Spin ice Thin Film: Surface Ordering, Emergent Square ice, and Strain Effects

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(Received 24 June 2016; published 19 May 2017)

Motivated by recent realizations of $Dy_2Ti_2O_7$ and $Ho_2Ti_2O_7$ spin ice thin films, and more generally by the physics of confined gauge fields, we study a model spin ice thin film with surfaces perpendicular to the [001] cubic axis. The resulting open boundaries make half of the bonds on the interfaces inequivalent. By tuning the strength of these inequivalent "orphan" bonds, dipolar interactions induce a surface ordering equivalent to a two-dimensional crystallization of magnetic surface charges. This surface ordering may also be expected on the surfaces of bulk crystals. For ultrathin films made of one cubic unit cell, once the surfaces have ordered, a square ice phase is stabilized over a finite temperature window. The square ice degeneracy is lifted at lower temperature and the system orders in analogy with the well-known *F* transition of the 6-vertex model. To conclude, we consider the addition of strain effects, a possible consequence of interface mismatches at the film-substrate interface. Our simulations qualitatively confirm that strain can lead to a smooth loss of Pauling entropy upon cooling, as observed in recent experiments on $Dy_2Ti_2O_7$ films.

DOI: 10.1103/PhysRevLett.118.207206

Introduction.—Highly frustrated magnets host an astonishing array of exotic many-body phenomena, taking us far from the conventional paradigms of collective magnetic behavior [1]. The formulation of the local frustrated constraints in terms of gauge fields has revolutionized our perspective on these systems. Depending on the system, such gauge structure can take the form of electromagnetism [2–5] with photon and magnetic-monopole excitations [5–13], or be similar to quantum chromodynamics [14] and linearized general relativity [15], as well as support phase transitions lying outside the Ginzburg-Landau-Wilson framework [16–19].

As vividly exposed in classic texts on electromagnetism [20], boundaries dramatically influence the behavior of gauge fields. It is therefore natural to ask what may be the role of boundary conditions in frustrated magnets described by gauge theories. Classical spin ice [21] presents itself as a nascent paragon to lay the foundations of such concepts, motivated by recent experimental studies of spin ice thin films of Dy₂Ti₂O₇ [22,23] and Ho₂Ti₂O₇ [24,25], along with the promising possibilities offered by spin ice heterostructures [26,27] and pyrochlore-iridates thin films [28–34]. This work was further motivated by the repeated reports of exotic phenomena arising at the interfaces of oxide heterostructures [35].

In this paper, we study the dipolar spin ice (DSI) model [36] in thin film (slab) geometry (slab DSI) defined by Eq. (1) below. As illustrated in Fig. 1, the slab geometry

renders the nearest-neighbor bonds on the surface inequivalent: either a bond belongs to a "bulk tetrahedron" or it is an "orphan bond" belonging to a "virtual tetrahedron" which has been "cleaved away" at the interface. Each virtual tetrahedron may therefore carry a magnetic surface charge that can propagate to and from the surface into the bulk. The orphan bonds act as effective chemical potentials for surface charges, allowing the slab DSI to undergo a surface charge-ordering transition. This is monopole crystallization [37] in two dimensions, driven by the magnetic Coulomb potential between surface charges, which originates from the dipolar interactions. This crystallization is limited to the microscopic surface layer without penetration into the bulk. The thinnest slab where this phenomenology can be explored is one cubic unit-cell thick, containing three layers of tetrahedra. For such thickness, below the surface ordering temperature, T_{so} , the central layer emerges in the form of a constrained square ice system with extensive degeneracy and correlations characteristic of a two-dimensional Coulomb phase [38]. Dipolar interactions ultimately lift the degeneracy of the Coulomb phase similarly to the venerable antiferroelectric *F*-model [39]. To conclude, we explore the possibility for strain effects to lift the residual Pauling entropy, as suggested by experiments on Dy₂Ti₂O₇ films [22].

