Fate of partial order on the trillium and distorted windmill lattices

Sergei V. Isakov,1 John M. Hopkinson,2 and Hae-Young Kee3,4

1 Institute for Theoretical Physics, ETH Zurich, CH-8093 Zurich, Switzerland
2 Brandon University, Brandon, Manitoba, Canada, R7A 6A9
3 University of Toronto, Toronto, Ontario, Canada, M5S 1A7
4 Korea Institute for Advanced Study, Seoul 130-722, Korea

The classical Heisenberg model on the trillium and distorted windmill lattices exhibits a degenerate ground state within large- \( N \) theory, where the degenerate wavevectors form a surface and line, in 3-dimensional space, respectively. We name such states partially ordered to represent the existence of long-range order along the direction normal to these degenerate manifolds. We investigate the effects of thermal fluctuations using Monte Carlo (MC) methods, and find a first order transition to a magnetically ordered state for both cases. We further show that the ordering on the distorted windmill lattice is due to order by disorder, while the ground state of the trillium lattice is unique. Despite these different routes to the realization of low temperature ordered phases, the static structure factors obtained by large- \( N \) theory and MC simulations for each lattice show quantitative agreement in the cooperative paramagnetic regime at finite temperatures. This suggests that a remnant of the characteristic angle-dependent spin correlations of partial order remains above the transition temperatures for both lattices. The possible relevance of these results to \( \beta \)-Mn, CeIrSi, and MnSi is discussed.

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I. INTRODUCTION

Geometrically frustrated (GF) magnetic systems offer a rich avenue to the search for emergent or collective phases of matter. Frustrated magnetism arises when spins between nearby magnetic sites cannot form a unique alignment to minimize their magnetic interactions, and leads to the possibility of a macroscopically degenerate classical ground state. Materials with antiferromagnetic nearest neighbor interactions between local moments lying at the sites of corner-sharing tetrahedra or triangles comprise a considerable fraction of the GF magnetic systems known to date.

Most magnetic materials select a particular ordering wavevector, representative of their magnetic structure, upon the onset of magnetic order. In contrast, in frustrated magnets, it is not uncommon to find within the mean field approximation a set of degenerate continuously connected “ordering wavevectors” which span the three-dimensional (3-D) space. For example, the disordered ground state of the classical Heisenberg model on the pyrochlore lattice features such a wavevector manifold. Partially ordered magnets lie between these two extremes, forming degenerate wavevector (2-D) surfaces or (1-D) lines in a 3-D space. The name of “partially ordered” comes from the fact that there is a long-range order along the direction normal to this degenerate manifold. In other words, in such systems, to very low temperatures, a system may appear disordered/ordered to measurements probing parallel/perpendicular directions to this line or surface. A growing number of frustrated spin models have been shown to exhibit such a partial order. Experimental evidence for partial order has been observed in single-crystal neutron scattering measurements of the correlated metals, MnSi under pressure, and CeCu\(_{5.9}\)Au\(_{0.1}\), although in these materials it is unlikely to arise from considerations as simple as here considered.

Our interest in this problem arose from the realization that partial order has been found using large- \( N \) theory on both the trillium and distorted windmill lattices. The key question we address in this paper is whether these partially ordered states can be captured beyond the large- \( N \) theory. If not, to what extent can the results of large- \( N \) theory be regained due to thermal fluctuations? We answer this question by carrying out large scale classical Monte Carlo simulations of the AF Heisenberg model on the trillium and distorted windmill lattices.

Large- \( N \) theory and MC simulations of the classical Heisenberg model give consistent results for both highly frustrated systems (ex: on the pyrochlore lattice a disordered ground state is found), and for unfrustrated lattices, (ex: choosing the correct ordering wavevector). Despite this trend, the finding of a partially ordered ground state in large- \( N \) theory does not necessarily translate into a partially ordered ground state within MC simulations. We find that the classical Heisenberg model on both the trillium and distorted windmill lattices shows a first order phase transition to a coplanar magnetically ordered state featuring neighboring spins rotated by 120°. We further show that this model on the trillium lattice does not have a macroscopic ground state degeneracy. The chosen 120° coplanar state is a unique ground state, and the partial order is an artificial effect of the spherical approximation constraint used in the large- \( N \) theory.

