Understanding Reentrance in Frustrated Magnets: the Case of the Er$_2$Sn$_2$O$_7$ Pyrochlore

D. R. Yahne,$^1$ D. Pereira,$^2$ L. D. C. Jaubert,$^3$ L. D. Sanjeeva,$^4$ M. Powell,$^5$ J. W. Kolis,$^5$ Guangyong Xu,$^6$ M. Enjalran,$^7$ M. J. P. Gingras,$^{2,8}$ and K. A. Ross$^{1,8}$

$^1$Department of Physics, Colorado State University, 200 W. Lake St., Fort Collins, CO 80523-1875, USA
$^2$Department of Physics and Astronomy, University of Waterloo, Waterloo, Ontario N2L 3G1, Canada
$^3$CNRS, Université de Bordeaux, LOMA, UMR 5798, 33400 Talence, France
$^4$Department of Materials Science and Engineering, University of Tennessee, Knoxville, TN 37996, USA
$^5$Department of Chemistry, Clemson University, Clemson, South Carolina 29634-0973, USA
$^6$NIST Center for Neutron Research, National Institutute of Standards and Technology, Gaithersburg, Maryland 20899, USA
$^7$Physics Department, Southern Connecticut State University, 501 Crescent Street, New Haven, Connecticut 06515-1355, USA
$^8$CIFAR, MaRS Centre, West Tower 661 University Ave., Suite 505, Toronto, ON, M5G 1M1, Canada

(Dated: January 22, 2021)

Reentrance, the return of a system from an ordered phase to a previously encountered less-ordered one as a controlled parameter is continuously varied, is a recurring theme found in disparate physical systems, from condensed matter to black holes. While diverse in its many incarnations and generally unsuspected, the cause of reentrance at the microscopic level is often not investigated thoroughly. Here, through detailed characterization and theoretical modeling, we uncover the microscopic mechanism behind reentrance in the strongly frustrated pyrochlore antiferromagnet Er$_2$Sn$_2$O$_7$. Taking advantage of the recent advance in rare earth stannate single crystal synthesis, we use heat capacity measurements to expose that Er$_2$Sn$_2$O$_7$ exhibits multiple instances of reentrance in its magnetic field $B$ vs. temperature $T$ phase diagram for magnetic fields along three cubic high symmetry directions. Through classical Monte Carlo simulations, mean field theory and classical linear spin-wave expansions, we argue that the origins of the multiple occurrences of reentrance observed in Er$_2$Sn$_2$O$_7$ are linked to soft modes. Depending on the field direction, these arise either from a direct $T = 0$ competition between the field-evolved ground states, or from a field-induced enhancement of the competition with a distinct zero-field antiferromagnetic phase. In both scenarios, the phase competition enhances thermal fluctuations which entropically stabilize a specific ordered phase. This results in an increased transition temperature for certain field values and thus the reentrant behavior. Our work represents a detailed examination into the mechanisms responsible for reentrance in a frustrated magnet and may serve as a template for the interpretation of reentrant phenomena in other physical systems.

PACS numbers:

Within the field of magnetism, frustration refers to a system’s inability to simultaneously satisfy all of its energetic preferences. Strong frustration can result in a variety of exotic phenomena such as spin liquids, spin ice, emergent quasi-particles, topological phases and order-by-disorder [1–7]. Most of the research focus in this area over the past thirty years has been devoted to investigating the physics near zero temperature, considering finite temperatures as a necessary modus operandi to search for signatures of the low-energy properties. However, even when subject to high frustration, a majority of frustrated magnetic materials ultimately develop long-range order or display spin-glass freezing at a nonzero critical temperature $T_c$, albeit often at a very low one compared to the spin-spin interactions. In this context, it therefore seems natural to ask what behavior near $T_c$ may be a witness of the zero-temperature ground state physics. This is particularly important when $T_c$ is just above the experimental baseline temperature, so that temperatures which are low relative to $T_c$ cannot be reached. Here we precisely consider such a situation, as arises in the Er$_2$Sn$_2$O$_7$ pyrochlore antiferromagnet, and which provides an opportunity to study a recurrent aspect of frustrated magnetic systems observed at nonzero temperature: reentrance [8–16].

Reentrance occurs when a system, after having developed an ordered phase of some sort, returns to its original less-ordered (e.g., paramagnetic) state as some parameter (e.g., temperature, field, pressure, stoichiometry) is continuously varied. Reentrance has been found in spin glasses [17, 18], liquid mixtures [19, 20], protein thermodynamics [21], liquid crystals [22, 23], bilayer graphene [24], superconductors [25], modulated phases [26, 27] and even in black hole thermodynamics [28]. Despite its ubiquity, reentrance is typically unexpected and its explanation in terms of entropic contributions to the free-energy from the underlying microscopic degrees of freedom is usually subtle. In this context, while frequently observed in frustrated magnets, the microscopic mechanism leading to reentrance often remains obscure [8–15]. Two mechanisms have commonly been invoked: a field-dependent suppression of quantum fluctuations [29–31] and the partial disorder of an intervening phase [18, 32–34]. Here, we present an alternative scenario of a generic nature which illustrates how the observation of reentrance may be used as a fingerprint of the frustration at play in the ground state.

In this article, we show that Er$_2$Sn$_2$O$_7$ represents a tractable material example where the intricate microscopic mechanisms responsible for reentrance in frustrated magnets can be rigorously studied experimentally and theoretically. Er$_2$Sn$_2$O$_7$ holds a special place among rare earth pyrochlores [5, 35]: it is well-characterized, has a suppressed critical temperature...
and is one of the few materials with a simple Palmer-Chalker (PC) antiferromagnetic ground state [36–38] [Fig. 1(a)]. Its estimated exchange and single-ion susceptibility parameters are highly anisotropic and theory suggests a proximity to another competing antiferromagnetic phase [37–39] known as \( \Gamma_5 \) [5, 6, 39] [Fig. 1(b,c)]. Because of this anisotropy, the response of \( \text{Er}_2\text{Sn}_2\text{O}_7 \) to an applied field is expected to differ with field direction, as has been poignantly illustrated with the experimental exploration of rare earth pyrochlore titanates [5, 6, 11, 12, 35]. As such, the present study critically relies on the recently gained ability to synthesize pyrochlore stannate single crystals [40], including \( \text{Er}_2\text{Sn}_2\text{O}_7 \).

We report herein the discovery of multiple occurrences of reentrance in the \( B-T \) phase diagram of \( \text{Er}_2\text{Sn}_2\text{O}_7 \) for fields along the \([100], [110] \) and \([111] \) cubic directions using heat capacity measurements. By thoroughly investigating this experimental phase diagram using mean field theory, classical linear spin-wave expansions and Monte Carlo simulations, we have uncovered the various microscopic origins of reentrance in this system. In short, we find that different types of multi-phase competitions at \( T = 0 \) result in enhanced thermal fluctuations that entropically stabilize the ordered phase, and thus increase \( T_c(B) \) over a certain \( B \) field range. This produces \( T_c(B) \) reentrant phase boundaries whose maximum temperature extent corresponds to the zero-temperature field-driven phase transitions [see Fig. 2(a-c)]. This multi-phase competition is in some cases a direct consequence of the competition of the field-evolved PC states while in others it is inherent to \( \text{Er}_2\text{Sn}_2\text{O}_7 \)’s zero-field ground state being in close proximity to the phase boundary between the PC and \( \Gamma_5 \) phases. See the Supplementary Material [41] for technical details on the experiments, simulations and analytics.

Heat capacity \( (C_p) \) measurements were performed on single crystal samples, grown via the hydrothermal method described in Ref. [40], down to 50 mK with varying magnetic fields, \( B \), oriented in the \([111], [110] \) and \([100] \) directions, using a dilution refrigerator insert in a Quantum Design Physical Properties Measurement System. Two measurement techniques were used: the conventional quasi-adiabatic thermal relaxation method (called "short pulses" hereafter), as well as "long pulses", both of which are described in detail in Ref. [45]. The long pulse technique allows faster and higher point-density measurements across phase transitions, enabling an accurate mapping of a phase diagram by measuring the field dependence of the transition temperature, \( T_c(B) \) [Fig. 1 (d)].

