Mean-field phase diagram and spin glass phase of
the dipolar Kagome Ising antiferromagnet

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We derive the equilibrium phase diagram of the classical dipolar Ising antiferromagnet at the
mean-field level on a geometry that mimics the two dimensional Kagome lattice. Our mean-field
method is based on the combination of the cluster variational Bethe-Peierls formalism and the
cavity method, developed in the context of the glass transition, and is complementary to the Monte
Carlo simulations realized in [Phys. Rev. B 98, 144439 (2018)]. Our results confirm the nature of
the low temperature crystalline phase which is reached through a weakly first-order phase transition.
Moreover, they allow us to interpret the dynamical slowing down observed in the work of Hamp &
al. as a remnant of a spin glass transition taking place at the mean-field level (and expected to be
avoided in 2 dimensions).

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I. INTRODUCTION

Many interesting classes of classical and quantum magnetic systems are extremely constrained. Hard local con-
straints lead to frustration and to the impossibility of satisfying all competing interactions simultaneously [1],
giving rise to the existence of highly degenerate ground states [2, 3]. Under certain conditions, these features produce a rich variety of collective behaviors [2, 3], unconventional phase transitions [4, 5], the emergence of a Coulomb phase with long-range correlations [9, 7], and other remarkably unusual and exotic phenomena.

On the other hand, frustration is also one of the key properties of glassy systems [8,13], where it also arises from the fact that minimizing some local interactions leads to the impossibility of minimizing other ones [1]. This feature can generate rugged energy landscapes and slow dynamics even in the absence of disorder [14,31]. It is therefore surprising at first sight that very little is known on glassy phases in geometrically frustrated magnetic systems. One of the first tentative investigations on this subject has been performed in Ref. [32], where glassy behavior was observed in nonrandomly frustrated Ising models with competing interactions. More recently, strong nonequilibrium effects, slow dynamics, and super-Arrhenius relaxation have also been reported in two-dimensional spin systems with competing long-range and short-range interactions [33,41].

On a different front, a thermodynamic theory, called the “frustration-limited domain theory” of the properties of supercooled liquids, and of the extraordinary increase of their characteristic structural relaxation times as the temperature is lowered, was formulated in terms of the postulated existence of a narrowly avoided thermodynamic phase transition due to geometric frustration [34] (see Ref. [19] for a review). In this context frustration describes an incompatibility between extension of the locally preferred order in a liquid and tiling of the whole space. This picture is consistent with appropriate minimal statistical mechanical models, such as three-dimensional Ising Coulomb frustrated lattice models, which display a slowing down of the relaxation in Monte Carlo simulations [15,35] and an ideal glass transition within mean-field approximations [18,20]. However, numerical simulations of these models in 3d are limited by the presence of a first-order transition to a modulated, defect-ordered phase [18], and cannot be performed at sufficiently low temperatures.

Several frustrated spin (or Potts) lattice models without quenched disorder have also been introduced and studied over the past years to describe the key features of the glass transition. However, most of them are either mean-field in nature (and cannot be easily generalized to finite dimensions) [11,16], or are characterized by (unphysical) multi-body interactions [31]. The classical three-coloring model on the two-dimensional hexagonal lattice have been shown to undergo a dynamical freezing in metastable states very similar to the one observed in structural glasses [37,38]. Slow dynamics also appears in electronic Coulomb liquids on the triangular lattice at quarter-filling [39], as well as in spin-ice systems both in 2d [40,41] and in 3d [42]. On the quantum side, it was shown in Ref. [43] that a valence bond glass phase emerges in the SU(N) Hubbard-Heisenberg model on a Bethe lattice in the large-N limit due to the interplay of
strong magnetic frustration and quantum fluctuations.

Yet, despite all these efforts over the past years, a clear and coherent picture of the glassy behavior that can arise due to the effect of geometric frustration in finite dimensional magnetic systems at low temperature is still missing.

Recently Hamp & al. [44] studied an Ising model on the Kagome lattice with short range antiferromagnetic interactions and dipolar interactions decaying as $1/r^3$—the Dipolar Kagome Ising Antiferromagnet (DKIAFM) introduced in [45]. By means of extensive Monte Carlo simulations the authors first showed evidence for a first-order transition from the high temperature paramagnetic phase to a low temperature crystal state that breaks time-reversal and sublattice symmetries, and coincides with the one previously proposed in Ref. [45] as the ground state. Furthermore, upon cooling below the first-order transition, the system enters a supercooled liquid regime which exhibits all the characteristic features of fragile glasses: two-time autocorrelation functions decay as stretched exponentials and the relaxation time grows in a super-Arrhenius fashion as the temperature is decreased. However, these conclusions were drawn out of numerical simulations of relatively small systems (about 300 spins) and might be affected by both strong finite-size effects and the difficulty of reaching thermal equilibrium in a reliable fashion due to strong metastability effects. Moreover, a consistent picture of the physical origin of the dynamical slowing down at low temperatures has not been convincingly established yet.

In order to overcome, at least partially, these issues, in this paper we perform an analytical study of the equilibrium phase diagram of the DKIAFM in the thermodynamic limit at a mean-field level, focusing both on the ordered state and the glassy phase. Our results essentially confirm, support, and elucidate the observations reported in Ref. [45]. Upon decreasing temperature, we first find a transition to a six-fold degenerate crystal state which breaks time reversal and rotation sublattice symmetry as the one observed in [44] [45]. The mean-field analysis indicates that the transition is indeed discontinuous. However, its first-order nature turns out to be extremely weak: the spinodal point of the crystal phase is very close to the transition point, resulting in a very large jump of the specific heat at the transition. This feature provides a possible explanation of the fact that the finite-size scaling of the numerical data of the maximum of the specific heat with the system size performed in Ref. [44] did not find the usual behavior ($C_{\text{max}} \propto N$) expected at a first-order transition due to very large finite-size effects.

When the system is supercooled below the first-order transition, we find that the paramagnetic state becomes unstable below a temperature at which the spin glass susceptibility diverges. Here, a continuous spin glass transition takes place at the mean-field level [46].

Note that the fact that the model displays a continuous spin glass transition in mean-field instead of a Random First-Order Transition [47] [48] of the kind found in structural glasses (such as hard spheres in infinite dimensions [27] and lattice glass models on the Bethe lattice [21] [22] [24] [26]) is perhaps not surprising. In fact, Ising spins with antiferromagnetic couplings on high-dimensional frustrated lattices and other related frustrated mean-field models with pairwise interactions are known to undergo a continuous transition to a spin glass phase when the temperature is lowered below the critical temperature of the antiferromagnetic phase [20] [49].