On the other hand, the selection of a particular coplanar ordering on the distorted windmill lattice proceeds due to an order by disorder mechanism.

In the cooperative paramagnetic state (for an intermediate temperature regime \( T_c < T < \theta_{CW} \) extending...
FIG. 1: (Color online) Magnetic sites of (a) the trillium lattice and (b) the distorted windmill lattice. In both cases three corner-shared equilateral triangles meet at each spin. The unit cell of the trillium lattice is only 4 sites and the next smallest spin loop is 5 sided. The unit cell of the distorted windmill lattice is 12 sites and the smallest spin loop is 4 sided. Atoms within the first unit cell are depicted in boldface (black) and labeled ((b) as in Table I).

II. MODEL, LATTICE STRUCTURES, AND APPROACHES

We study the classical $O(N)$ model on the trillium and distorted windmill lattices given by the following Hamiltonian

$$H = J \sum_{\langle ij \rangle} S_i \cdot S_j,$$

where $J > 0$ is the antiferromagnetic exchange coupling constant, the sum runs over the nearest neighbors only, and $S = (S^1, \ldots, S^N)$ is an $N$-component classical spin.

It was shown in Ref. 3 that the magnetic lattice of MnSi forms a three-dimensional network of corner-sharing equilateral triangles with the cubic $P2_13$ symmetry shown in Fig. 1 (a), which we have named the trillium lattice. The trillium lattice is common to many systems including the antiferromagnetically correlated Ce local moments of CeIrSi. The magnetic lattice of $\beta$-Mn, the distorted windmill lattice ($P4_132$ symmetry), bears a remarkable qualitative resemblance to the trillium lattice as shown in Fig. 1 (b). In particular, both lattice structures feature three corner-shared equilateral triangles joined at a common site. The coordinates of each site within a unit cell for both the trillium lattice and the distorted windmill lattice have been previously presented. For completeness, these are listed in Table I.

To carry out a quantitative comparison between large-$N$ theory and $N = 3$ Monte Carlo results, we have carried out large scale classical Monte Carlo simulations of the $N=3$ Heisenberg model on the trillium and distorted windmill lattices for lattices with $n_s = n \times L \times L \times L$ spins, where $n$ is the number of sites (or spins) in the unit cell and $L$ is the number of spins along each dimension of a cube. On the trillium lattice ($n=4$), we have considered $L = \{6, 9, 12, 18\}$. On the distorted windmill lattice ($n = 12$), we have considered $L = \{6, 8, 12\}$. The standard Metropolis algorithm has been used in which we...
and set a uniform $\lambda$ with the action $\phi$ attempt to update a spin within a small angular range $\delta$ around its original direction. We choose $\delta$ in such a way that around 50% of attempted spin updates are accepted. Starting with a random configuration, we usually perform $2 \cdot 10^5$ Monte Carlo steps (MCS) for equilibration and $10^6$ MCS for measurements with one MCS consisting of $\sim n_s/T$ single spin updates, where $T$ is the temperature.

Within large-$N$ theory, one writes the partition function for the $N$-component spin model\cite{99} and solves for the spin-spin correlations\cite{99,100,101} in the limit $N \to \infty$. That is we rewrite the Hamiltonian as

$$H = T \sum_{i,j} M_{ij} S_i \cdot S_j,$$

where $M_{ij}$ is the interaction matrix and the spins are subject to the constraint $S_i^2 = N$. The corresponding partition function is given by

$$Z = \int D\phi D\lambda \ e^{-S(\phi, \lambda)} \tag{3}$$

with the action

$$S(\phi, \lambda) = \sum_{i,j} \left[ \frac{1}{2} M_{ij} \phi_i \cdot \phi_j + \lambda \frac{N}{2} \delta_{ij} (\phi_i \cdot \phi_i - N) \right], \tag{4}$$