Slab dipolar spin ice.—We consider thin films of the pyrochlore lattice of corner-sharing tetrahedra and whose surfaces are normal to the [001] cubic axis (Fig. 1). The

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FIG. 1. Spin ice thin film, thickness $L_h = 1$ in the [001] direction. Open boundaries lead to orphan bonds (green) on the surfaces. Each spin amounts to a pair of \oplus and \ominus effective magnetic charges ($\pm q$) [8]. Negative δ_o favors magnetic surface charges $Q = \pm 2q$. Below the surface ordering temperature, at $T < T_{so}$, the central tetrahedra (orange) develop into a square ice phase, as seen by the projection of the spins in plane. Shown, one of the ground states below the *F* transition ($T < T_F$). r_{nn} is the nearest-neighbor distance.

magnetic moments are described by classical Ising pseudospins $S_i \equiv \sigma_i \hat{z}_i$ [40], where \hat{z}_i is the local easy axis and $\sigma_i = \pm 1$ [41]. The spins interact via nearest-neighbor couplings *J*, dipolar interactions *D*, surface effects δ_o on orphan bonds, and strain effects δ_d on all bonds in the *xy* planes parallel to the surfaces over the whole system:

$$\mathcal{H} = J \sum_{\langle ij \rangle} \sigma_i \sigma_j + \delta_o \sum_{\langle ij \rangle_{\text{orphan}}} \sigma_i \sigma_j + \delta_d \sum_{\langle ij \rangle_{xy}} \sigma_i \sigma_j + Dr_{nn}^3 \sum_{i>j} \left(\frac{\hat{z}_i \cdot \hat{z}_j}{r_{ij}^3} - \frac{3(\hat{z}_i \cdot \vec{r}_{ij})(\hat{z}_j \cdot \vec{r}_{ij})}{r_{ij}^5} \right) \sigma_i \sigma_j.$$
(1)

 δ_o and δ_d are perturbations *in addition* to the coupling J.

The bulk DSI model ($\delta_o = \delta_d = 0$) is characterized by a low energy band of states with extensive entropy [42,43] in which each tetrahedron satisfies the ice rules with two spins pointing inwards and two spins pointing outwards (two-intwo-out) [21]. This is a Coulomb phase with Pauling entropy [38]. Topological excitations out of the Coulomb phase carry either a single (three-in-one-out or three-outone-in) or a double (four-in or four-out) gauge charge. Dipolar interactions endow these topological excitations with an effective (monopole) magnetic charge $\pm Q$. This is the dumbbell model where each moment is recast as a pair of positive or negative magnetic charges $\pm q$ sitting in the centers of the adjoining tetrahedra such that Q = 2q [Fig. 1] [8]. Within the dumbbell model, the low energy ice states are exactly degenerate. The ordering in bulk DSI [44,45] is due to corrections to this dumbbell description [8,43].

Slab DSI differs from bulk DSI [46] because of the open boundaries in the [001] direction, the presence of orphan bonds ($\delta_o \neq 0$), and strain effects ($\delta_d \neq 0$). Given the rapid variation of exchange constants with distance and geometry [40], we consider here a full range of parameter values, of the order of the bulk couplings themselves. Nevertheless, the dumbbell model remains a useful description if one also considers surface charges (Fig. 1). To set an experimental context, we use J = -1.24 K and D = 1.41 K, as in a minimal model of Dy₂Ti₂O₇ [36].

Method.—We use Monte Carlo simulations with parallel tempering [47,48] and a loop algorithm adapted to account for dipolar interactions [44,45] and monopoles [37,49]. The system size is $L \times L \times L_h$ in units of the 16-site cubic unit cell, with thickness $L_h < L = 8$. We take open boundary conditions along the [001] cubic axis and periodic ones along the other two cubic axes. As in other simulations of dipolar spin ice [45], we use the Ewald summation method to describe the long-range dipolar interactions. The slab geometry is implemented here by inserting an empty space, as large as needed, between replicas in the [001] direction [50,51]. We chose an empty space of $L_e = 1000$ unit cells. No difference was observed over the temperature range considered upon varying L_e for $L_e \gg L > L_h$.