In the zero-field heat capacity [Fig. 1 (d) inset], we find a sharp lambda-like anomaly indicative of a phase transition at \( T_c = 118 \pm 5 \) mK, which is consistent with previous measurements on powder samples reported in Ref. [44] (130 mK, from heat capacity, data shown in Fig. 1 (d) inset) and Ref. [38] (108 \pm 5 mK, from DC magnetic susceptibility). The extremely high point density of the long pulse measurements allow for the observation of subtle features in the peak shape, which are typically not resolved by conventional short pulse measurements. This reveals a low temperature shoulder of the \( C_p \) peak in the zero-field data at \( 97 \pm 5 \) mK [41]. We performed elastic neutron scattering measurements to determine the magnetic structure between the sharp high temperature peak and the low temperature shoulder to check for an intermediate magnetic phase [41]. We found that the magnetic structure is of Palmer-Chalker type at all measured temperatures throughout the transition range with no sign of other magnetic phases. It is not clear what causes this structure in the heat capacity anomaly, but we note that similar (though not identical) broadening is observed in all five crystals we have measured as well as in published data on a powder sample [41, 44] (Fig. 1 (d) inset). Thus, it seems to be a feature of all \( \text{Er}_2\text{Sn}_2\text{O}_7 \) samples, but is likely due to (or influenced by) slight inhomogeneities rather than being purely intrinsic in origin. Although it may be worth future investigation, its presence does not affect any of the conclusions of this work.

To model \( \text{Er}_2\text{Sn}_2\text{O}_7 \), we use the generic nearest-neighbor Hamiltonian on the pyrochlore lattice [39, 46],

\[
\mathcal{H} = \sum_{(i,j)} J_{ij}^{\alpha\beta} S_i^\alpha S_j^\beta - \mu_B \sum_i g_i^{\alpha\beta} B^\alpha S_i^\beta.
\]

\( S_i = (S_i^x, S_i^y, S_i^z) \) is a three-component pseudo-spin of length \( |S_i| = 1/2 \) and \( B \) is the external magnetic field. The \( g \)-tensor represents the single-ion anisotropy, with local easy-plane \( g_\perp \) and easy-axis \( g_\parallel \) components at lattice site \( i \). Given

![FIG. 1: Example of 6-fold degenerate states: (a) Palmer-Chalker [36] and (b) \( \psi_2 \) and (c) \( \psi_3 \) basis states of \( \Gamma_5 \) [42]. The \( \psi_2 \) and \( \psi_3 \) states are connected by a rotation of the spins by an angle \( \phi \) within their local easy-planes (yellow circles): \( \phi \equiv n\pi/3 \) (\( +\pi/6 \)) for \( n = 0, ..., 5 \) correspond to \( \psi_2 (\psi_3) \) [43]. Panels (b) and (c) are for \( \phi = 0 \) and \( \pi/2 \), respectively. The manifold with \( U(1) \) degeneracy, \( \phi \in [0,2\pi] \), forms the so-called \( \Gamma_5 \) states that appear in the \([111] \) phase diagram. (d) Heat capacity, \( C_p(T) \), vs temperature, \( T \), of \( \text{Er}_2\text{Sn}_2\text{O}_7 \) with the magnetic field along \([100] \), showing the reentrant nature of the transition. Curves at different fields are offset vertically for clarity. (Inset) \( C_p(T) \) \( \text{Er}_2\text{Sn}_2\text{O}_7 \) in zero-field, with short and long pulse measurements on crystal samples overlaid. Powder data from Shirai et al. [44] is also overlaid to demonstrate agreement between sample types.](image-url)
FIG. 2: B-T phase diagrams of Er2Sn2O7 in the (a) [100], (b) [110] and (c) [111] field directions, comparing experimental data with sharp (□) and smooth (■) heat-capacity peaks [Fig. 1(d)], to Monte Carlo results with 1st (□) and 2nd (■) order transitions. Experiments and simulations are notably similar, showing the same (multiple) reentrance. The degeneracy \( Z_n \) found in simulations is given for each phase. The width of the red rectangles at 0 and 0.1 T represent the position of the double peaks. (d) In a [111] field, each of the six FEPC ground states has a \( \Gamma_5 \) contribution described by an angle \( \phi \) [Fig. 1(b,c)], that can be computed exactly by minimizing the energy of one tetrahedron as a function of \( B \).

The symmetries of the pyrochlore lattice, the anisotropic exchange matrix \( J_{ij}^{\alpha \beta} \) is parameterized by four independent coupling constants: \( \langle J_1, J_2, J_3, J_4 \rangle \) [46, 47]. Er2Sn2O7 has been previously parameterized using inelastic neutron scattering on powder samples [37, 38]. Here we choose to remain within the error bars of Ref. [38], selecting a set of coupling parameters where simulations find \( T_c \sim 180 \) mK at 0.4 T for a [111] field to match the experimental result [52]: \( \langle J_1, J_2, J_3, J_4 \rangle = (+0.079, +0.066, -0.111, +0.032) \) meV and \( g_{\perp} = 7.52, g_{\parallel} = 0.054 \).

To proceed, we first analyze this model using classical Monte Carlo simulations, with the results summarized in the B-T phase diagrams of Fig. 2. Most importantly, with \( T_c(B = 0.4 \text{T}) \) fitted (for \( B \) along the [111] direction), the simulations reproduce the number of reentrant “lobes” for each field direction (e.g. one and two for a [100] and [110] field, respectively), as well as, at each lobe, the rough magnitude of the increase of \( T_c \) at the corresponding value of \( B \). Moreover, simulations find that the transition always evolves from discontinuous to continuous when increasing the field. This is consistent with the shape of the experimental heat capacity peaks, evolving from sharp to smooth [Fig. 1(d) and Ref. [41]]. We suspect that a fine-tuning of the \( J_{ij}^{\alpha \beta} \) coupling parameters and incorporating quantum fluctuations and dipolar interactions should account for the quantitative disagreements. Nevertheless, the semi-quantitative match between experiments and simulations confirms the validity of Eq. (1) as a minimal model for Er2Sn2O7, suggesting it robustly encapsulates the key physics behind the experimentally observed multiple occurrences of reentrance.

The results in Figs. 2(a-c) raise multiple questions. Why are there multiple instances of reentrance and why are they so strongly dependent on the field direction? More fundamentally, why does Er2Sn2O7 demonstrate reentrance in the first place? As a set of clues, simulations bring to light a variety of phases that vie for ordering. In the rest of this article, we explain how soft modes induced by this multi-phase competition are linked to reentrance, using a combination of mean field theory and classical linear spin-wave expansions.

The zero-field ground state of Er2Sn2O7 is the sixfold-degenerate PC phase. However, the ground states naturally deform and evolve under the application of a magnetic field. For sufficiently large fields, some of these field-evolved PC (FEPC) states may become partially polarized into the same spin configuration. We therefore label the resulting phase according to the number of DEPC states that minimize the free energy but have distinct spin configurations (e.g. \( Z_6 \) at \( B = T = 0 \), for the six degenerate PC states, and \( Z_3 \) at sufficiently large \( B \) for the trivial field-polarized paramagnet). Phase transitions then occur whenever distinct FEPC states “merge” into the same spin configuration at a given field value \( B_c \).

First, consider the [100] field phase diagram in Fig. 2(a). At \( T = 0 \), the FEPC states merge at \( B_c = 0.82 \) T, giving rise to the yellow \( Z_2 \) region. Fig. 3 displays the classical spin-wave dispersions \( \kappa_{\nu}(q) \) for a number of field values below and above \( B_c \), calculated from the corresponding \( T = 0 \) FEPC ground states. As the merger transition is approached at \( B_c = 0.82 \) T, the bottom of the dispersive bands drop below the energy scale set by \( T_c(B = 0) \approx 160 \) mK, becoming soft and gapless at \( B = B_c \). This decrease indicates a propensity for stronger thermal fluctuations at \( B_c \) than at other field values. More precisely, since \( s = -\frac{1}{8N^*} \sum_q \sum_{\nu=1}^8 \ln(\kappa_{\nu}(q)) \) quantifies the entropy contribution from classical spin-waves, the decrease in \( \kappa_{\nu}(q) \) on approaching \( B_c \) from above or below (as shown in
Finally, the mechanisms behind reentrance for a [110] field are reminiscent of the other two field directions [Fig. 2(b)]. Below $B \leq 0.1$ T, simulations are difficult to thermalize, but above $B \geq 1.0$ T, we find two FEPC ground states that merge at $B_c = 0.42$ T. This merging gives rise to gapless soft modes, the subsequent violet $Z_1$ phase, and reentrance at finite temperatures. It is the same mechanism as in a [100] field. However, as opposed to the [100] scenario, this newly merged ground state vanishes immediately once $B > B_c$ (i.e., it becomes an excited state). The system is then found in two other ground states (corresponding to the yellow $Z_2$ phase). The vanishing of the merged state corresponds to the $Z_2$ phase abruptly disappearing above $B_c$, and the removal of the aforementioned gapless soft modes. This causes the rapid collapse of the reentrant lobe at $B \sim 0.42$ T. At higher field, there are no ground state FEPC mergers, but the gap in spin-wave excitations remains small for a broad field region around 0.7 T (see Section S5 in Ref [41]). These low-energy modes, concomitantly with the collapse at $B_c = 0.42$ T, are the reason for the higher field reentrant lobe at $\sim 0.7$ T.