Beyond the fact that both spin glasses and structural glasses exhibit a pronounced slowing down of the dynamics upon cooling and aging in the low temperature phase, several important qualitative and quantitative differences characterize the dynamical behavior of these systems: In structural glasses two-time autocorrelation functions generically exhibit a two-step relaxation, characterized by a relatively fast decay to a plateau (i.e., the Edwards-Anderson order parameter) which appears discontinuously upon lowering the temperature, followed by a much slower decay, described by a stretched exponential. Moreover, the structural relaxation time is found to grow extremely fast, in a super-Arrhenius fashion, as the temperature is decreased [8] [13]. Conversely, in spin glasses the Edwards-Anderson order parameter is continuous at the transition and vanishes in the paramagnetic phase. Hence two-time autocorrelation functions should display a simple exponential decay when the transition is approached from the high temperature phase, and an algebraic decay at the critical point. Furthermore, the relaxation time is expected to diverge (only) as a power-law at the critical point (and to stay infinite in the whole low temperature phase) [49]. Nonetheless these differences are not clearly visible in numerical simulation of relatively small samples. A clear example of that is provided by the analysis of the dynamics of 3d Ising spin glasses performed in Ref. [50] [51] using Monte Carlo simulations of systems with up to 64$^3$ spins. The two-time autocorrelation function was found to be very well fitted by stretched exponentials, with an exponent $\beta$ that exhibits a temperature dependence extremely similar to the one reported in [44] for the DKIAFM. Moreover, although the divergence of the relaxation time as a power law, $\tau \sim (T - T_c)^{-\nu}$, is consistent with the numerics, a Vogel-Fulcher law, $\tau \sim e^{E_0/(T-T_c)}$, was also found to account reasonably well for the data.

The lower-critical dimension of the spin glass transition is expected to be $d_L \approx 2.5$ [52] (at least in the case of short-range interactions). Hence on general grounds we do not expect a genuine spin glass phase for the DKIAFM in 2$d$. Yet, the manifestations of the vestige of the transition can be very strong also in two dimensional systems: The spin glass amorphous order can establish over very long (although not infinite) length scales, the spin glass susceptibility can become very large (although not infinite), and the relaxation time can grow very fast at low temperature. Several experimental realizations of two-dimensional spin glasses using thin films do indeed show the same behavior as 3$d$ spin glasses at sufficiently
low temperature \cite{53,55}. In this sense, the existence of the spin glass phase in higher dimension, accompanied by the growth of long-range amorphous order and a rough free-energy landscape, provides a possible and natural explanation of the slow dynamics observed in the numerical simulations of Ref. \cite{44} of the 2d model at low temperatures.

Despite the fact that our mean-field approach consists in studying the model on a random sparse graph of triangular Kagome plaquettes, and cutting-off the dipolar interactions beyond the second nearest-neighbour plaquettes (i.e., the 5th nearest-neighbor spins), it provides a remarkably good approximation for the equilibrium properties of the 2d DKIAFM. For instance, the mean-field approach yields a zero temperature entropy density of the nearest-neighbor Kagome spin ice model obtained for $D = 0$ \cite{56} equal to $s_{GS} \approx 0.75204$, which turns out to be extremely close to the Pauling estimate $s_{GS} \approx 0.75225$. Similarly, the ground state energy density of the crystal ground state within the mean-field approximation is $e_{GS} \approx -1.6116$, which accounts reasonably well for the one found in the Monte Carlo simulations of systems with 300 spins, $e_{GS} \approx -1.515 \ 57$. As expected, the transition temperature to the crystalline phase is overestimated (by about a factor 3) by the mean-field treatment. Yet, the temperature dependence of the specific heat, the energy density, and the magnetization are remarkably similar, also at a quantitative level, to the ones found with Monte Carlo simulations (see Fig. 3).

The results presented here can serve at least two purposes: (i) They help to support, understand, and clarify the numerical results of Ref. \cite{44}. (ii) They provide a first step to bridge the gap between the slow dynamics observed in geometrically frustrated magnetic systems and the theory of the glass transition formulated in terms of rough free-energy landscapes. We believe that this analysis is of particular interest, especially in the light of the experimental relevance of the model, which could serve in geometrically frustrated magnetic systems and as a step to bridge the gap between the slow dynamics observed in the numerical results of Ref. \cite{44}.

In the following we will be interested in the case in which both interactions are antiferromagnetic, i.e., $J > 0$ and $D > 0$. The case $D = 0$ is known to be fully frustrated and does not order down to zero temperature \cite{67}.

The phase diagram of the $J = 0$ model is less well understood but the system is again strongly frustrated with any ordering (if present at all) suppressed down to temperatures $T < D \ 69$.

The previous studies of the model \cite{44,45} considered the coupling parameters $D = 1 \text{K}$ and $J = 0.5 \text{K}$ (setting $k_B = 1$ and measuring all energies in Kelvin). A further advantage of developing an analytic (although approximate) treatment is that it is relatively simple to explore the parameter space. Without loss of generality we set $J = 0.5 \text{K}$ throughout (as in \cite{44,45}) and study the phase diagram of the model and the constitutive equations in the different phases varying the dipolar coupling $D$ and the temperature $T$.

III. THE MEAN-FIELD ANALYSIS

Our mean-field treatment is based on the combination of the cluster variational Bethe-Peierls formalism (already successfully employed in the study of the equilibrium properties of geometrically frustrated magnetic systems \cite{4,45,71}) and the cavity method \cite{49}, developed in the context of glassy and disordered systems described by replica symmetry breaking (RSB). The latter concept is related to a complex free-energy landscape with special structure and the calculational meaning of it, in the context of the cavity method, will become clear below. In particular, we define the model on a Random Regular Graph \cite{72} (RRG) of $N_\Delta$ triangular plaquettes of total coordination three (see Fig. 1 for a sketch). In this case, the number of spins is equal to $N = 3N_\Delta/2$ since since each spin belongs to two plaquettes and each plaquette contains three spins. RRGs are a special class of sparse graphs, whose elements are chosen at random with uniform probability over the ensembles of all graphs of $N_\Delta$ nodes, such that each node (i.e., a triangular plaquette) has exactly three neighbors. RRGs have a local tree-like structure, which allows one to obtain exact self-consistent recursion relations for the probability distributions of the spin configurations on each plaquette of the graph. Yet, they have large loops, whose typical

II. THE MODEL

We consider the DKIAFM \cite{44,45} in which $N$ classical spins $S_i = \pm 1$ are placed on the vertices of a two-dimensional Kagome lattice and point in a direction perpendicular to the plane. The Hamiltonian comprises an antiferromagnetic exchange term of strength $J$ between spins at nearest-neighbor lattice sites $(ij)$ and long-range dipolar interactions of characteristic strength $D$ between all pairs of spins:

$$
\mathcal{H} = J \sum_{(ij)} S_i S_j + \frac{D}{2} \sum_{ij,j} S_i S_j \frac{r_{ij}}{r_{ij}^3},
$$

where the distance $r_{ij} = |\mathbf{r}_i - \mathbf{r}_j|/a$ between the spins $i$ and $j$ is measured in units of the lattice spacing $a$ (that we set equal to 1 throughout).