where $\phi_i = (\phi_{i1}, ..., \phi_{iN})$ is an $N$-component real vector field and $\lambda_i$ the Lagrange multiplier for the constraint $\phi_i \cdot \phi_i = N$. To proceed, we take the $N \to \infty$ limit and set a uniform $\lambda_i = \lambda_0$. The locations $i = (l, \mu)$ of spins can be labeled by those of the cubic unit cell $l = 1, \ldots, n_c$ and the lattice sites $\mu = 1, \ldots, n$ within the unit cell ($n_c = L \times L \times L$ is the total number of the unit cells in the lattice). The Fourier transform with respect to the positions of the unit cells leads to

$$S(\phi) = \sum_{\mathbf{q}} \frac{1}{2} M_{\mathbf{q}\mathbf{q}} \phi_{\mathbf{q},\mu} \cdot \phi_{\mathbf{q},\nu} \tag{5}$$

with $\sum_{\mathbf{q}} = M_{\mathbf{q}\mathbf{q}} + \delta_{\mathbf{q}, \mathbf{0}} \lambda_0$. Performing Gaussian integrations over the $\phi$ fields in Eq. (3) one finds that $\lambda_0$ is determined by the saddle point equation,

$$nn_c = \sum_{\mathbf{q}} \sum_{\rho=1}^n \frac{1}{\beta \epsilon_q^\rho + \lambda_0} \tag{6}$$

where $\beta \epsilon_q^\rho$ are the eigenvalues of the $n \times n$ interaction matrix $M_{\mathbf{q}\mathbf{q}}$ (which has been shifted such that $\text{Min}(\epsilon_q^\rho) = 0$) and $\beta$ the inverse temperature. Note that the sum over $q$ is carried out for finite size periodic lattices to effectively compare with the Monte Carlo results. Plugging $\lambda_0$ back into Eqs. (3) and (4), we readily deduce the spin-spin correlation functions. For example, one can be found by calculating the second derivative of $Z$ with respect to an auxiliary field which couples to the spin and can be added to Eq. (4). The static structure factor is found to be

$$S(q) \propto \sum_{\kappa, \kappa'} \langle S_\kappa^\nu S_{-\kappa'}^\nu \rangle = \frac{1}{\beta \epsilon_q^\kappa + \lambda_0} \tag{7}$$

where $U$ is the matrix that diagonalizes the interaction matrix $M_{\mathbf{q}\mathbf{q}}$, $\kappa$ and $\rho$ are the sublattice indices. Note that, for simplicity, we will normalize spins to $\sqrt{N}$ in the large-$N$ theory\cite{99}. To compare with MC results ($N = 3$), one needs to fix the energy scale. Here we rescale $J$ to $J/3$ in $\epsilon_q^\rho$.

### III. THE TRILLIUM LATTICE

Within large-$N$ theory, we reported\cite{99} that the classical antiferromagnetic (AF) Heisenberg model on the trillium lattice has a partially ordered ground state, with a surface of degenerate wavevectors following

$$\cos^2 \left( \frac{9\pi}{2} \right) + \cos^2 \left( \frac{9\pi}{2} \right) + \cos^2 \left( \frac{9\pi}{2} \right) = \frac{9}{4}, \tag{8}$$