In summary, we have presented the first exploration of the field-direction dependence of the thermodynamics of stannate pyrochlores, which, despite decades of effort, were not available as single crystals until very recently [40]. Access to these crystals has proven to be crucial since the phase diagram of $\text{Er}_2\text{Sn}_2\text{O}_7$ is highly sensitive to the field direction, and exhibits several reentrant lobes with sundry underlying mechanisms. These features result from the competition of several orders, especially the zero-field Palmer-Chalker, the field-induced ferromagnetic and the neighboring $\Gamma_5$ states. In particular, most instances of reentrance in the phase diagram can be traced to zero-temperature field-induced merging of distinct degenerate states, leading to soft modes which entropically enhance the transition temperatures. In this light, reentrance is a useful and experimentally accessible fingerprint of a zero-temperature phase transition.

Given that multi-phase competition is a common feature of frustrated magnetism, we expect the mechanisms we have uncovered to be widespread among magnetic systems displaying reentrance: especially since it does not require the accidental presence of an exotic partially-disordered phase [18, 33]. In semi-classical and quantum systems, our mechanism may work together with the field-induced suppression of quantum fluctuations [29] to produce even larger reentrant lobes. We hope our work will motivate others to pursue a microscopic interpretation of future observations of reentrance (and possibly to revisit old ones [8–16]) in light of zero-temperature transitions. Since magnetic systems often
afford us with minimal models to understand other areas of physics, our results raise a more general question: if reentrance is observed by varying a given parameter, when is it actually due to a nearby transition in a broader parameter space?

We acknowledge Natalia Perkins for useful discussions and Rob Mann for comments on reentrance in black holes. We thank Allen Scheie and Tom Hogan for their help with analyzing the long pulse measurements, and acknowledge the use of the LongHC Pulse program for this analysis. We thank I. Zivkovic and R. Freitas for providing us their C_p data previously published in Ref. [44]. We acknowledge the support of the National Institute of Standards and Technology, U.S. Department of Commerce, in providing the neutron research facilities used in this work. This research was partially supported by CIFAR. DRY, KAR, and JWK acknowledge funding from the Department of Energy award DE-SC0020071 during the preparation of this manuscript. The work at the University of Waterloo was supported by the Natural Sciences and Engineering Research Council (NSERC) and by the Canada Research Chairs Program (M.G, Tier 1). L. J. acknowledges financial support from CNRS (PICS No. 228338) and from the French ‘‘Agence Nationale de la Recherche’’ under Grant No. ANR-18-CE30-0011-01.

[1] L. Balents, “Spin liquids in frustrated magnets,” Nature 464, 199 (2010).
[2] M. J. P. Gingras and P. A. McClarty, “Quantum spin ice: a search for gapless quantum spin liquids in pyrochlore magnets,” Rep. Prog. Phys. 77, 056501 (2014).
[3] L. Savary and L. Balents, “Quantum spin liquids: a review,” Rep. Prog. Phys. 80, 016502 (2017).
[4] M. Hermanns, I. Kimchi, and J. Knolle, "Physics of the Kitaev model: Fractionalization, dynamic correlations, and material connections," Annu. Rev. Condens. Matter Phys. 9, 17 (2018).
[5] A. M. Hallas, J. Gaudet, and B. D. Gaulin, "Experimental insights into ground-state selection of quantum XY pyrochlores," Annu. Rev. Condens. Matter Phys. 9, 105 (2018).
[6] J. G. Rau and M. J. Gingras, “Frustrated Quantum Rare-Earth Pyrochlores,” Annu. Rev. Condens. Matter Phys. 10, 357 (2019).
[7] J. Knolle and R. Moessner, “A field guide to spin liquids,” Annu. Rev. Condens. Matter Phys. 10, 451 (2019).
[8] M. V. Gvozdikova, P.-E. Melchy, and M. E. Zhitomirsky, "Magnetic phase diagrams of classical triangular and kagomé antiferromagnets," J. Phys. Condens. Matter 23, 164209 (2011).
[9] R. Rawl, M. Lee, E. S. Choi, G. Li, K. W. Chen, R. Baumbach, C. R. dela Cruz, J. Ma, and H. D. Zhou, "Magnetic properties of the triangular lattice magnets A_2B_2B_2O_12 (A = Ba, Sr, La; B^2 = Co, Ni, Mn; B = W, Re)," Phys. Rev. B 95, 174438 (2017).
[10] A. Scheie, J. Kindervater, S. Säubert, C. Duvignac, C. Pfeiderer, H. J. Changlani, S. Zhang, L. Harriger, K. Arpino, S. M. Koolen, G. Grenier, J. Sichelschmidt, H. Yasuoka, A. M. Strydom, Y. Skourski, J. Wosnitza, et al., "Reentrant phase diagram of Yb_2Ti_2O_7 in a (111) magnetic field," Phys. Rev. Lett. 119, 127201 (2017).
[11] S. Säubert, A. Scheie, C. Duvignac, J. Kindervater, S. Zhang, H. J. Changlani, G. Xu, S. M. Koolen, G. Grenier, J. Sichelschmidt, H. Yasuoka, A. M. Strydom, Y. Skourski, J. Wosnitza, et al., "Anisotropic field-induced ordering in the magnetic phase diagram of Yb_2Ti_2O_7," Phys. Rev. B 101, 174434 (2020).
[12] O. A. Petrenko, M. R. Lees, G. Balakrishnan, and D. M. Paul, "Magnetic phase diagram of the antiferromagnetic pyrochlore Gd_2Ti_2O_7," Phys. Rev. B 70, 012402 (2004).
[13] L. Seabra, T. Momoi, P. Sindzingre, and N. Shannon, "Phase diagram of the classical Heisenberg antiferromagnet on a triangular lattice in an applied magnetic field," Phys. Rev. B 84, 214418 (2011).
[14] L. Seabra, P. Sindzingre, T. Momoi, and N. Shannon, "Novel phases in a square-lattice frustrated ferromagnet: ½-magnetization plateau, helicoidal spin liquid, and vortex crystal," Phys. Rev. B 93, 085132 (2016).
[15] M. Li, I. Rousochatzakis, and N. B. Perkins, "Reentrant incommensurate order and anomalous magnetic torque in the Kitaev magnet β-Li_2IrO_3," Phys. Rev. Res. 2, 033328 (2020).
[16] A. Rousseau, J.-M. Parent, and J. A. Quilliam, "Anisotropic phase diagram and spin fluctuations of the hyperkagome magnet Gd_2Ir_2O_7 as revealed by sound velocity measurements," Phys. Rev. B 96, 060411 (2017).
[17] M. J. P. Gingras and E. S. Sørensen, "Evidence for a genuine ferromagnetic to paramagnetic reentrant phase transition in a Potts spin-glass model," Phys. Rev. B 57, 10264 (1998).
[18] H. T. Diep, Magnetic Systems with Competing Interactions (Frustrated Spin Systems) (World Scientific Publishing, 1994).
[19] C. A. Vause and J. S. Walker, "Reappearing Phases," Sci. Am. 256, 98 (1987).
[20] C. A. Vause and J. S. Walker, "Effects of orientational degrees of freedom in closed-loop solubility phase diagrams," Phys. Lett. A 90, 419 (1982), ISSN 0375-9601.
[21] C. L. Dias, T. Ala-Nissila, J. Wong-ekkabut, I. Vattulainen, M. Grant, and M. Karttunen, "The hydrophobic effect and its role in cold denaturation," Cryobiology 60, 91 (2010).
[22] S. Singh, "Reentrant phase transitions in liquid crystals," Phase Transit. 72, 183 (2000).
[23] A. N. Berker and J. S. Walker, "Frustrated spin-gas model for doubly reentrant liquid crystals," Phys. Rev. Lett. 47, 1469 (1981).
[24] I. V. Lebedeva and A. M. Popov, "Two phases with different domain wall networks and a reentrant phase transition in bilayer graphene under strain," Phys. Rev. Lett. 124, 116101 (2020).
[25] T. H. Lin, X. Y. Shao, M. K. Wu, P. H. Hor, X. C. Jin, C. W. Chu, N. Evans, and R. Bayuzick, "Observation of a reentrant superconducting resistive transition in granular BaPbO_0.75BiO_{2.53}O_3 superconductor," Phys. Rev. B 29, 1493 (1984).
[26] A. Mendoza-Coto, L. Nicolao, and R. Díaz-Méndez, "On the mechanism behind the inverse melting in systems with competing interactions," Sci. Rep. 9, 2020 (2019), ISSN 2045-2322.
[27] A. Mendoza-Coto, D. E. B. de Oliveira, L. Nicolao, and R. Díaz-Méndez, "Topological phase diagrams of the frustrated Ising ferromagnet," Phys. Rev. B 101, 174438 (2020).
[28] N. Altamirano, D. Kubizňák, and R. B. Mann, "Reentrant phase transitions in rotating anti–de Sitter black holes," Phys. Rev. D 88, 101502 (2013).
[29] B. Schmidt and P. Thalmeier, "Néel temperature and reentrant H–T phase diagram of quasi-two-dimensional frustrated magnets," Phys. Rev. B 96, 214443 (2017).
[30] M. Skoulas, F. Rucker, G. J. Nilsen, A. Bertin, E. Pomjakushina, J. Ollivier, A. Schneidewind, R. Georgii, O. Zarkhoro, L. Keller, et al., "Putative spin-nematic phase in BaCdVO(PO_4)_2," Phys. Rev. B 100, 014405 (2019).
[31] K. M. Ranjith, S. Luther, T. Reinmann, B. Schmidt, P. Schlender, J. Sichelschmidt, H. Yasuoka, A. M. Strydom, Y. Skourski, J. Wosnitza, et al., "Anisotropic field-induced ordering in the
triangular-lattice quantum spin liquid NaYbSe$_2$,” Phys. Rev. B 100, 224417 (2019).