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length scales as \( \log N_\Delta \) and diverges in the thermodynamic limit. Hence, a RRG is locally a tree, but it is frustrated, does not have a boundary, and is statistically translational invariant. For these reasons RRGs are suitable lattices to study the thermodynamics of glassy and disordered systems at a mean-field level [21, 22, 24, 25]. However, in order to be tractable, the cavity approach is formulated for systems with finite range interactions. Hence, before proceeding further we need to treat the dipolar interactions of Eq. (1) in an approximate fashion. In practice, in the analytic calculations described below we choose to cut-off the dipolar couplings up to second nearest-neighboring plaquettes (i.e., the interactions between the spins belonging to the red plaquette \( \alpha \) of Fig. 1 and the spins belonging to green plaquettes are set to zero).

Consider now the cavity triangle \( \alpha \) (red) in absence of one of its neighboring plaquettes \( \beta \) (dashed black). We define \( p_{\alpha \rightarrow \beta}({\{S_\alpha\}}|{\{S_\beta\}}) \) as the probability to observe the spin configuration \( {S_\alpha} \equiv \{S_{\alpha,1}, S_{\alpha,2}, S_{\alpha,3}\} \) on the cavity triangle \( \alpha \) of the (rooted) RRG, given that the spin configuration of the plaquette \( \beta \) is \( {S_\beta} \equiv \{S_{\beta,1}, S_{\beta,2}, S_{\beta,3}\} \). We have adopted the convention that spin 1 is the root of the cavity plaquette and spins 2 and 3 are labeled anticlockwise. Using this convention one has that \( S_{\beta,1} \equiv S_{\alpha,1} \).

The probabilities \( p_{\alpha \rightarrow \beta}({\{S_\alpha\}}|{\{S_\beta\}}) \) can be written in terms of the marginal probabilities defined on the cavity triangles \( \gamma \) and \( \delta \) in absence of the triangle \( \alpha \), times the Gibbs’ weight associated to each spin configuration:

\[
p_{\alpha \rightarrow \beta}({\{S_\alpha\}}|{\{S_\beta\}}) = \left( Z^{(\text{iter})}_{\alpha \rightarrow \beta} \right)^{-1} \sum_{\{|\gamma,\delta\}\}} p_{\gamma \rightarrow \alpha}({\{S_\gamma\}}|{\{S_\alpha\}}) p_{\delta \rightarrow \alpha}({\{S_\delta\}}|{\{S_\alpha\}}) e^{-\beta \hat{H}_{\alpha \rightarrow \beta}({\{S_\alpha,S_\gamma,S_\delta\}}|{\{S_\beta\}})}, \tag{2}
\]

where \( Z^{(\text{iter})}_{\alpha \rightarrow \beta} \) is a normalization factor ensuring that \( \sum_{\{|\gamma,\delta\}\}} p_{\gamma \rightarrow \alpha}({\{S_\gamma\}}|{\{S_\alpha\}}) p_{\delta \rightarrow \alpha}({\{S_\delta\}}|{\{S_\alpha\}}) = 1 \) and is associated to the “free-energy shift” involved in the iteration process: \( -\beta \Delta F^{(\text{iter})}_{\alpha \rightarrow \beta} \equiv \log Z^{(\text{iter})}_{\alpha \rightarrow \beta} \). Here we introduce the notation \( \sum_{\{|\gamma,\delta\}\}} \) that indicates the sum over all possible configurations \( \{S_\gamma\} \) and \( \{S_\delta\} \) of the spin degrees of freedom of the plaquettes \( \gamma \) and \( \delta \), compatible with the constraints imposed by the spin configuration \( \{S_\alpha\} \) on the plaquette \( \alpha \), i.e., \( S_{\gamma,1} = S_{\alpha,2} \) and \( S_{\delta,1} = S_{\alpha,3} \) [see Eq. (10)]. We will use this notation throughout this section. The Hamiltonian \( \hat{H}_{\alpha \rightarrow \beta}({\{S_\alpha,S_\gamma,S_\delta\}}|{\{S_\beta\}}) \)
appearing in the Gibbs factor of Eq. 2 is a modified Hamiltonian, Eq. 1, restricted to the cavity plaquette $\alpha \rightarrow \beta$: 

$$\tilde{H}_{\alpha \rightarrow \beta}(\{\alpha, S_\gamma, S_\delta\}) = \tilde{H}_{\alpha}^{(AF)} + \tilde{H}_{\alpha}^{(2D)} + \tilde{H}_{\alpha}^{(3D)} + \tilde{H}_{\alpha}^{(2NND)} + \tilde{H}_{\alpha}^{(2NND)},$$

where 

$$\tilde{H}_{\alpha}^{(AF)} = (J + D) [S_{\alpha,1}S_{\alpha,2} + S_{\alpha,1}S_{\alpha,3} + S_{\alpha,2}S_{\alpha,3}],$$

$$\tilde{H}_{\alpha}^{(2D)} = D \left[ \frac{S_{\alpha,1}S_{\alpha,2} + S_{\alpha,3}S_{\gamma,3} + S_{\alpha,1}S_{\gamma,3} + S_{\alpha,2}S_{\gamma,2}}{8} \right],$$

$$\tilde{H}_{\alpha}^{(3D)} = D \left[ \frac{S_{\alpha,1}S_{\alpha,3} + S_{\alpha,2}S_{\delta,2} + S_{\alpha,1}S_{\delta,2} + S_{\alpha,2}S_{\delta,3}}{8} \right],$$

$$\tilde{H}_{\alpha}^{(2NND)} = D \left[ \frac{S_{\gamma,2}S_{\delta,2} + S_{\gamma,3}S_{\beta,3}}{\sqrt{7}} + \frac{S_{\gamma,2}S_{\delta,3}S_{\beta,3}}{27} \right].$$

The meaning of this decomposition is the following. $\tilde{H}_{\alpha}^{(2D)}$ contains the 4 dipolar interaction terms between the spins belonging to the plaquette $\alpha$ and the spin belonging to its nearest-neighbor plaquette $\gamma$ attached to the spin $S_{\alpha,2}$, which are not already contained in $\tilde{H}_{\alpha}^{(AF)}$. $\tilde{H}_{\alpha}^{(3D)}$ contains the 4 dipolar interaction terms between the spins of the second nearest-neighbor plaquettes $\gamma$ and $\delta$ which are not already contained in $\tilde{H}_{\alpha}^{(2D)}$ and $\tilde{H}_{\alpha}^{(3D)}$.