where we set the lattice constant $a = 1$. However, in Ref.\cite{99} we relaxed the hard spin constraint of $S_i^2 = 1$ to a soft constraint, $\sum_{i=1}^n S_i^2 = n$, where $n$ is the number of sites in the unit cell. Using energy minimization on spin clusters of classical spins (with $N = 3$), we effectively imposed the hard spin constraint show that the lowest energy state actually exhibits a coplanar magnetic order with wavevector $(\frac{3\pi}{2}, 0, 0)$\cite{99}. We speculated that the soft constraint of conventional large-$N$ theory is crucial to the realization of partial order at $T \to 0$.
The position of the sites in a cubic unit cell. In this and all transition temperature is strongly first order transition. Our best estimate for the temperatures as the size of the system increases. We have effects, the location of this jump appears to scale to lower temperatures as the size of the system increases. We have carried out large scale MC simulations. The energy as a function of temperature, shown in Fig. 2 (a), exhibits a smooth behavior before an abrupt jump, which indicates a first order transition to an ordered state. The heat capacity (b) and magnetic order parameter (c) vs. temperature for the Heisenberg model on the trillium lattice. These results are independent of the lattice parameter \( u \) which determines the position of the sites in a cubic unit cell. In this and all other plots, error bars are smaller than the symbol size if not visible.

To find out whether a finite temperature transition to a magnetically ordered state occurs, we have carried out large scale MC simulations. The energy as a function of temperature, as seen in Fig. 2 (a), exhibits a smooth behavior before an abrupt jump, which indicates a first order transition to an ordered state. The heat capacity as a function of temperature, shown in Fig. 2 (b), clearly shows a peak corresponding to the onset of magnetic order on the trillium lattice. Because of strong hysteresis effects, the location of this jump appears to scale to lower temperatures as the size of the system increases. We have not attempted to determine the precise location of this strongly first order transition. Our best estimate for the transition temperature is \( T_c = 0.21(1)J \). In Fig. 2 (c), we plot the magnetic order parameter of this transition, which is defined as the structure factor at the ordering wavevector \( \mathbf{Q} = (\frac{2\pi}{3}, 0, 0) \):

\[
(m)^2 = \frac{S(q = \mathbf{Q})}{n_s}.
\]

We see that this order parameter exhibits a sharp onset at the transition temperature as would be expected for a first order phase transition.

It was anticipated that the MC would find a transition to a 120° coplanar magnetic state with the wavevector \( (\frac{2\pi}{3}, 0, 0) \), where the spins on each triangle form at 120° to each other, since it was shown (by minimization) that this is the lowest energy state. Thus this ground state does not have a macroscopic degeneracy and the observed transition is not expected to result from an order by disorder mechanism. An intuitive way to understand the uniqueness of this ground state is to start with a coplanar state with spins labeled \( \alpha \), \( \beta \) and \( \gamma \) where the letters \( \alpha \), \( \beta \) and \( \gamma \) denote 120° rotated spins on every triangle. It is then natural to ask whether one can generate degenerate states by rotating \( \beta \) and \( \gamma \) spins around the axis of spin \( \alpha \). The high connectivity of this lattice in comparison with the kagome and hyperkagome lattices\(^{20}\) where such degeneracies naturally arise, prevents any such rotations which are not consistent with the crystal symmetries of the lattice. Therefore, the rotation of \( \beta \) and \( \gamma \) spins around \( \alpha \) spin axis cannot generate distinctly different states. This argument itself is not sufficient to prove that the ground state is unique because, in principle, there might be other ground states that are not connected to the \( (\frac{2\pi}{3}, 0, 0) \) state by simple spin rotations. However, our previous minimization showed that the state with the wavevector \( (\frac{2\pi}{3}, 0, 0) \) is the lowest energy state. Combining our current MC result, we conclude that the ground state of Heisenberg model on the trillium lattice is unique.

In the disordered state at temperatures above this ordering transition, however, the spins fluctuate, and one might expect to recover features found by large-\( N \) theory with the soft constraint. Now the question is to what extent is the partial order of large-\( N \) theory recovered due to temperature fluctuations?