[32] V. Vaks, A. Larkin, and Y. N. Ovchinnikov, “Ising model with interaction between nonnearest neighbors,” Soviet Physics JETP 22, 820 (1966).

[33] P. Azaria, H. T. Diep, and H. Giacomini, “Coexistence of order and disorder and reentrance in an exactly solvable model,” Phys. Rev. Lett. 59, 1629 (1987).

[34] E. H. Boubcheur, R. Quartu, H. T. Diep, and O. Nagai, “Non-collinear XY spin system: First-order transition and evidence of a reentrance,” Phys. Rev. B 58, 400 (1998).

[35] J. S. Gardner, M. J. P. Gingras, and J. E. Greedan, “Magnetic pyrochlore oxides,” Rev. Mod. Phys. 82, 53 (2010).

[36] S. E. Palmer and J. T. Chalker, “Order induced by dipolar interactions in a geometrically frustrated antiferromagnet,” Phys. Rev. B 62, 488 (2000).

[37] S. Guitteny, S. Petit, E. Lhotel, J. Robert, P. Bonville, A. Forget, and I. Mirebeau, “Palmer-Chalker correlations in the XY pyrochlore antiferromagnet Er$_2$Sn$_2$O$_7$,” Phys. Rev. B 88, 134408 (2013).

[38] S. Petit, E. Lhotel, F. Damay, P. Boutrouille, A. Forget, and D. Colson, “Long-range order in the dipolar antiferromagnet Eu$_2$Sn$_2$O$_7$,” Phys. Rev. Lett. 119, 187202 (2017).

[39] H. Yan, O. Benton, L. Jaubert, and N. Shannon, “Theory of multiple-phase competition in pyrochlore magnets with anisotropic exchange with application to Y$_2$Ti$_2$O$_7$, Eu$_2$Ti$_2$O$_7$, and Er$_2$Sn$_2$O$_7$,” Phys. Rev. B 95, 094422 (2017).

[40] M. Powell, L. D. Sanjeeva, C. D. McMillen, K. A. Ross, C. L. Sarkis, and J. W. Kolis, “Hydrothermal crystal growth of rare earth tin cubic pyrochlores, RE$_2$Sn$_2$O$_7$ (RE = La-Lu); site ordered, low defect single crystals,” Cryst. Growth Des. 19, 4920 (2019).

[41] See Supplemental Materials for additional technical details and which also includes Refs. [? ? ? ].

[42] A. Poole and A. S. Wills and E. Lelièvre-Berna, “Magnetic order-
Supplemental Material for: “Understanding Reentrance in Frustrated Magnets: the Case of the Er$_2$Sn$_2$O$_7$ Pyrochlore”

D. R. Yahne,¹ D. Pereira,² L. D. C. Jaubert,³ L. D. Sanjeewa,⁴ M. Powell,⁵ J. W. Kolis,⁵ Guangyong Xu,⁶ M. Enjalran,⁷ M. J. P. Gingras,² and K. A. Ross¹,⁸

¹Department of Physics, Colorado State University, 200 W. Lake St., Fort Collins, CO 80523-1875, USA
²Department of Physics and Astronomy, University of Waterloo, Waterloo, Ontario N2L 3G1, Canada
³CNRS, Université de Bordeaux, LOMA, UMR 5798, 33400 Talence, France
⁴Department of Materials Science and Engineering, University of Tennessee, Knoxville, TN 37996, USA
⁵Department of Chemistry, Clemson University, Clemson, South Carolina 29634-0973, USA
⁶NIST Center for Neutron Research, National Institute of Standards and Technology, Gaithersburg, Maryland 20899, USA
⁷Physics Department, Southern Connecticut State University, 501 Crescent Street, New Haven, Connecticut 06515-1355, USA
⁸CIFAR, MaRS Centre, West Tower 661 University Ave., Suite 505, Toronto, ON, M5G 1M1, Canada

(Dated: January 22, 2021)

This document is a supplement to our main article and contains a number of brief sections to assist the reader with details. Specifically, we provide in turn:

- Section S1: A definition of the Palmer-Chalker states.
- Section S2: Additional heat capacity data for the different field directions.
- Section S3: Neutron scattering experimental details and results.
- Section S4: Details of the mean field theory used to describe the field-induced Palmer-Chalker state merger transitions.
- Section S5: Details of the classical spin-wave expansion.
- Section S6: Details of the Monte Carlo simulations and choice of model Hamiltonian parameters.
- Section S7: Details of the analysis for the phase diagram for fields along the [110] direction.
- Section S8: Details of the Monte Carlo simulations at low fields along the [110] direction.
- Section S9: Details of the Monte Carlo simulations for fields along the [111] direction.
S1. DEFINITION OF PALMER-CHALKER STATES

There are six Palmer-Chalker states in zero field [Fig. S1.1]; the spin configurations for three of these (on the four sublattices of a tetrahedron) are outlined in Table S1.1. The remaining three, denoted as \( \langle xy \rangle, \langle xz \rangle, \text{ and } \langle yz \rangle \), can be obtained from the listed three by reversal of the spins.

![Image](image_url)

FIG. S1.1: The six Palmer-Chalker states, labeled from left to right \( \langle xz \rangle, \langle yz \rangle, \langle xy \rangle \) (first row) and \( \langle xz \rangle, \langle yz \rangle, \langle xy \rangle \) (second row).

| \( \langle xy \rangle \) | \( \langle xz \rangle \) | \( \langle yz \rangle \) |
|-----------------|-----------------|-----------------|
| \( m_0 = \frac{1}{\sqrt{2}}(1, -1, 0) \) | \( m_0 = \frac{1}{\sqrt{2}}(1, 0, -1) \) | \( m_0 = \frac{1}{\sqrt{2}}(0, 1, -1) \) |
| \( m_1 = \frac{1}{\sqrt{2}}(-1, -1, 0) \) | \( m_1 = \frac{1}{\sqrt{2}}(-1, 0, -1) \) | \( m_1 = \frac{1}{\sqrt{2}}(0, -1, 1) \) |
| \( m_2 = \frac{1}{\sqrt{2}}(1, 1, 0) \) | \( m_2 = \frac{1}{\sqrt{2}}(-1, 0, 1) \) | \( m_2 = \frac{1}{\sqrt{2}}(0, -1, 1) \) |
| \( m_3 = \frac{1}{\sqrt{2}}(-1, 1, 0) \) | \( m_3 = \frac{1}{\sqrt{2}}(1, 0, 1) \) | \( m_3 = \frac{1}{\sqrt{2}}(0, 1, 1) \) |

TABLE S1.1: Spin configurations for three of the six Palmer-Chalker states on the four sublattices \( i = 0, 1, 2, 3 \) of a tetrahedron. Each state lies within a plane in the global frame of reference (e.g. \( \langle xy \rangle \) lies in the \( xy \)-plane). The remaining three Palmer-Chalker states are obtained by spin reversal. The convention for labeling the sublattices follows the one from Ref. [1].
S2. ADDITIONAL HEAT CAPACITY DATA

Figure S2.1 shows the raw zero-field long pulse heat capacity data for three different single crystals polished in the three high symmetry directions. As discussed in the main text, we find a sharp peak at $118 \pm 5 \text{ mK}$ with a low-temperature shoulder around $97 \pm 5 \text{ mK}$. A similar broad feature can be seen in polycrystalline measurements by Ref. [2], but the broadness of the peak is not discussed therein. The heat capacity measurements in field are shown in Fig. S2.2 for fields applied along the three cubic directions, the transition temperatures of which are included in Fig. 2 in the main text.