In order to obtain the marginal probabilities of the spin configurations on each plaquette of the (unrooted) RRG (where each triangular plaquette has exactly three neighbors), one needs to merge three cavity plaquettes (e.g., plaquettes $\beta$, $\gamma$, and $\delta$ of Fig. 1) onto their neighboring plaquette (e.g., plaquette $\alpha$ of Fig. 1). In this way one obtains:

$$P_\alpha(\{\alpha\}) = \left( Z_\alpha^{(s)} \right)^{-1} \sum_{[\beta,\gamma,\delta]} p_{\beta \rightarrow \alpha}(\{\beta\})p_{\gamma \rightarrow \alpha}(\{\gamma\})p_{\delta \rightarrow \alpha}(\{\delta\})e^{-\beta \tilde{H}_\alpha(\{\alpha, S_\beta, S_\gamma, S_\delta\})},$$

where $Z_\alpha^{(s)}$ is a normalization factor ensuring that $\sum_\alpha P_\alpha(\{\alpha\}) = 1$, and is associated to the “free-energy shift” involved in the process of joining three cavity plaquettes ($\beta$, $\gamma$, and $\delta$) to a central plaquette ($\alpha$): $-\beta \Delta F^{(s)}_\alpha \equiv \log Z_\alpha^{(s)}$.

The plaquette Hamiltonian $\tilde{H}_\alpha(\{\alpha, S_\beta, S_\gamma, S_\delta\})$ reads

$$\tilde{H}_\alpha(\{\alpha, S_\beta, S_\gamma, S_\delta\}) = \tilde{H}_\alpha^{(AF)} + \tilde{H}_\alpha^{(1D)} + \tilde{H}_\alpha^{(2D)} + \tilde{H}_\alpha^{(3D)} + \tilde{H}_\alpha^{(2NND)} + \tilde{H}_\alpha^{(2NND)},$$

and the other terms are given in Eq. 3.

The equilibrium averages of all local observables which involve the spin degrees of freedom of a given plaquette, including, e.g., the magnetization, can be expressed in terms of these marginal probabilities:

$$\langle O_\alpha \rangle = \sum_{\{S_\alpha\}} O(\{S_\alpha\})P_\alpha(\{S_\alpha\}).$$

Similarly, the contribution to the average energy due to the plaquette $\alpha$ can be expressed as

$$\langle e_\alpha^{(s)} \rangle = \frac{\sum_{[\alpha \rightarrow (\beta, \gamma, \delta)]} p_{\beta \rightarrow \alpha}(\{\beta\})p_{\gamma \rightarrow \alpha}(\{\gamma\})p_{\delta \rightarrow \alpha}(\{\delta\})\tilde{H}_\alpha(\{\alpha, S_\beta, S_\gamma, S_\delta\})e^{-\beta \tilde{H}_\alpha(\{\alpha, S_\beta, S_\gamma, S_\delta\})}}{\sum_{[\alpha \rightarrow (\beta, \gamma, \delta)]} p_{\beta \rightarrow \alpha}(\{\beta\})p_{\gamma \rightarrow \alpha}(\{\gamma\})p_{\delta \rightarrow \alpha}(\{\delta\})e^{-\beta \tilde{H}_\alpha(\{\alpha, S_\beta, S_\gamma, S_\delta\})}}.$$}

The process of joining two neighboring cavity plaquettes (e.g., plaquettes $\alpha$ and $\beta$ of Fig. 1) involves another “free-energy shift”, defined as

$$e^{-\beta \Delta F^{(l)}_{\alpha \rightarrow \beta}} \equiv Z_{\alpha \rightarrow \beta}^{(l)} = \sum_{[\alpha \rightarrow \beta]} p_{\alpha \rightarrow \beta}(\{\alpha\})p_{\beta \rightarrow \alpha}(\{\beta\})e^{-\beta \tilde{H}_{\beta \rightarrow \alpha}^{(l)}},$$

where the Gibbs’ factor $\tilde{H}_{\beta \rightarrow \alpha}^{(l)}$ has been defined in Eq. 5. Similarly, the contribution to the average energy coming from the interactions between two neighboring cavity plaquettes is given by

$$\langle e^{(l)}_{\alpha \rightarrow \beta} \rangle = \frac{\sum_{[\alpha \rightarrow \beta]} p_{\alpha \rightarrow \beta}(\{\alpha\})p_{\beta \rightarrow \alpha}(\{\beta\})\tilde{H}_{\beta \rightarrow \alpha}^{(l)}e^{-\beta \tilde{H}_{\beta \rightarrow \alpha}^{(l)}}}{\sum_{[\alpha \rightarrow \beta]} p_{\alpha \rightarrow \beta}(\{\alpha\})p_{\beta \rightarrow \alpha}(\{\beta\})e^{-\beta \tilde{H}_{\beta \rightarrow \alpha}^{(l)}}}.$$
We recall here the convention adopted for the notation of the summation over the spin degrees of freedom in the expressions above:

\[
\sum_{\|[\gamma, \delta]||_\alpha} = \sum_{\{S_\gamma, S_\delta\}}^{S_{\gamma,1}=S_{\gamma,2}}_{S_{\delta,1}=S_{\delta,2}} \frac{1}{2} \sum_{\{S_\gamma, S_\delta\}}^{S_{\gamma,1}=S_{\gamma,2}}_{S_{\delta,1}=S_{\delta,2}}
\]

The free-energy of the system can be obtained by combining the free-energy shifts involved in the different processes, as explained in [24, 49]:

\[
F = \sum_{\alpha=1}^{N_\Delta} \Delta F_{\alpha}^{(s)} - \sum_{\langle\alpha, \beta\rangle} \Delta F_{\alpha \leftrightarrow \beta}^{(l)} = \frac{1}{2} \sum_{\langle\alpha, \beta\rangle} \left( \Delta F_{\alpha \rightarrow \beta}^{(\text{iter})} + \Delta F_{\beta \rightarrow \alpha}^{(\text{iter})} \right) - \frac{1}{2} \sum_{\alpha=1}^{N_\Delta} \Delta F_{\alpha}^{(s)}
\]

where \(\langle\alpha, \beta\rangle\) denotes the sum over the \(3N_\Delta/2\) nearest-neighbors plaquettes on the graph (the last equality simply comes from the fact that \(\Delta F_{\alpha}^{(s)} = \Delta F_{\alpha \rightarrow \beta}^{(\text{iter})} + \Delta F_{\beta \rightarrow \alpha}^{(\text{iter})}\) by construction). The average entropy of the system is then given by \(\langle S \rangle = \beta(\langle E \rangle - F)\). Analogously, the total average energy can be written as

\[
\langle E \rangle = \sum_{\alpha=1}^{N_\Delta} \langle e_{\alpha}^{(s)} \rangle - \sum_{\langle\alpha, \beta\rangle} \langle e_{\alpha \leftrightarrow \beta}^{(l)} \rangle.
\]

