basis of the transitions between the mean-field states \( |\phi_{i,\mu}\rangle \) of energy \( E_{i,\mu} \). These transitions occur between the set of CEF modes, perturbed or split by the molecular field.

Because of the interactions between the magnetic moments, these transitions acquire a dispersion. Two quantities are essential, namely the matrix element of \( \mathbf{J} \) that connects the two states |\( \phi_{i,\mu}\rangle \) and |\( \phi_{i,\nu}\rangle \):

\[ \tilde{a}_{\mu\nu} = \langle \phi_{i,\mu}|\mathbf{J}_i - \langle \mathbf{J}_i|\phi_{i,\nu}\rangle \]

and the energy of the transition \( (E_{i,\mu} - E_{i,\nu}) \).

Following the generalized susceptibility approach, it is useful to introduce labels associated with a transition \( s = (i, \mu \rightarrow \nu) \), as well as matrices \( A_s \) and energies \( \Delta_s \) so that:

\[ \Delta_s = E_{i,\nu} - E_{i,\mu} \]

and:

\[ A_s^{ab} = (n_{i,\mu} - n_{i,\nu}) \ u_{i\mu\nu}^a u_{i\nu\mu}^b \]

with \( n_{i,\mu} = \exp (E_{i,\mu}/k_B T)/Z \). The RPA generalized spin-spin susceptibility tensor \( \tilde{\chi}_{s,s'}(\mathbf{Q},\omega) \) is a solution of:

\[ \tilde{\xi}_s \delta_{s,s'} = \sum_{s''} \left[ \tilde{\xi}_{s,s''} - \tilde{\xi}_s \tilde{\chi}_{s,s''}(\mathbf{Q}) \right] \tilde{\chi}_{s'',s'}(\mathbf{Q}) \]

Here, \( \tilde{\xi}_s(\omega) \) is the \( 3 \times 3 \) single-ion spin-spin susceptibility:

\[ \tilde{\xi}_s(\omega)^{\alpha\beta} = \frac{A_s}{\hbar\omega + i\sigma^0 - \Delta_s} \]

The spin-spin correlation function \( S(\mathbf{Q},\omega) \) is then given by:

\[ S(\mathbf{Q},\omega) = \frac{|F(\mathbf{Q})|^2}{1 - \exp(-\hbar\omega/k_B T)} \sum_{\alpha\beta=x,y,z} \left( \delta_{\alpha\beta} - \frac{Q^\alpha Q^\beta}{Q^2} \right) \text{Im} \left[ \sum_{s,s'} \tilde{\chi}_{s,s'}(\mathbf{Q},\omega)e^{i\mathbf{Q}(\mathbf{r}_s - \mathbf{r}_s')} \right] \]

where \( \mathbf{r}_s' \) denotes the position of the ion at site \( i \) and \( F(\mathbf{Q}) \) is the magnetic form factor of the \( \text{Er}^{3+} \) magnetic ions.

We proceed by rewriting the RPA Eq. (17) using Eq. (18):

\[ A_s \delta_{s,s'} = \sum_{s''} \left( (\hbar\omega - \Delta_s)\delta_{s,s''} - A_s K_{s,s''} \right) \tilde{\chi}_{s'',s'}(\mathbf{Q}) \]

Owing to the definition of \( A_s \), it is possible to show that this matrix has just one non zero eigenvalue \( \alpha_s \):

\[ \alpha_s = \sum_a A_s^{aa} = (n_{i,\mu} - n_{i,\nu}) \sum_a |u_{i\mu\nu}^a|^2 \]

while the corresponding (normalized) eigenvector is nothing but \( \tilde{U}_s = \frac{\tilde{a}_s}{|\tilde{a}_s|} \). Multiplying Eq. (20) on the left and on the right by \( U_s^+ \) and \( U_{s'} \) respectively, we get:

\[ \alpha_s \delta_{s,s'} = \sum_{s''} \left( (\hbar\omega - \Delta_s)\delta_{s,s''} - \alpha_s U_{s'}^+ K_{s,s''} U_{s''} \right) \times (U_{s'}^+ \tilde{\chi}_{s'',s'} U_{s''}) \]