FIG. S2.1: Raw zero field heat capacity data for the three crystals polished in each of the high symmetry directions. We find that all samples show a low temperature shoulder, but the pronouncement of the shoulder varies between samples. Inset: Example of a typical crystal that was used in heat capacity measurements.

FIG. S2.2: Raw long pulse heat capacity data for fields along the three high symmetry cubic directions, leading to the phase diagram shown in the main text. Note, all data at nonzero field have been shifted vertically for clarity.
Elastic neutron scattering measurements were performed at the NIST Center for Neutron Research using the SPINS triple axis spectrometer. Approximately 4 g of crystals were ground into a fine powder and placed in a copper cigar type foil construction which was subsequently filled with 10 atm of He gas at room temperature. We collected elastic scattering scans at 50 mK, 110 mK, and 8 K on the (111), (002), (220), (113) and (222) Bragg peaks using $E = 5$ meV neutrons with an energy resolution of approximately 0.25 meV. The collimation settings were guide-open-80$'$-open, with Be filters placed before and after the sample.

The 50 mK data was taken to corroborate the Palmer-Chalker ground state found from powder samples by Ref. [3], as it should be well into the ordered phase. Data at 110 mK was taken as it lies in between the high temperature and low temperature peaks we find in the heat capacity curve (118 and 97 mK respectively), as discussed in the main text, and could possibly show evidence for an intermediate phase. We find that the correlations are Palmer-Chalker throughout the broad transition, as evidenced by the $Q = (111), (002)$ and (220) Bragg peaks. In particular, there was no hint of $\Gamma_5$ order, which may have been a likely intermediate phase candidate due to the material’s proposed proximity to the phase boundary between PC and $\Gamma_5$ [3,4]. No (002) intensity is expected for the $\Gamma_5$ phase (Fig S3.1 (b)), and we do not see any evidence of an enhanced (111) nor (220) peak that could indicate coexistence of the $\Gamma_5$ and PC ($\Gamma_7$) phase. The fit to the Palmer-Chalker structure is shown in Fig S3.1 (a) and is in agreement with the Ref. [3] powder diffraction data. We note that only intermediate phases with ordering wave vector $q = 0$ were investigated, and it is still possible that a $q \neq 0$ intermediate phase could exist.

![Elastic neutron scattering data (black) on SPINS. Data was taken at 50 mK and 110 mK, and 8 K data was used for background subtraction. The fit to the PC phase is shown (red), as well as the difference between the data and fit (blue). (b) Calculated magnetic diffraction patterns for the $\Gamma_5$ and PC ($\Gamma_7$) configurations (nuclear contributions not shown). The $\Gamma_5$ pattern has been shifted vertically for clarity.](image)

**FIG. S3.1:** (a) Elastic neutron scattering data (black) on SPINS. Data was taken at 50 mK and 110 mK, and 8 K data was used for background subtraction. The fit to the PC phase is shown (red), as well as the difference between the data and fit (blue). (b) Calculated magnetic diffraction patterns for the $\Gamma_5$ and PC ($\Gamma_7$) configurations (nuclear contributions not shown). The $\Gamma_5$ pattern has been shifted vertically for clarity.
The following derivation of variational mean field theory for a spin system on the pyrochlore lattice, with an exchange Hamiltonian and an applied magnetic field, follows Refs. [5–7]. The exchange Hamiltonian can be written as

\[ H_0 = \frac{1}{2} \sum_{a,b} \sum_{i,j} \sum_{\mu,\nu} J_{ia,jb}^{\mu \nu} S_{ia}^{\mu} S_{jb}^{\nu}, \]  

(S4.1)

where \( a \) and \( b \) denote the chosen tetrahedra, \( i \) and \( j \) denote the sublattices chosen within those tetrahedra, and \( \mu \) and \( \nu \) denote the components of the (classical) spin vectors. The exchange matrix \( J_{ia,jb}^{\mu \nu} \) is completely general in this form. In this work, we have predominantly only taken into account the global-frame nearest-neighbor exchange parameters that are defined for rare-earth pyrochlore systems with a Kramers ground state doublet [1]. However, this general form can also include the dipole-dipole interaction (see further below at the end of this section). With an applied magnetic field, this Hamiltonian becomes

\[ H = \frac{1}{2} \sum_{a,b} \sum_{i,j} \sum_{\mu,\nu} J_{ia,jb}^{\mu \nu} S_{ia}^{\mu} S_{jb}^{\nu} - \sum_{i,a} h_{ia} \cdot S_{ia}, \]  

(S4.2)

where

\[ h_{ia}^{\mu} = \mu g_{ia}^{\mu \nu} B_{\nu}, \]  

(S4.3)

is the scaled magnetic field for a given applied magnetic field \( B_{\nu} \), \( g_{ia}^{\mu \nu} \) is the g-tensor of the ion at the given sublattice \( i \), expressed in the global frame of Ref. [1] and \( \mu_B \) is the Bohr magneton. This g-tensor is independent of the tetrahedron \( a \) and only depends on which sublattice \( i \) is considered [1]. Henceforth, all physical quantities with energy dimension are measured in K (Kelvin) units and, as such, we correspondingly set the Boltzmann constant \( k_B = 1 \).

The mean field free energy is given by

\[ F_\rho = Tr(\rho H) + T Tr(\rho \ln \rho), \]  

(S4.4)

where \( \rho \) is the many-body density matrix and the trace is computed over all spin configurations. The variational mean field approximation assumes \( \rho(\{S_{ia}\}) = \prod_{i,a} \rho_{ia}(S_{ia}), \) where \( \rho_{ia} \) is the density matrix for a single site’s spin. These single-site density matrices are then treated as variational parameters, subject to the constraints of normalization (\( Tr(\rho_{ia}) = 1 \)) and the order parameter definition (\( Tr(\rho_{ia} S_{ia} = m_{ia}) \)). Enforcing these constraints using the sets of Lagrange multipliers \( \{\xi_{ia}\} \) and \( \{A_{ia}\} \) yields

\[ F(\{\rho_{ia}\}, \{\xi_{ia}\}, \{A_{ia}\}) = Tr(\rho H) + T Tr(\rho \ln \rho) - T Tr \left\{ \sum_{i,a} \xi_{ia}(\rho_{ia} - 1) \right\} - T Tr \left\{ \sum_{i,a} (\rho_{ia} S_{ia} - m_{ia}) \cdot A_{ia} \right\}. \]  

(S4.5)

Minimizing the free energy with respect to the variational parameters, the single-site density matrices are found to be \( \rho_{ia} = \frac{1}{Z_{ia}} e^{A_{ia} \cdot S_{ia}}. \) The partition function \( Z_{ia} \) is computed by integrating over all spin configurations in spherical coordinates, given that the spins are treated classically (that is, as continuous vectors); this yields \( Z_{ia} = \frac{\sqrt{2}}{\beta} \sinh(A_{ia}), \) where \( A_{ia} = |A_{ia}|. \) Using all of the above relations and computing the traces in Eq. (S4.5), the free energy simplifies to

\[ F = \frac{1}{2} \sum_{a,b} \sum_{i,j} \sum_{\mu,\nu} J_{ia,jb}^{\mu \nu} m_{ia}^{\mu} m_{jb}^{\nu} - \sum_{i,a} h_{ia} \cdot m_{ia} + \sum_{i,a} (H_{ia} \cdot m_{ia} - \frac{1}{\beta} \ln(Z_{ia})), \]  

(S4.6)

where \( H_{ia} \equiv A_{ia} \) can be considered as a local field. Explicitly, by minimizing the free energy with respect to the order parameters \( m_{ia}, \frac{\partial F}{\partial m_{ia}} = 0, \) it can be shown that

\[ H_{ia}^{\mu} = -\sum_{j,b,\nu} J_{ia,jb}^{\mu \nu} m_{jb}^{\nu} + h_{ia}^{\mu}. \]  

(S4.7)

Using this expression for \( H_{ia}^{\mu}, \) the free energy, \( f, \) averaged over all \( N \) sites of the lattice is

\[ f = \frac{F}{N} = \frac{X(m)}{N} - \frac{1}{N} \sum_{i,a} \ln(Z_{ia}), \]  

(S4.8)

where

\[ X(m) \equiv -\frac{1}{2} \sum_{a,b} \sum_{i,j} \sum_{\mu,\nu} J_{ia,jb}^{\mu \nu} m_{ia}^{\mu} m_{jb}^{\nu}, \]  

(S4.9)

Lastly, using the identity \( \frac{\partial f}{\partial m_{ia}} = -m_{ia} \) for the order parameter at each site, one finds

\[ m_{ia} = \frac{H_{ia}}{|H_{ia}|} \left[ \coth(\beta |H_{ia}|) - \frac{1}{\beta |H_{ia}|} \right]. \]  

(S4.10)
This equation relates the local field $H_{ia}$ with the order parameter $m_{ia}$ at each site, which can be calculated self-consistently.