Equations [2] can be written for arbitrary (large) RRGs and are expected to become exact in the thermodynamic limit. On each triangle of the RRG one can define three cavity plaquettes by removing one of its three neighbors. Thus, Eqs. (2) represent a set of \(32 \times 3 \times N_\Delta\) coupled nonlinear algebraic equations for the 32 marginal probabilities \(p_{\alpha \rightarrow \beta}(\{S_\alpha\}|\{S_\beta\})\) associated to the 32 possible configurations of the spins \(\{S_\alpha\}\) on the cavity plaquette \(\alpha\), given the configuration of the spins \(\{S_\beta\}\). Once the fixed points of these equations are found, one can compute the marginal probabilities on each plaquette of the graph from Eq. (4), along with the free-energy and all observables. In the following, we will discuss three specific solutions of the equations in the thermodynamic limit, corresponding to the (RS) homogeneous paramagnet, the (RS) ordered crystalline state, and the (RSB) glassy phase.

### B. The paramagnetic phase

The paramagnetic phase is characterized by translational invariance and corresponds to the homogeneous and RS solution of the recursion relations:

\[
p_{\alpha \rightarrow \beta}(\{S_\alpha\}|\{S_\beta\}) = p_{\text{para}}(\{S_\alpha\}|\{S_\beta\}) \quad \forall \alpha, \beta.
\]

The probabilities \(p_{\text{para}}(\{S_\alpha\}|\{S_\beta\})\) are given by the fixed point of Eqs. (2), which in this limit become a simple system of 32 coupled nonlinear algebraic equations. The free-energy, the energy, and the magnetization (which is identically zero by \(Z\) inversion symmetry in the paramagnetic phase, which implies that \(p_{\text{para}}(\{S_\alpha\}|\{S_\beta\}) = p_{\text{para}}(-\{S_\alpha\}|\{-S_\beta\})\), can be easily computed from Eqs. (4). This phase is expected to be stable at high temperature. However, the average entropy density \(\langle s \rangle = \beta(\langle e \rangle - f)\) becomes negative below a certain temperature, \(T_{\text{para}}(D)\). This indicates that the homogeneous solution is certainly not appropriate to describe the low temperature region of the phase diagram.

### C. The stability of the paramagnetic phase

The manifestation of the failure of the RS solution also shows up via a loss of stability of the RS fixed point, as given by a simple linear analysis. To describe this instability one needs to introduce a probability distribution \(P[p]\), where \(p\) is a short-hand notation for the 32 marginal probabilities \(p_{\alpha \rightarrow \beta}(\{S_\alpha\}|\{S_\beta\})\) and \(P[p]\) is defined as the probability density that the probabilities \(p_{\alpha \rightarrow \beta}(\{S_\alpha\}|\{S_\beta\})\) on the cavity plaquette \(\alpha\) are equal to \(p(\{S_\alpha\}|\{S_\beta\})\).

In the homogeneous phase from Eq. (2) one has that the probability distributions of the marginal probabilities on the triangular plaquettes must satisfy the following
self-consistent equation:
\[ \mathcal{P}[\vec{p}] = \int d\vec{p}_{\vec{r}} \mathcal{P}[\vec{p}_{\vec{r}}] \delta[\vec{p}_{\vec{r}} - \vec{p}(\vec{p}_{\vec{r}}, \vec{p}_{\vec{b}})], \]
where \( \vec{p}(\vec{p}_{\vec{r}}, \vec{p}_{\vec{b}}) \) is a short-hand notation for the r.h.s. term of the recursion relations \[2\]. Close to the homogeneous paramagnetic solution we have, to first order, \( p(\{S_a\}|\{S_b\}) \approx p_{\text{para}}(\{S_a\}|\{S_b\}) + \delta p(\{S_a\}|\{S_b\}) \).
Starting with \( \delta \vec{p} \) identically and independently distributed and injecting the expression above into Eq. \[12\], one has that the deviation of the marginal probabilities from the homogeneous solution evolves under iteration as
\[ \langle \delta \vec{p} \rangle = 2 \frac{\partial \vec{p}(\vec{p}_{\vec{r}}, \vec{p}_{\vec{b}})}{\partial \vec{p}_{\vec{r}}} \bigg|_\text{para} \langle \delta \vec{p} \rangle, \]
where \( \langle \cdot \rangle \) refers to the average using the distribution \( \mathcal{P}(\vec{p}) \). \( \delta \vec{p}/\partial \vec{p}_{\vec{r}} \) is actually a 32 × 32 Jacobian matrix. If \( \lambda_{\text{max}} \) denotes the eigenvalue of largest modulus of that matrix, the stability criterion simply reads \( 2|\lambda_{\text{max}}| \leq 1 \). When \( 2|\lambda_{\text{max}}| > 1 \), the paramagnetic solution is instead unstable with respect to a “modulation” instability, corresponding to a transition to a regime with successive (homogeneous) generations of the tree carrying different values of the marginal probabilities. Such modulation instability is thus a manifestation of an instability toward an ordered phase, which breaks translational invariance.