Surface ordering.—We first set $\delta_d = 0$. By varying δ_o with respect to a threshold value $\delta_o^c = -J - 1.945(2)D$, one can favor either antiferromagnetic or ferromagnetic configurations on orphan bonds. For $\delta_o > \delta_o^c$, this extra surface energy scale manifests itself in the specific heat as a Schottky anomaly at $T \sim \delta_o$ (Fig. 2). Bulk physics is then rapidly recovered as L_h increases. For $\delta_o < \delta_o^c$, a sharp feature appears in the specific heat at an intermediate temperature T_{so} whose height decreases with increasing L_h , suggesting a surface rather than a bulk transition, as discussed below.

One can think of δ_o as a chemical potential shift compared with the bulk, inciting charges on the surface. Indeed, the ferromagnetic (respectively, antiferromagnetic) alignment of the pseudospins on an orphan bond amounts to a net magnetic charge, $\pm Q$ (respectively, vacuum) placed above the orphan bond at its mid point (Fig. 1). The charge sites form a square array with a lattice constant of $2r_{nn}$. The dipolar interaction between spins generates an effective Coulomb interaction between the surface charges [8], so that each surface of the thin film can be thought of as a (square) Coulomb lattice gas. The surface charges can propagate into the bulk, which increases the density of monopoles above that of the bulk DSI at intermediate temperature [Fig. 3(b)]. At low temperature, when varying δ_o below δ_o^c , the surface state transforms from charge



FIG. 2. Specific heat C_h for films ($\delta_o = +4$ K in orange and -4 K in violet) of different thickness [$L_h = 1$ (solid triangle), 2 (solid circle), 3 (square with dot inside)] and for bulk DSI (bulk DSI, black). Inset: low temperature C_h for $\delta_o = -4$ K. Temperature axes: log (main), linear (inset). As the thickness of the film L_h increases, the transition T_F approaches the bulk $T_c \approx 0.18$ K and becomes first order [44,45].

vacuum to charge crystal with a checkerboard pattern in order to minimize the Coulomb potential (Figs. 1 and 3). A suitable order parameter is $M_{so} = |M^b| + |M^t|$, with $M^{b,t} \equiv \sum_{\alpha} Q_{\alpha}^{b,t} \epsilon_{\alpha}/(2L^2)$, where $Q_{\alpha}^{b,t}$ is a bottom (b) or top (t) surface charge and $\epsilon_{\alpha} = \pm 1$ accounts for the bipartite nature of the square array. Finite size effects are discussed in the Supplemental Material [52].

Such surface order is not limited to thin films and could also occur on a sufficiently pristine [100] surface of bulk crystals. The enhancement of monopole density at intermediate temperature should also persist within a thin layer below the surface of spin ice compounds, which could be manipulated by a magnetic field [10]. Near-surface dynamical properties could possibly be probed by β -NMR [53,54] or μ SR with slow muons [55].

Emergent square ice for $L_h = 1$.—Below T_{so} , the system enters a temperature regime in which most tetrahedra respect the ice rules. Considering a tetrahedron on the top layer, the orientation of the two surface spins becomes fixed by the surface ordering (Fig. 1). The two lower spins are not fixed, but are coupled by the ice rules, forming a composite \mathbb{Z}_2 degree of freedom with projection along \pm [110]. The same holds for the tetrahedra on the bottom layer, with composite spins projected along $\pm [\bar{1}10]$. These composite projections now form the famous 6-vertex model [56] whose vertices correspond to the middle-layer tetrahedra respecting the ice rules (Fig. 1). This is confirmed by an entropy plateau for intermediate temperature at $S_{sq} =$ $\frac{1}{4}(\frac{3}{4}\ln\frac{4}{3})$ [Fig. 3(a)], which is the exact square ice entropy [57] (the composite spins account for 1/4 of the original degrees of freedom), and is visible as pinch points in the structure factor [52].