Magnetic susceptibility (not shown) has been calculated within both approaches and shows good agreement until very close to the ordering temperature \( T_c \approx 0.21J \), well below the Curie-Weiss temperature \( \Theta_{CW} = 2J \), and in the zero temperature limit. Qualitative comparisons between the large-\( N \) theory and Monte Carlo results for the structure factor are shown in Fig. 3 in the cooperative paramagnetic phase at temperatures slightly above the onset of magnetic order. We see excellent qualitative agreement between these two approaches. Within the cooperative paramagnetic window we see that the static structure factor on the trillium lattice peaks around the surface of a sphere-like shape determined by Eq. 8 although part of this sphere is obliterated by geometric (rather than energetic) effects. Within both approaches the geometric factor (that is cancellations between spin contributions within the unit...
FIG. 3: Contour plots of the intensity of the structure factor in the \( \text{hhk} \) ((a) and (b)) and \( \text{hk0} \) ((c) and (d)) planes for the trillium lattice in the cooperative paramagnetic regime (for \( u = 0.138 \), \( L = 12 \) and \( T = 0.25J \)) show prominent features near these surfaces. Classical Monte Carlo ((a) and (c)) agrees well with large-\( N \) results ((b) and (d)). The maximal intensity is shown in white, and the axes run from \(-4\pi\) to \(4\pi\) along \( k(001) \) (vertical), \( h(110) \) (horizontal), \( k(010) \) (vertical), \( h(100) \) (horizontal). Notice the prominent features near the zero energy surfaces given by Eq. 8 of the large-\( N \) theory.

IV. DISTORTED WINDMILL LATTICE

Mean field calculations on the distorted windmill lattice also revealed a partially ordered ground state, with a line of degenerate wavevectors along the \( \{qqq\} \) direction. In this paper we show that Monte Carlo simulations of the AF Heisenberg model on this lattice find an ordered ground state as on the trillium lattice. However, the physical origin of the two phase transitions differs. While the ordering in the trillium lattice can be understood by minimization, being the result of energetically different states, the ordering in the distorted windmill lattice cannot. Rather, it must proceed via an order by disorder mechanism. As on the trillium lattice it is interesting to ask whether the partial ordering features obtained by large-\( N \) (or mean field) theory can be found at finite \( T \), where one might expect the spins to fluctuate strongly.

Again, on the distorted windmill lattice, MC simulations find a jump in the energy as a function of temperature indicative of a first order phase transition. Nonetheless, this transition is qualitatively different from that seen on the trillium lattice, as it does not show strong hysteresis effects and much variation with system size. This is reflected in a sharp transition from one smooth energy vs. temperature curve to another at the transition temperature as seen in Fig. (a). The heat capacity as a function of temperature, shown in Fig. (b), clearly shows a peak corresponding to the onset of magnetic order. The variation of the order parameter shows

FIG. 4: (Color online) A quantitative comparison of the angle-dependent structure factor is shown along three high symmetry directions between the Monte Carlo (Heisenberg model) and large-\( N \) results for classical spins in the cooperative paramagnetic phase on the trillium lattice (\( u = 0.138 \), \( L = 12 \)). The solid line is the large-\( N \) result, while the symbols are from MC simulations.
an abrupt onset at the transition temperature. Here the order parameter has been defined in terms of the structure factor at the ordering wave vector \( Q_0 = (0, 0, 0) \) as

\[
\langle m \rangle^2 = \frac{S(q = Q_0)}{n_s}. \tag{10}
\]

As in the case of the trillium lattice, we have not attempted to determine the precise location of this first order transition. Our best estimate for the transition temperature is \( T_c = 0.155(5) J \).