This instability criterion can also be obtained by studying response functions to a perturbation (which is related to correlations through the fluctuation-dissipation theorem) \[24\]. In this setting the instability is detected by means of the divergence of the linear magnetic susceptibility in the paramagnetic phase, defined as:
\[ \chi = \frac{1}{N^2} \sum_{\alpha, \beta} \langle S_\alpha S_\beta \rangle_c = \frac{1}{N^2} \sum_{\alpha, \beta} \left( \frac{\partial \langle S_\alpha \rangle}{\partial h_\beta} \right)_{h_\gamma = 0}, \]
where \( S_\alpha = \sum_{i \in \alpha} S_i \) is a short-hand notation for the magnetization of the plaquette \( \alpha \) and \( h_\beta \) is an external magnetic field conjugated to the magnetization of the plaquette \( \beta \). Making use of the homogeneity of the paramagnetic solution and the tree-like structure of the lattice, the susceptibility can be rewritten as
\[ \chi = 1 + 3 \sum_{r=1}^{\infty} 2^{r-1} \langle S_{\alpha_0} S_{\alpha_r} \rangle_c, \]
where \( S_{\alpha_0} \) and \( S_{\alpha_r} \) are two plaquettes taken at distance \( r \) on the tree. The series converges provided that \( \lim_{r \to \infty} \log(\langle S_{\alpha_0} S_{\alpha_r} \rangle_c)/r < \log 2 \). To evaluate \( \langle S_\alpha S_\beta \rangle_c \), we invoke the fluctuation-dissipation relation:
\[ \langle S_{\alpha_0} S_{\alpha_r} \rangle_c = \left. \frac{\partial \langle S_{\alpha_0} \rangle}{\partial h_{\alpha_0}} \right|_{h_\gamma = 0}, \]
where \( h_{\alpha_0} \) denotes the external magnetic field conjugate to \( S_{\alpha_0} \). Since \( h_{\alpha_0} \) is a function of (the components of) \( P(\{S_{\alpha_0}\}) \), we can use the chain rule along the branch of the tree which connects the plaquette \( \alpha_0 \) with the plaquette \( \alpha_r \) through the plaquettes \( \alpha_l \), \( l = 1, \ldots, r - 1 \):
\[ \frac{\partial \langle S_{\alpha_r} \rangle}{\partial h_{\alpha_0}} = \left( \prod_{l=1}^{r} \frac{\partial \langle S_{\alpha_l} \rangle}{\partial h_{\alpha_{l-1}}} \right) \frac{\partial \langle S_{\alpha_r} \rangle}{\partial \vec{p}_{\alpha_r}} \frac{\partial \vec{p}_{\alpha_r}}{\partial h_{\alpha_0}}. \]
In the paramagnetic phase, all the intermediate marginal cavity probabilities are equal and the previous equation factorizes, leading again to \( 2|\lambda_{\text{max}}| \leq 1 \).

The maximal eigenvalue \( \lambda_{\text{max}} \) increases as the temperature is lowered and the linear susceptibility of the paramagnetic phase diverges at a certain temperature, signaling a modulation instability of the paramagnetic phase toward a crystalline phase (see Sec. \[III \]D) at a temperature \( T_{\text{mod}}(D) \).

One can also look for another kind of instability, namely a spin glass instability, which manifests itself as a divergence of the non-linear susceptibility \[23\], which is defined as
\[ \chi_{\text{sg}} = \frac{1}{N^2} \sum_{\alpha, \beta} \langle S_\alpha S_\beta \rangle_c^2. \]
Equivalently, this instability appears as a widening of the variance \( \langle (\delta \vec{p})^2 \rangle \) under the recursion of Eq. \[12\]. Both approaches lead to a stability criterion \( 2|\lambda_{\text{max}}| \leq 1 \). Note that this condition is always weaker than that for the modulation instability, \( 2|\lambda_{\text{max}}| \leq 1 \), associated to the crystalline order. However, it is the relevant one in the case of glassy phases, characterized by the establishment of long-range amorphous order.

Solving the recursion relations \[2\] in the paramagnetic phase, we find that the homogeneous solution becomes unstable below a temperature \( T_{\text{sg}}(D) \), at which the spin glass susceptibility diverges (with \( T_{\text{sg}}(D) < T_{\text{mod}}(D) \)).

This requires either a phase transition before the spin glass local instability is reached \[21\]–\[26\] (as occurs in the mean-field models of fragile glasses, described by a Random First-Order Transition \[17\]–\[18\], or a continuous (possibly spin glass) transition at \( T_{\text{sg}} \). We will show below that the latter scenario is the correct one for the DKIAFM. In order to do this in Sec. \[III \]E we look for a solution of the recursion relations which breaks the replica symmetry, corresponding to a glassy phase where many local minima of the free-energy exist and where the local marginal probabilities fluctuate from a plaquette to another.

### D. The crystal phase

One can look for a crystalline RS solution, where the marginal probabilities do not fluctuate from site to site, but are different in different sites (break-down of translational invariance). The (sixfold degenerate) crystalline
state proposed in [45] and observed numerically in [44] is characterized by a 12-spin unit cell and breaks (twofold) time-reversal symmetry and (threefold) rotation symmetry (see Refs. [44,45] for more details). In order to be able to account for such ordered phase we need to introduce 8 sublattices of triangular plaquettes (see Fig. 2), corresponding to 24 different sets of cavity plaquettes. The merging of the 24 cavity plaquettes is done taking into account the structure of the crystalline phase, as explained in the caption of Fig. 2. The recursion equations (2) become then a set of $24 \times 32$ coupled nonlinear algebraic equations for the marginal probabilities on the 24 cavity plaquettes on each sublattice. The solution of these equations appears discontinuously at a spinodal point $T_{sp}(D)$, and becomes thermodynamically stable when the corresponding free-energy crosses the paramagnetic one, at the melting temperature $T_{m}(D)$. At that temperature we observe a first-order phase transition characterized by a spontaneous breakdown of the translational, rotational, and spin inversion invariance, accompanied by a discontinuous jump of the energy density and of the entropy density. Decreasing further the temperature, the energy in the crystalline phase approaches quickly the ground state value, $e_{GS} \approx -1.6116$ (which turns out to be remarkably close to the one found with Monte Carlo simulations of systems of 300 spins, $e_{GS} \approx -1.515$ [44], and the entropy quickly approaches zero.

Inspecting the (ground state) spin configuration of Fig. 2 it was noticed in [44] that one of the three spins of the Kagome triangles are completely polarized (i.e., the bottom spins of sublattices $a,b,c,d$ and the top spins of sublattices $A,B,C,D$), with the state having zero magnetization overall. Note that the need to introduce 8 sublattices of triangular plaquettes is due to the fact that the spin pattern on the two non-polarized rows of spins of the Kagome triangles (i.e., along the horizontal bonds in Fig. 2) has period four, with three spins $S = \mp 1$ followed by one spin $S = \pm 1$. Based on these observations, suitable order parameters for the transition to the ordered state are the sublattice magnetizations:

$$m_X = \sum_{i \in X} S_i,$$

with $X$ denoting the 8 different sublattices: $X = \{A,B,C,D,a,b,c,d\}$. These order parameters essentially coincide with the emergent effective charge variables introduced in Ref. [44], derived from the so-called dumbbell picture [73]. The (ground state) spin configuration of Fig. 2 corresponds to $m_a = m_c = m_A = m_B = \pm 1$ and $m_b = m_d = m_C = \mp 1$. Equivalently, one can choose as order parameter $m_{pol} = (S_{pol})$, the average magnetization of the spins of the Kagome triangles that are completely polarized, as done in [44]. Following this suggestion, we will use $m_{pol}$ as the order parameter jumping from zero to a finite value at the transition.