The evolution of a chosen Palmer-Chalker state (e.g. $\langle xy \rangle$) in the presence of an applied magnetic field $B$ is accomplished within variational mean field theory as follows. For a given magnitude and direction of the field (and at zero temperature), the tetrahedra of the pyrochlore lattice are initiated with an initial spin configuration. At zero-field, this initial spin configuration is that of the chosen Palmer-Chalker state. At finite field, this initial spin configuration is the mean field solution from the previous field magnitude. For example, if the mean field calculation is done for $B$ values incremented by 0.01 T, the initial configuration used for $B = 0.11$ T is the mean field solution from $B = 0.10$ T. In this way, the evolution of the chosen zero-field Palmer-Chalker state can be tracked as a function of $B$. The self-consistency equation Eq. (S4.10) is then solved iteratively for zero temperature until convergence is attained. The resulting spin configuration is denoted with a subscript $h$ (e.g. $\langle xy \rangle_h$) to represent that it is the field-evolved configuration of the originally-chosen Palmer-Chalker state. This field-evolved Palmer-Chalker (FEPC) state for a given $B$ can then be used to tile the pyrochlore lattice and solve the self-consistency equation for finite temperatures for the same $B$ value, showing how the chosen Palmer-Chalker state evolves with temperature as well. This process is followed for each of the six parent $B = 0$ Palmer-Chalker states, in order to track their field- and temperature-evolution individually.

The mean field calculations are performed for a pyrochlore lattice of size $L = 2$ (in terms of cubic unit cells) for a total of $16L^3 = 128$ sites. At all fields and temperatures, the spin configuration is always found to be $q = 0$ ordering wave vectors. The phases can be labelled by (i) how many of the six FEPC states minimize the free energy (which may be less than six), and (ii) how many of these degenerate FEPC states have distinct spin configurations. If there are $n$ distinct spin configurations out of all degenerate FEPC states, then phase is labelled as a $Z_n$ phase, reflecting the order of the discrete degeneracy. Phase transitions are then indicated by a reduction in this discrete symmetry as the field increases, referred to as “merger” transitions. The mean field phase diagram for the $[100]$, $[111]$, and $[110]$ field directions, as well as a pictorial representation of the merger transitions at zero temperature, are illustrated in Fig. S4.1.

Variational mean field theory including dipolar interactions has also been performed by use of the Ewald summation method [6, 8, 9]. When including the dipolar interactions, the $g$-tensor must be properly incorporated to map the interaction (between dipoles) onto the pseudospin $S = \frac{1}{2}$ representation. As well, it should be noted that the nearest-neighbor part of the dipolar interaction is already included in the nearest-neighbor exchange couplings ($J_1, J_2, J_3, J_4$). With the inclusion of dipolar interactions, the critical temperatures and fields are reduced relative to Fig. S4.1, but the $q = 0$ orders, $Z_n$ phases, and overall topology of the phase diagrams remain unchanged.
FIG. S4.1: Zero temperature representations of field-evolved Palmer-Chalker (FEPC) mergers ((a), (c), and (e)) and mean field $B$-$T$ phase diagrams ((b), (d), and (f)) for the [100], [111], and [110] field directions, respectively. In the FEPC merger diagrams, vertical grey lines represent critical fields of the merging transitions; the vertical blue line represents field-induced selection of the $\psi_2$ phase out of the accidentally-degenerate $\Gamma_{5}$ manifold; vertical red lines represent changes in the set of degenerate FEPC states. Solid lines denote the FEPC states that are degenerate and minimize the energy; dotted lines represent FEPC states that are excited and do not minimize the energy. In the mean field phase diagrams, filled black squares represent a phase transition to a phase of different discrete symmetry due to merging; open red squares represent a change in the set of degenerate FEPC states. In both sets of diagrams, $Z_n$ denotes the discrete symmetry of the phase, which has $n$ distinct spin configurations out of the degenerate FEPC states. FP-PM denotes the field-polarized paramagnetic phase; PC denotes Palmer-Chalker order. Note that the zero temperature phases in the [110] field direction differ from the finite temperature phases due to entropic effects, for reasons discussed further in Section S7.
S5. CLASSICAL SPIN-WAVE EXPANSION

A classical spin-wave expansion for \( q = 0 \) spin ordering on the pyrochlore lattice and without an applied field is detailed in Ref. [1]; the derivation below for a classical spin-wave expansion in a finite field is similar. Informed by the mean field results at zero temperature, \( q = 0 \) ordering is assumed and dipolar interactions, which do not change the topology of the phase diagram, are excluded. Under these assumptions, the Hamiltonian is

\[
\mathcal{H} = \frac{1}{2} \sum_{ij} J_{ij}^{\mu\nu} S_i^{\mu} S_j^{\nu} - \mu_B \sum_i \varepsilon_i S_i^{\mu} B^{\mu},
\]

(S5.1)

where \( J_{ij}^{\mu\nu} \) is now restricted to only nearest neighbors, and the tetrahedra indices are suppressed because of the \( q = 0 \) assumption.

Starting with the classically-ordered ground state spin configuration on a single tetrahedron at \( T = 0 \), define the local coordinate system at each sublattice with \( \{ u_i, v_i, w_i \} \), where the classical spin vector is of length \( S \) and points along \( w_i \). The other two local unit vectors are defined arbitrarily, so long as mutual orthogonality is satisfied. The fluctuations about this ordered spin vector, on sublattice \( i \) and tetrahedron \( a \), can then be expressed as

\[
S_{ia} = \begin{bmatrix} \sqrt{S} \delta u_{ia} \\ \sqrt{S} \delta v_{ia} \end{bmatrix} \approx \begin{bmatrix} \sqrt{S} \delta u_{ia} \\ S - \frac{1}{2} \delta u_{ia}^2 - \frac{1}{2} \delta v_{ia}^2 \end{bmatrix}.
\]

(S5.2)

Substituting this perturbed spin into the original Hamiltonian and expanding to quadratic order in the fluctuations yields

\[
\mathcal{H} = \frac{1}{2} N_s \sum_{ij} S^2 (u_i \cdot \mathbf{J}_{ij} \cdot w_j) - \mu_B S N_s \sum_i (u_i \cdot \mathbf{g}_i \cdot \mathbf{B}) + \frac{S}{2} \sum_{ij} \delta u_{ia} \delta u_{ja} (u_i \cdot \mathbf{J}_{ij} \cdot u_j) + \frac{S}{2} \sum_{ij} \delta v_{ia} \delta v_{ja} (v_i \cdot \mathbf{J}_{ij} \cdot v_j) + \frac{1}{2} \mu_B \sum_{ij} (\delta u_{ia}^2 + \delta v_{ia}^2) (u_i \cdot \mathbf{g}_i \cdot \mathbf{B}).
\]

(S5.3)

The first two terms just represent the ground state energy, \( \varepsilon_0 \), where \( N_s \) is the number of tetrahedra in the system (\( N_s = \frac{N}{4} \), where \( N \) is the number of spins in the system). The rest of the expression encapsulates the effect of fluctuations from exchange and Zeeman interactions.