Ground states.—Just as in bulk DSI [44,45], the degeneracy of the Coulomb phase is eventually lifted at sufficiently low temperature, here T_F , to give a long-range



FIG. 3. Two-step ordering in thin films of thickness $L_h = 1$ and $\delta_o = -4$ K: (a) specific heat and entropy, (b) order parameters and density ρ of single charges (blue) inside the thin film, i.e., not on the surfaces. The latter is noticeably higher than for the bulk-DSI model (black). The surface ordering at $T_{so} \approx 900$ mK stabilizes a square ice model in the middle of the slab, as confirmed by the entropy in the intermediate region delimited by the dashed lines. Dipolar interactions ultimately lift the square-ice entropy at $T_F \approx 300$ mK (orange) in favor of the same antiferromagnetic ground state as the *F* model [39]. Temperature axes: log.

ordered ground state [Figs. 1 and 2 (inset)]. However, the 12-fold degenerate ground states of bulk-DSI are inequivalent in slab geometry, as only four of them support surface charge order. When $\delta_o < \delta_o^c$, these four states are quasidegenerate ground states; the energy remains quasidegenerate if all spins on either one or both surfaces of the film are flipped. When $\delta_o > \delta_o^c$, orphan bonds favor four of the bulk-DSI ground states with no surface charges.

When $L_h = 1$, the transition at T_F is equivalent to the \mathbb{Z}_2 symmetry breaking of the antiferroelectric F model [39] [Figs. 1 and 3(b)]. The order parameter is $M_F = \sum_i \sigma_i \eta_i / N'$ where $\eta_i = \pm 1$ transcribes the \mathbb{Z}_2 ordering of the $N' = 8L^2$ spins of the middle layer of tetrahedra (Fig. 1). The transition is characterized by a very smooth specific heat, as opposed to the sharp first-order singularity for thicker films where the square-ice mapping does not hold (inset of Fig. 2).

Substrate and strain effects.—Since films are grown on a substrate, the surfaces, and their respective orphan-bond couplings δ_o^{bot} , δ_o^{top} , will be asymmetric unless the top surface is capped with the substrate material. Surfaces with different couplings will order at different temperatures and one could have frustrated ordering for $\delta_o^{\text{top}} < \delta_o^c < \delta_o^{\text{bot}}$ (or vice versa). In particular, one might expect a regime of parameters where the charge order on one surface hinders the ordering of the entire film at lower temperature.

If the lattice parameters of the substrate are incommensurate with those of the spin ice film, lattice distortions will invariably be induced on the surface which could penetrate into the film, inducing a "bond distortion field." In experiments on $Dy_2Ti_2O_7$ films [22] with surfaces normal to the [110] axis, such a substrate-induced strain has been suggested as a possible mechanism for the loss of residual Pauling entropy [22] compared to bulk samples. We also note that for films of ionic crystals, termination with a polar surface often leads to a "polar catastrophe" with structural repercussions over the *entire* film [58].

We introduce bond distortions through δ_d in Eq. (1) which is applied uniformly throughout the sample. For $\delta_d > 0$, the distortion enhances the degeneracy lifting of the Coulomb phase driven by the dipolar interactions [44,45], such that the ordering occurs at a higher temperature than in the bulk (see results for $\delta_d = +0.2$ K in Fig. 4). As for negative δ_d , it induces a topological "KDP" transition (KDP stands for Potassium Di-hydrogen phosphate) whose ground state is ferromagnetic [59-61]. The description of the diverse ground states resulting from the competition with dipolar interactions, as a function of δ_d and L_h , is beyond the scope of this Letter. Nevertheless, in a first theoretical step towards understanding the loss of Pauling entropy observed in experiments [22], we simulated a thin film with $L_h = 5$ corresponding to the thinnest sample of Ref. [22] (Fig. 4). Indeed, for weak distortion $-0.2 \text{ K} \lesssim \delta_d < 0$, we find that the entropy dips smoothly below the Coulomb phase plateau as the temperature is reduced, without any phase transition down to 0.4 K. Our model is thus in qualitative agreement with the results of Ref. [22], even if it fails to reproduce the precise shape of the specific heat, whose Schottky peak is broader in experiments. The difference could be due to the difference of surface orientation, a strain gradient in the film, or the possible presence of disorder, common in rare-earth pyrochlores [62,63]. Because the KDP transition is topological in nature, the addition of a dedicated directed worm algorithm might be necessary for thermalization down to the lowest temperatures [61,64]. The absence of such global coherent updates, which is likely to be the case