We have also looked for other plausible competing ordered phases, which break the translational and rotational symmetries in different ways and have a different unit cells. However, such alternative crystalline states turn out to be less favourable (i.e., they have a higher free-energy) compared to the crystalline phase of [44,45]. Yet, if one does not include the dipolar interactions between the second nearest-neighboring plaquettes, the crystalline phase depicted in Fig. 2 disappears (i.e., no physically relevant fixed point of the recursion relations is found corresponding to the sublattice structure of Fig. 2), and another completely different (fourfold degenerate) ordered phase emerges. This observation highlights the importance of accounting for the dipolar interaction as accurately as possible, in order to recover the correct dep-
to Eq. (11), the 1RSB free-energy density functional is given by
\[ \phi(m) = \Delta \phi^{(s)}(m) - \frac{3}{2} \Delta \phi^{(l)}(m), \] (14)

with
\[ e^{-\beta m \Delta \phi^{(s)}} = \int d\mathcal{P}_m(\bar{p}\gamma) d\mathcal{P}_m(\bar{p}\beta) e^{-\beta m \Delta F^{\text{iter}}}, \]
\[ e^{-\beta m \Delta \phi^{(l)}} = \int d\mathcal{P}_m(\bar{p}\alpha) d\mathcal{P}_m(\bar{p}\beta) e^{-\beta m \Delta F^{\text{iter}}}, \]
where the free-energy shifts have been defined in Sec. IIIA. The other relevant thermodynamic observables, such as, e.g., the average energy, can be obtained in a similar fashion [24, 49]. The parameter \( m \) is fixed by the maximization of the free-energy functional with respect to it [24, 49], which allows to recover the complexity as a Legendre transform of \( \phi(m) \):
\[ m \phi(m) = m f - \frac{1}{\beta} \Sigma(f). \]

The RS high-temperature homogeneous description of the phase is recovered by taking \( \mathcal{P}_m(\bar{p}) = \delta(\bar{p} - \bar{p}_{\text{para}}) \) and \( m = 1 [75] \).

Since Eq. (13) is a functional relation, an analytical treatment is not possible in general. Yet the self-consistent equation can be efficiently solved numerically with arbitrary precision using a population dynamics algorithm (for all technical details see [49]). For high values of the temperature \( (T > T_{sg}(D)) \) we recover the paramagnetic solution. Lowering the temperature, a nontrivial solution of the 1RSB equation appears continuously exactly at \( T_{sg}(D) \). Right below \( T_{sg} \) the probability distribution \( \mathcal{P}_m(\bar{p}) \) acquires an infinitesimal widening of the variance \( \langle (\delta \bar{p})^2 \rangle \). This scenario corresponds to a continuous transition to a spin glass phase at the temperature at which the spin glass susceptibility diverges. The order parameter of the spin glass transition is the Edwards-Anderson order parameter, \( \eta_{EA} = (1/N) \sum_i (\langle S_i \rangle)^2 \), which vanishes linearly as \( \eta_{EA} \sim (T_{sg} - T) \) for \( T \rightarrow T_{sg} [49] \).

As it is known, the low-temperature spin glass phase should be described by full RSB [49]. However, any new level of RSB will require considering a more sophisticated situation, namely a distribution over the probability distribution of the previous level. For instance, the two-step RSB will be written as a distribution \( \mathcal{Q}[\mathcal{P}(\bar{p})] \) over distributions \( \mathcal{P}(\bar{p}) \). Describing with this formalism a finite connectivity system with full RSB is therefore too complicated, and we will limit ourselves to the 1RSB Ansatz. Moreover, since solving the self-consistent functional equation (13) via population dynamics is quite computationally demanding, we did not perform the maximization of the free-energy functional (13) with respect to \( m \). For these reasons, our approach only provides an approximate description of the equilibrium properties of the spin glass phase and we have not pushed the 1RSB calculations far below \( T_{sg} \) (essentially we only consider few values of the temperature in the vicinity of the critical point).
IV. PHASE DIAGRAM AND THERMODYNAMIC BEHAVIOR

In this Section we discuss the main results found within the mean-field treatment of the DKIAFM described in the previous sections. In order to compare with the numerical results of the Monte Carlo simulations of \cite{44}, we start by fixing the parameter $D$ to 1, as in Refs. \cite{44,45}, and measure several observables such as the average energy density $\langle e \rangle = \langle E \rangle / N_{\Delta}$, the (intensive) specific heat $c = \partial \langle e \rangle / \partial T$, the magnetization of the polarized spin in one of the sixfold degenerate ground state configurations $|m_{\text{pol}}\rangle$, and the average entropy density $\langle s \rangle = (S) / N_{\Delta}$, as a function of the temperature $T$ in the paramagnetic, crystal, and 1RSB glass solutions of the recursive cavity equations. The results are shown in Fig. 3. At high temperature the system is found in the paramagnetic phase. Upon lowering the temperature, a first-order transition to the crystalline phase proposed in Refs. \cite{44,45} (see Fig. 2) occurs at $T_m$. The order parameter $|m_{\text{pol}}\rangle$ presents a finite jump at $T_m$, where the average energy and entropy densities also display an abrupt decrease. The transition to the ordered state turns out to be very weakly first-order, in the sense that the spinodal point of the crystalline solution, $T_{sp} \approx 0.16$ K, is very close to the transition temperature $T_m \approx 0.1566$ K where the free-energies of the paramagnetic phase and the crystal phase cross. These temperatures are also numerically close to the modulation instability temperature $T_{\text{mod}} \approx 0.147$ K. Since the specific heat of the crystal solution diverges at the spinodal point, the vicinity of $T_m$ and $T_{sp}$ results in a very large jump (of about a factor 3) of the intensive specific heat at the transition. This feature might explain the deviations observed in the numerical simulations of the expected scaling of the peak of the (extensive) specific heat as $C_{\text{max}} \propto N_{\Delta}$ \cite{44}.

Although approximate, our approach accounts remarkably well for the numerical results of Ref. \cite{44}. As expected, the transition temperature is overestimated by the mean-field approximation (by about a factor 3). Yet, the temperature dependencies of the specific heat, the energy, and the magnetization are, also at a quantitative level, very similar to the ones found in Ref. \cite{44} (recall the energy, entropy, and specific heat per spin are obtained by multiplying the energy, entropy, and specific heat per plaquette by a factor 2/3).