Performing a Fourier transform over the reciprocal lattice vectors \( q \) of the FCC lattice and defining \( u(q) \equiv (\delta u_1(q), \delta u_2(q), \delta u_3(q), \delta u_4(q), \delta v_1(q), \delta v_2(q), \delta v_3(q), \delta v_4(q)) \), the fluctuation contribution to the Hamiltonian can be written as:

\[
\mathcal{H} = \varepsilon_0 + \frac{1}{2} \sum_q u(-q) \left( \mathcal{M}(q) + \mathcal{N}(q) \right) u(q).
\]

(S5.4)

The matrices \( \mathcal{M}(q) \) and \( \mathcal{N}(q) \) can be written in block matrix form, composed of four separate \( 4 \times 4 \) blocks. They are:

\[
\mathcal{M}^{(\mu\nu)} = 2S \begin{bmatrix} M^{11}(q) & M^{12}(q) \\ M^{21}(q) & M^{22}(q) \end{bmatrix}
\]

(S5.5)

\[
M^{11}_{ij} = \cos (q \cdot r_{ij})(u_i \cdot \mathbf{J}_{ij} \cdot u_j - \delta_{ij} \sum_l w_l \cdot \mathbf{J}_{lj} \cdot w_j)
\]

\[
M^{12}_{ij} = \cos (q \cdot r_{ij})(u_i \cdot \mathbf{J}_{ij} \cdot v_j)
\]

\[
M^{21}_{ij} = \cos (q \cdot r_{ij})(v_i \cdot \mathbf{J}_{ij} \cdot u_j)
\]

\[
M^{22}_{ij} = \cos (q \cdot r_{ij})(v_i \cdot \mathbf{J}_{ij} \cdot v_j - \delta_{ij} \sum_l w_l \cdot \mathbf{J}_{lj} \cdot w_j)
\]

\[
\mathcal{N}^{(\mu\nu)} = \mu_B \begin{bmatrix} N^{11}(q) & 0 \\ 0 & N^{22}(q) \end{bmatrix}
\]

(S5.6)

\[
N^{11}_{ij} = N^{22}_{ij} = \delta_{ij} w_i \cdot \mathbf{g}_i \cdot \mathbf{B}.
\]

Diagonalizing this harmonic spin-wave Hamiltonian yields the classical spin-wave dispersions \( \kappa_{\mu i}(q) \) as a function of the wavevector \( q \). Since the classical spin-wave Hamiltonian is quadratic, the partition function and free energy of the classical
spin-waves can be calculated exactly. Assuming the above diagonalization has been performed to find the normal modes \( \phi_\nu(q) \) and dispersions \( \kappa_\nu(q) \), the partition function and free energy are given by

\[
Z = \int \left[ \prod_q \prod_{\nu=1}^8 d\phi_\nu(q) \right] e^{-\frac{1}{T}(\varepsilon_0 + \frac{1}{2} \sum_q \sum_{\nu=1}^8 \kappa_\nu(q) \phi_\nu(q) \phi_\nu(-q))}
\]

\[
= e^{-\frac{2\varepsilon_0}{T}} \prod_q \prod_{\nu=1}^8 \sqrt{\frac{2\pi T}{\kappa_\nu(q)}}
\]

\[\Rightarrow F = \varepsilon_0 + \frac{T}{2} \sum_q \sum_{\nu=1}^8 \ln(\kappa_\nu(q)) - NT \ln(2\pi T).\]  

(S5.7)

(S5.8)

(S5.9)

If there are \( N_q = \frac{N}{4} \) wavevectors in the sum, then the free energy per spin is

\[f = \frac{\varepsilon_0}{N} + \frac{T}{8N_q} \sum_q \sum_{\nu=1}^8 \ln(\kappa_\nu(q)) - T \ln(2\pi T).\]  

(S5.10)

Note that the entropy per spin \( s \) can be calculated using \( s = -\frac{\partial f}{\partial T} \). Computing this, keeping only the terms that depend on the spin-wave dispersions \( \kappa_\nu(q) \), yields

\[s = -\frac{1}{8N_q} \sum_q \sum_{\nu=1}^8 \ln(\kappa_\nu(q)).\]  

(S5.11)

The dispersions \( \kappa_\nu(q) \) for each of the FEPC states are shown below at various relevant choices of the applied field (e.g. at the merger transitions).
FIG. S5.1: Classical spin-wave dispersions for all Palmer-Chalker states that are degenerate in their energy, for a given field direction and magnitude. The shaded grey box represents energy scales below the zero-field Monte Carlo critical temperature of $T_c \approx 160$ mK. The chosen wavevectors are taken from the Brillouin zone of the FCC lattice. Note that some curves may overlap at high-symmetry points or due to their degeneracies in a field (e.g. $\langle xy \rangle_h$ and $\langle xy \rangle_h$ for $B \geq 0.82$ T along the [100] direction).
Monte Carlo simulations are performed on systems of classical Heisenberg spins with $N = 16L^3$ sites, where $L^3$ is the number of cubic unit cells. The spin length is $|S| = 1/2$. Several update algorithms are used together: the heatbath method, over-relaxation and parallel tempering. Parallel tempering is done every 100 Monte Carlo steps (MCS) and overrelaxation is done at every MCS. Thermalization is made in two steps: first a slow annealing from high temperature to the temperature of measurement $T$ during $t_e$ MCS followed by $t_e$ MCS at temperature $T$. After thermalization, measurements are done every 10 MCS during $t_m = 10 t_e$ MCS.

The characteristics of our simulations are typically:

- $4 < L < 10$,
- $10^6 \leq t_m \leq 5 \times 10^7$ MCS,
- 100 different temperatures (regularly spaced) between 0 and 200 mK for parallel tempering.

Please note that different exchange parameters for Er$_2$Sn$_2$O$_7$ have been investigated, within the error bars of Ref. [3]. In particular, a systematic search in the [111] direction has shown a consistent increase of the transition temperature between 0 and 0.4 T, confirming the robustness of the reentrance in this exchange-parameter region.
S7. ANALYSIS OF THE [110] RESULTS

Here we discuss the Monte Carlo phase diagram in a [110] field (Fig. 2(b) of the main text). There are two occurrences of reentrance, as well as a pronounced dip in $T_c$ that occurs between them. As with the [100] and [111] phase diagrams, these three features can also be understood as originating from soft modes that arise at the merging of $T = 0$ FEPC states. In this field and at $T = 0$, the six FEPC states do not all minimize the energy (for reasons discussed later on in this section). Therefore, when determining the merging of FEPC states, we primarily consider those ground states which simultaneously minimize the energy (as shown in Fig. S4.1(e)), since these are the states the system can be found in. Hence, as $B$ is increased from 0 and approaches a merging transition (between $(\mathbf{x}_\perp B)_h$ and $(\mathbf{y}_\perp B)_h$) around $B \approx 0.4$ T, soft modes arise (see Fig. S5.1(e)) and increase the thermal fluctuations within the ordered phase, producing the first (lower) reentrance. As $B$ continues to increase to and above $B \approx 0.42$ T, however, the merging FEPC states that produced these soft modes (namely, $(\mathbf{x}_\perp B)_h$ and $(\mathbf{y}_\perp B)_h$) are no longer energetically preferred (see Fig. S4.1(e) and Fig. S5.1(f)). These soft modes are therefore removed from the system, taking away their entropic support and producing a dip in $T_c$. As $B$ is increased further and enters the region $0.57 \, T < B < 1.12 \, T$, soft modes still arise (see Fig. S5.1(g)), although there are no merger transitions between FEPC ground states, only between ground and excited states (as shown in Fig. S4.1(e)). These soft modes give rise to the second (upper) reentrance. Since they are sustained for a range of $B$, the upper reentrant lobe is broad in the $B$ direction.

Although the mechanisms of reentrance are similar to the other two directions, the mean field theory results for the [110] field direction do stand out among the three field directions we studied, given that the zero temperature phases differ from those at finite temperature. The objective of the remainder of this section is to clarify this difference.

Consider very low temperatures $T = \epsilon$ where $\epsilon \to 0^+$. When $T = 0$ exactly, Eq. (S4.8) reduces to just the energy, whereas Eq. (S4.10) implies that $m_{ia} = \frac{H_{ia}}{T}$. For $T = \epsilon$, $m_{ia} \approx \frac{H_{ia}}{T}$. As well, the free energy now gains a contribution from the entropy, namely $-\frac{1}{N\beta} \sum_{i,a} \ln(Z_{ia})$. Using $A_{ia} = \beta |H_{ia}|$ in the partition function:

$$-\frac{1}{N\beta} \sum_{i,a} \ln(Z_{ia}) = -\frac{1}{N\beta} \sum_{i,a} \ln\left(\frac{4\pi e}{\beta |H_{ia}|} \sinh(\beta |H_{ia}|)\right)$$

$$= -\frac{\epsilon}{N} \sum_{i,a} \ln\left(\frac{4\pi e}{|H_{ia}|}\right) + \ln\left(\sinh\left(\frac{|H_{ia}|}{\epsilon}\right)\right).$$

As $\epsilon \to 0^+$, $\epsilon \ln(\epsilon) \to 0$ as well. Hence, the first term in the above summation is not an important contribution to the entropy at $T = 0^+$. On the other hand, $\sinh\left(\frac{|H_{ia}|}{\epsilon}\right)$ scales roughly as $e^{\frac{|H_{ia}|}{\epsilon}}$ for $\epsilon \to 0^+$. Hence:

$$-\frac{1}{N\beta} \sum_{i,a} \ln(Z_{ia}) \approx -\frac{\epsilon}{N} \sum_{i,a} \ln\left(e^{\frac{|H_{ia}|}{\epsilon}}\right) \approx -\frac{1}{N} \sum_{i,a} |H_{ia}|. \quad (S7.1)$$

When temperature becomes finite, the entropic contribution is therefore related to the average magnitude of the local fields.