FIG. 4. Strain effects smoothly lifts the Pauling entropy for $-0.2 \text{ K} \leq \delta_d < 0$ down to 0.4 K, while promoting bulk-DSI order for $\delta_d > 0$ (e.g. $\delta_d = +0.2$ K). Here $\delta_o = 0$. Thickness: $L_h = 5$ (5 nm) as in the thinnest sample of Ref. [22].

in real experiments, makes the study of out-of-equilibrium effects in strained thin films a promising open problem.

Discussion.—In this work, we have provided a theoretical benchmark for spin ice thin films. A surface termination perpendicular to the [001] direction generates strains and surface effects captured by orphan bonds. The latter plays the role of a chemical potential for emergent magnetic monopoles on the surfaces. We observe surface ordering, driven by long-ranged dipolar interactions, resulting in the crystalization of the magnetic surface charges. Over a finite temperature range, this surface ordering does not penetrate into the bulk, providing a magnetic example of partial wetting (see Fig. 2). For ultrathin films $(L_h = 1)$, the surface charge order stabilizes an effective square ice phase in the intervening layers of spins. This offers a solid-state route of this exotic phase, recently reported in experiments on confined water ice [65] and artificial lattices designed by nanolithography [66-68]. Last but not least, our simulations also confirm that strains coming from interface mismatch can lead to a smooth loss of Pauling entropy in Dy₂Ti₂O₇ films, as proposed in Ref. [22]. Our model thus offers a useful first step and guide for future ab initio calculations [27] aimed at developing a quantitative understanding of thin film experiments.

It is interesting to compare the partial wetting observed here to that occurring in water ice. Ice crystals show a thin surface layer stabilized down to -40 °C with remarkably high charge mobility [69–72]. Although often described as a liquid layer, its conductivity is orders of magnitude *larger* than that of bulk liquid water. This is admittedly a vastly more complex problem than our model spin ice [73–75]. However, on cleaving a surface from an ice crystal, one might expect surface charges to be induced in an analogous manner. While the putative charge crystal could then melt under the high Coulombic pressure, it has also been proposed that the high surface conductivity could come through a super-ionic mechanism in an *ordered* ionic array [70], reminiscent of the surface ordering we found here.

In conclusion, we believe the theory presented here is just the tip of the iceberg of possibilities for spin ice films and surfaces of single crystals. We hope our work motivates further efforts in the investigation of surface and confinement phenomena in frustrated magnetism and other strongly correlated systems described by an emergent gauge theory.

We thank Bruce Gaulin, Gabriele Sala, Etienne Lantagne-Hurtubise, Patrick Fournier, Rob Kiefl, Andrew MacFarlane, Jeff Rau, Nic Shannon, George Sawatzky, and Tommaso Roscilde for useful discussions. We acknowledge the hospitality of the Ecole Normale Supérieure de Lyon, Université Lyon 1 and CNRS (L. D. C. J. and M. J. P. G.) and of the Okinawa Institute of Science and Technology Graduate University (P. C. W. H. and M. J. P. G.). L. D. C. J. is supported by the Okinawa Institute of Science and Technology Graduate University.

The work at the University of Waterloo was supported by the NSERC of Canada, the Canada Research Chair program (M. J. P. G., Tier 1) and by the Perimeter Institute (PI) for Theoretical Physics. Research at the Perimeter Institute is supported by the Government of Canada through Innovation, Science and Economic Development Canada and by the Province of Ontario through the Ministry of Research, Innovation and Science.

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