We are then left with an inversion problem to determine \( U_{s'}^+ \tilde{\chi}_{s'',s'} U_{s''} \) and then \( \tilde{\chi}_{s'',s'} \). From this equation, it is also worth noting that the pole of the susceptibility are given by the eigenvalues of the matrix \( R \):

\[ R_{s,s'} = \Delta_s \delta_{s,s'} + \alpha_s U_{s'}^+ K_{s,s''} U_{s''} \]
we therefore consider $\mathcal{K}_4 = 0$. Exhaustive calculations of
the scattering function $S(Q, \omega)$ have been performed as a
function of the remaining parameters $\mathcal{K}_{a,b,c}$, assuming
equipopulated $\psi_2$ magnetic domains at zero magnetic
field and only one domain at $H = 2.5$ T. Experimental
data were fitted by matching the location of the maxi-
mum neutron intensity in several directions. The spe-
tral weight repartition has also been taken into account,
checking that numerical intensities roughly match the
experimental ones. All the considered parameters giv-
ing a good agreement are reported in blue (resp. green)
for $H = 0$ T (resp. 2.5 T) in Figure 3. Interestingly,
while many sets of parameters capture the zero field
data, the ones obtained at $H = 2.5$ T appear to be
much restrictive: only a very small portion of param-
eter space accounts for the corresponding experimental
dispersions. This also points out very simple linear cor-
relations between parameters (which is better evidenced in
the inset): the relevant coupling constants are con-
tained into a plane, thus reducing the parameter space
to a surface. Different relations between the coupling
constants (or surfaces) are obtained considering the zero
$(\mathcal{K}_c = 1.90(2)\mathcal{K}_a - 2.53(5)\mathcal{K}_b + 0.218(2))$ or the finite mag-
netic field data $(\mathcal{K}_c = -2.0(3)\mathcal{K}_a - 4.4(6)\mathcal{K}_b + 0.37(5))$.
Combining these two latter relations reduces the relevant
parameter space to a line represented in red in the Fig.
3 whose parametric equations is:

\begin{align}
\mathcal{K}_a &= 0.138\mathcal{K}_c \quad (23) \\
\mathcal{K}_b &= 0.085 - 0.291\mathcal{K}_c \quad (24) \\
\mathcal{K}_c &= \mathcal{K}_c. \quad (25)
\end{align}

Along this line, the best set of param-
eters, fitting all the experimental data, is finally obtained
for $(\mathcal{K}_a, \mathcal{K}_b, \mathcal{K}_c, \mathcal{K}_d) = (0.003(5), 0.075(5), 0.034(5), 0.000(5))$ K (red point
in Fig. 3). We finally checked that this set of parameters
allows one to capture the magnetization curve measured
at $T = 110$ mK (see Fig. 4) except near the quantum
critical field driven transition at 1.6 T, where the mean-field
theory is not expected to be accurate.

VI. TIME OF FLIGHT EXPERIMENTS

Data were collected on the IN5 time-of-flight instru-
ment (ILL) in its single-crystal set up. Measurements
were carried out with an incident neutron wavelength
of 6 and 8.5 Å, in zero field and under an applied magnetic
field of 1.5 and 2.5 T along $[1, -1, 0]$ at 50 mK. Note that
Ref. [19] shows data along $[0, 0, L]$ while the main body
of Ref. [2] shows in field data only. Figure 5 shows
selected maps from the 8.5 Å data, allowing one to observe
the spin gap at the zone center $(1, 1, 1)$. Those data were
obtained by integrating the full 4D data-set in narrow
Q-range by steps of 0.05 “rlu”, and then reassembled
to produce the maps. The data by Ross et al. is shown

V. DETERMINATION OF THE COUPLING
CONSTANTS : FITTING PROCEDURE

A fitting procedure in a four parameters space may
be hazardous. Reducing the parameter space often re-
quires a combination of different experimental techniques
giving complementary information as well as a detailed
characterization of the system in function of external pa-
rameters, such as the temperature or the magnetic field.
Ref. [2] claims that Er$_2$Ti$_2$O$_7$ has a small Dzyaloshinskii-
Moriya interaction $\mathcal{K}_4 \ll \mathcal{K}_{a,b,c}$. Although it may slightly
improve the agreement between simulations and experi-
mental data, we checked that such a contribution has a
weak effect on the excitation spectrum. In what follows,
for comparison. Note that the latter was obtained with a resolution of 12 $\mu$eV$^{21}$, while it was 20 $\mu$eV in the present experiment. The two acoustic-like modes coming from different magnetic domains are clearly visible along (11$L$), along with the depletion of the intensity below the spin gap.

FIG. 5: (color online): (a) corresponds to TOF data by Ross et al. showing the filling of the spin gap with temperature at 50 and 700 mK, together with reconstructed fits$^{21}$. (b) and (c) shows the data obtained at 50 mK in the present work along (hh1) and (11$L$) respectively with an incident wavelength of 8.5 Å providing an energy resolution of 20 $\mu$eV.
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