As shown in Eq. (S4.7), there are two contributions to this local field: the local field resulting from the exchange between moments, and the local field resulting from the Zeeman coupling to the applied field $B$. When $B = 0$, only the first contribution is active. For the exchange parameters used here, the exchange couplings create local fields that move the spins into their Palmer-Chalker states, hence making these the ground state configurations. When $B$ is turned on, we must consider the second contribution, given by Eq. (S4.3). Assuming a pure easy-plane anisotropy on each sublattice, there are four $g$-tensors to consider:

$$\tilde{x}_y = \frac{g_1}{3} \begin{pmatrix} 2 & -1 & -1 \\ -1 & 2 & -1 \\ -1 & -1 & 2 \end{pmatrix}, \quad \tilde{y}_y = \frac{g_1}{3} \begin{pmatrix} 2 & 1 & 1 \\ 1 & 2 & -1 \\ 1 & -1 & 2 \end{pmatrix},$$

$$\tilde{x}_z = \frac{g_1}{3} \begin{pmatrix} 2 & 1 & -1 \\ 1 & 2 & 1 \\ -1 & 1 & 2 \end{pmatrix}, \quad \tilde{y}_z = \frac{g_1}{3} \begin{pmatrix} 2 & -1 & 1 \\ 1 & 2 & 1 \\ 1 & -1 & 2 \end{pmatrix}. $$

For a [110] field, a remarkable coincidence occurs: the Zeeman contribution $\mu_B g_{ia}^{\mu\nu} B^\nu$ for two sublattices (namely, sublattices 1 and 2) lie within the $xy$ plane, exactly parallel to the spin configuration of the $(\mathbf{x}_y)$ and $(\mathbf{y}_y)$ Palmer-Chalker states in Table S1.1. As such, for low fields in the [110] direction, these spins do not cant out of their original positions; the local fields from the exchange and Zeeman contributions add in parallel. Returning to Eq. (S7.1), this lack of canting is the origin of the slight entropy difference between the Palmer-Chalker states that lie in the $xy$-plane (where the two contributions add in parallel) and out of the $xy$-plane (where the two contributions are not parallel). A similar discrepancy should be expected at $T = 0$: $m_{ia} = \frac{H_{ia}}{T}$ at $T = 0$, so this lack of canting will discriminate between the $xy$-planar and non-$xy$-planar states. This effect is therefore the origin of the difference between the $T = 0$ and $T = 0^+$ phases. It should be noted that this is not an order by disorder effect. The $Z_2$ symmetry found at low fields and at $T = 0^+$ is not a subset of the original $Z_4$ symmetry found at low fields and at $T = 0$. Rather, the $Z_2$ symmetry corresponds to the two $xy$-planar Palmer-Chalker states, whereas the $Z_4$ symmetry corresponds to the other four Palmer-Chalker states.
FIG. S8.1: At low field in the [110] direction, there are four FEPC ground states $\{\langle xz \rangle_h, \langle yz \rangle_h, \langle xz \rangle_h, \langle yz \rangle_h \}$ and two FEPC excited states $\{\langle xy \rangle_h, \langle xy \rangle_h \}$ [see Fig. S4.1]. The very small energy gap between them means that it is not obvious which order is stabilised at finite temperature. The easiest way to differentiate between the ground and excited states in Monte Carlo simulations is via the $z$-component of the Palmer-Chalker order parameter, $m_{PCz}$ [see (a,b,c) panels]. $m_{PCz}$ is finite for the two excited states, while it is zero for the four ground states. Here we show that as the system size is increased, $m_{PCz}$ vanishes for $B = 0.1$ and 0.15 T, which means the FEPC excited states are not stable for $B \gtrsim 0.1$ T. For $B = 0.05$ T on the other hand, simulations are difficult to thermalize and the evolution with system size is not monotonic. Hence, we cannot rule out the presence of the FEPC excited states at finite temperature, as predicted by mean field theory in the previous section. That being said, this possible co-existence would only arise at very low field, $B \lesssim 0.1$ T, and thus not affect the mechanism for reentrance.

As for the four FEPC ground states, they can be divided into two pairs. For $\langle xz \rangle_h$ and $\langle yz \rangle_h$, the $x$-component of the Palmer-Chalker order parameter is positive, $m_{PCx} > 0$. For $\langle xz \rangle_h$ and $\langle yz \rangle_h$, $m_{PCx} < 0$. As shown in panels (d,e,f), $m_{PCx}$ always gets more and more positive for large system sizes, which means that the $Z_2$ states at low [110] field are $\langle yz \rangle_h$ and $\langle yz \rangle_h$. At very low temperatures, $T < 40$ mK, the data split into two groups because of the broken ergodicity in simulations between $\langle xz \rangle_h$ and $\langle yz \rangle_h$.

All of these simulations were done for $t_m = 5 \times 10^7$ Monte-Carlo steps.
S9. MONTE CARLO SIMULATIONS IN THE [111] DIRECTION

FIG. S9.1: Here we reproduce the phase diagrams in a [111] field as signalled via the (a) Palmer-Chalker and (b) $\Gamma_5$ order parameters, as well as the quantity (c) $m_{\phi6}$ (defined below) to differentiate between $\psi_2$ and $\psi_3$ states. The contour of the phase diagram is clearly visible in panels (a) and (b). However, as explained in the main text, in order to understand the reentrance at $B \sim 0.5\, T$, we need to consider the evolution of the angle $\phi$. The $\psi_2$ ($\psi_3$) states are characterized by $\phi \equiv n\pi/3$ ($+\pi/6$) for $n = 0, \ldots, 5$. Hence, the quantity $m_{\phi6} \equiv m_{\Gamma5} \cos \phi$ is equal to $+1$ ($-1$) for $\psi_2$ ($\psi_3$) states. The yellow region at finite temperature for $B \sim 0.5\, T$ thus indicates the dominance of $\psi_2$ order in this region, as discussed in the main text.

All of these simulations were done for $L = 6$ and $t_{\text{m}} = 5 \times 10^7$ Monte-Carlo steps.

---

1. H. Yan, O. Benton, L. Jaubert, and N. Shannon, "Theory of multiple-phase competition in pyrochlore magnets with anisotropic exchange with application to Yb$_2$Ti$_2$O$_7$, Er$_2$Ti$_2$O$_7$, and Er$_2$Sn$_2$O$_7," Phys. Rev. B 95, 094422 (2017).
2. M. Shirai, R. S. Freitas, J. Lago, S. T. Bramwell, C. Ritter, and I. Živković, "Doping-induced quantum crossover in Er$_2$Ti$_{2-x}$Sn$_x$O$_7," Phys. Rev. B 96, 180411(R) (2017), ISSN 24699969.
3. S. Petit, E. Lhotel, F. Damay, P. Boutrouille, A. Forget, and D. Colson, "Long-range order in the dipolar antiferromagnet Er$_2$Sn$_2$O$_7," Phys. Rev. Lett. 119, 187202 (2017).
4. S. Guitteny, S. Petit, E. Lhotel, J. Robert, P. Bonville, A. Forget, and I. Mirebeau, "Palmer-Chalker correlations in the XY pyrochlore antiferromagnet Er$_2$Sn$_2$O$_7," Phys. Rev. B 88, 134408 (2013).
5. J. N. Reimers, A. J. Berlinsky, and A.-C. Shi, "Mean-field approach to magnetic ordering in highly frustrated pyrochlores," Phys. Rev. B 43, 865 (1991).
6. M. Enjalran and M. J. P. Gingras, "Theory of paramagnetic scattering in highly frustrated magnets with long-range dipole-dipole interactions: The case of the Tb$_2$Ti$_2$O$_7$ pyrochlore antiferromagnet," Phys. Rev. B 70, 174426 (2004).
7. M. Enjalran, A. Del Maestro, and M. J. P. Gingras, "Mean-Field theory of the soft mode spectrum and field-driven transitions in a dipolar Heisenberg pyrochlore antiferromagnet model," unpublished.
8. P. P. Ewald, "Die Berechnung optischer und elektrostatischer Gitterpotentiale," Ann. Phys. (Leipzig) 64, 253 (1921).
9. J. Oitmaa, R. R. P. Singh, B. Javanparast, A. G. R. Day, B. V. Bagheri, and M. J. P. Gingras, "Phase transition and thermal order-by-disorder in the pyrochlore antiferromagnet Er$_2$Ti$_2$O$_7$: A high-temperature series expansion study," Phys. Rev. B 88, 220404 (2013).