The paramagnetic phase is metastable below $T_m$, corresponding to a supercooled regime. If one keeps lowering the temperature within the supercooled phase, the spin glass susceptibility grows and diverges at $T_{sg} \approx 0.1$ K, where a continuous transition to a spin glass phase takes place. Although the spin glass phase is presumably described by full RSB (at least at the mean-field level), our approach only allows one to perform an approximate 1RSB Ansatz for the low temperature glassy phase. Moreover, solving the self-consistent functional equation \cite{13} via population dynamics is computationally heavy. For these reasons, we did not push the calculations of the thermodynamic observables too deep into the spin glass phase, and only solved the equations for few points close to the critical temperature.

In Fig. 4 we plot the phase diagram of the DKIAFM, showing the position of the different phases when varying the temperature $T$ and the dipolar interaction $D$ ($J$ is fixed to $J = 0.5$ K). The effect of varying the dipolar interaction turns out to be particularly simple. In fact, we find that the phase boundaries, as well as all the characteristic temperature scales, vary linearly with $D$:

\begin{align*}
T_m &\approx 0.1566 D, \\
T_{sg} &\approx 0.1 D, \\
T_{sp} &\approx 0.16 D, \\
T_{\text{mod}} &\approx 0.147 D, \\
T_{s=0} &\approx 0.0713 D.
\end{align*}

As expected, in the limit $D = 0$ the paramagnetic phase is stable at all temperatures and corresponds to the only solution of the recursion relations. This is due to the fact that for $D = 0$ the system is much less frustrated and has a highly (i.e., extensively) degenerate...
ground state (i.e., \langle s \rangle\) approaches a finite value in the \(T \to 0\) limit, since each plaquette has a sixfold degenerate ground state which corresponds to the ice rule (two +1 and one -1 spins or two -1 and one +1 spins). In particular, for \(D = 0\) the model reduces to the nearest-neighbor Kagome spin ice model of Wills, Ballou, and Lacroix \cite{50}, for which a Pauling estimate yields the entropy \(s_{\text{GS}} = (3/2) \log[2(3/4)^{2/3}] \approx 0.75225\), while our mean-field approximation yields \(s_{\text{GS}} \approx 0.75204\). When the dipolar interactions are turned on (\(D > 0\)), such degeneracy is lifted, and a specific crystalline ground state structure emerges. The minimization of the local interactions produces a much stronger geometric frustration for \(T \lesssim D\) and gives rise to the emergence of a spin glass phase at low temperatures, characterized by an extremely rough free-energy landscape (at least at the mean-field level). The fact that all the relevant temperature scales of the problem show an apparent linear dependence on \(D\) is precisely due to the fact that the relevant energy scale is the energy difference between the ground state and the first excited states, which goes linearly to zero with \(D\).

\section{Conclusions}

In this paper we have developed an analytical mean-field treatment for the equilibrium properties of the DKIAFM introduced in \cite{45} and studied numerically in \cite{44}. Our mean-field approach is based on a cluster variational Bethe-Peierls formalism \cite{4} \cite{68–71} and on the cavity method \cite{49}, and consists in studying the model on a sparse random tree-like graph of triangular Kagome plaquettes, and cutting-off the dipolar interaction beyond the second nearest-neighbor plaquettes (i.e., the 5th nearest-neighbor spins). Our results essentially confirm and support the observations reported in Ref. \cite{44}, which were obtained using Monte Carlo simulations of relatively small system (ranging from 48 to 300 spins), and might be affected by both strong finite-size effects and by the difficulty of reaching thermal equilibrium in a reliable fashion due to strong metastability effects.

The summary of our results is the following. Upon decreasing the temperature we first find a transition to a sixfold degenerate crystal state which breaks time reversal, translation, and rotation symmetry as the one proposed in \cite{44} \cite{45}. Such transition is indeed discontinuous, as suggested in \cite{44}, although its first-order character turns out to be extremely weak, which might explain the strong finite-size effects observed in the finite-size scaling of the numerical data of the specific heat. When the system is supercooled below the first-order transition, we find that the paramagnetic state becomes unstable below a temperature at which the spin glass susceptibility diverges and a continuous spin glass transition takes place at the mean-field level.

On the one hand, the results presented here support and clarify the numerical findings of Ref. \cite{44}. On the other hand, they provide a first step to bridge the gap between the slow dynamics observed in geometrically frustrated magnetic systems and the mean-field theory of glassy systems formulated in terms of rough free-energy landscape. We believe that this analysis is of particular interest, especially in the light of the experimental relevance of the model, which could be potentially realized in several realistic set ups, including colloidal crystals \cite{58} \cite{59} \cite{76} \cite{79}, artificial nanomagnetic arrays \cite{60} \cite{61} \cite{80} \cite{81}, polar molecules \cite{62}, atomic gases with large magnetic dipole moments \cite{63}, and layered bulk Kagome materials \cite{64} \cite{66}.

Some comments are now in order. The lower-critical dimension of the spin glass transition is expected to be \(d_L \approx 2.5\) \cite{52} (at least in the case of short-range interactions). Hence we do not expect a genuine spin glass phase for the DKIAFM. Yet in \(2d\) the spin glass amorphous order can establish over very large (although not infinite) length scales and the spin glass susceptibility can become very large (although not infinite) at low temperature due to the vestiges of the transition. Indeed, there are plenty of experimental studies using thin films that at sufficiently low temperatures behave as the \(3d\) counterparts. See, e.g. \cite{54} for a very recent reference and \cite{53} for a more classical ones. The situation is similar concerning numerical simulations \cite{55}.

Concerning the dynamics, very early Monte Carlo simulations of the \(3d\) Edwards-Anderson model suggested that the spin auto-correlation function, close but above the expected critical temperature, decays as a stretched exponential \cite{50}. Therefore, although the model is ex-
pected to have a conventional second order phase transition with critical slowing down and algebraic decay of correlation functions, for the system sizes and time-scales accessed in this paper, the time-delayed correlations were satisfactorily fitted by such an anomalous form, with a stretching exponent decaying with decreasing temperature. Just a bit later, in [51] the conventional critical slowing down was recovered. A stretched exponential stretching exponent decaying with decreasing temperature. Just a bit later, in [51] the conventional critical slowing down was recovered. A stretched exponential relaxation of the self-correlation in the DKIAFM was reported in [44]. However, our results suggest that at sufficiently large time and length scales this behavior might be replaced by a conventional power law decay for control parameters in the critical region.

The case $J = 0$ and $D > 0$ [67] has been left over by the present investigation and might be an interesting subject for future studies. Possibly, the most interesting questions would be the investigation of how the properties of the model are affected by quantum fluctuations.

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Note that the mean-field value of the ground state energy is found to be lower than the 2d mean-field calculations. In particular, it can be shown that m = 1 always gives back the paramagnetic RS solution.