To understand this spin ordering we have carried out an energetic minimization of spin clusters with \( N = 3 \). The minimization of spin clusters on the distorted windmill lattice with \( qgg \) symmetry is found to admit only two types of magnetic spin substructures: one of which can repeat as it is from one unit cell to the next, and another which undergoes a 120° spin rotation in progressing by one unit cell along the (111) direction. Thus one has topologically distinct ground state sectors, unit cell structures which are shown in Table III and labeled \( s_q = 0 \) and \( s_q = 2\pi/3 \) respectively, where the ordering wavevector in \( s_q = 2\pi/3 \) is \( Q_{sp} = (2\pi, 2\pi, 2\pi) \). Curiously, as each spin structure features coplanar 120° rotated spins on each triangle, generic ground state spin structures are found to interchange between these two orderings. The only constraint arising for a \( s_q = 2\pi/3 \) spin structure is that the spin structure must complete three times an integral number of \( q = \frac{2\pi}{3} \) unit cells within its boundaries. There are a macroscopic number of degenerate ground states which can be generated by this mixing of these two states, all of which exhibit a coplanar order. This tendency to form a coplanar spin structure is reminiscent of the nematically ordered state recently found for the classical Heisenberg model on the hyperkagome lattice.\(^\text{22}\) We estimate that the degeneracy of the lattice grows exponentially in the linear lattice size, \( L \), roughly as \( e^{0.69L/6} \), as shown in Appendix A.

\[
\begin{array}{c|c|c|c|c|c}
\text{label} & \text{near. neighb. 1} & \text{near. neighb. 2} & \text{\( s_q = 0 \)} & \text{\( s_q = 2\pi/3 \)} \\
\hline
a & c_\perp g, d_{\perp - g}, f, k & j_{\perp - l} & \alpha & \alpha \\
b & q, h, j, l & d_{\perp - g}, f & \alpha & \alpha \\
c & a_\perp g, d_{\perp - g}, e, l & h_{\perp - l}, i_{\perp - l} & \beta & \alpha \\
d & a_\perp g, d_{\perp - g}, e, l & h_{\perp - l}, i_{\perp - l} & \gamma & \alpha \\
e & c, f_{\perp + g}, h_{\perp + g}, l & g_{\perp - l}, k_{\perp + l} & \alpha & \beta \\
f & a, e_{\perp - g}, h_{\perp - k}, b, d_{\perp - g} & \beta & \beta \\
g & b, d, i, l & e_{\perp - l}, k & \beta & \beta \\
h & b, e_{\perp - g}, f_{\perp + l}, i_{\perp - g}, j_{\perp - l} & \gamma & \beta \\
i & d, q, j_{\perp - l}, k_{\perp + l} & e_{\perp - l}, h_{\perp + l} & \gamma & \beta \\
j & b, h, i_{\perp + l}, k_{\perp + l}, a_{\perp - l}, l_{\perp + l} & \beta & \gamma \\
k & a, f_{\perp + g}, j_{\perp - l}, e_{\perp - g}, d_{\perp - g} & \gamma & \gamma \\
l & b, c, e, g & a_{\perp - l}, j_{\perp - l} & \gamma & \gamma \\
\end{array}
\]

TABLE II: Connections and ground state candidates of the classical Heisenberg model on the distorted windmill (\( \beta\)-Mn) lattice. When the lattice parameter, \( y = \frac{2\sqrt{3}a}{16} \), then the nearest neighbors 1 and 2 are at equivalent distances. The final column shows the two spin structures minimization finds. Here \( \{\alpha, \beta, \gamma\} \) refer to 120° rotated spins. In moving to the next unit cell, the \( s_q = 0 \) structure is unchanged, while the \( s_q = 2\pi/3 \) structure replaces \( \alpha \rightarrow \beta \rightarrow \gamma \rightarrow \alpha \).

Let us refer to spin configurations which order with ordering wavevectors, \( Q_0 \) and \( Q_{sp} \), as \( Q_0 \) and \( Q_{sp} \) states respectively. To understand the selection of the \( Q_0 \) state over other states, we carried out MC simulations with a \( Q_{sp} \) state and a \( Q_{mix} \) state that has three planes with \( s_q = 0 \) spin configurations and three planes with \( s_q = 2\pi/3 \) spin configurations for \( L = 6 \) as initial states, which allows the system to remain in these states. We found that the energy of the \( Q_{sp} \) state just below the transition temperature is lower than the energy of the \( Q_{mix} \) state, and the energy of the \( Q_{mix} \) state is lower than the energy of \( Q_0 \), as shown in Fig. 5 (a). However, the latter state is...
always selected as a ground state if an initial state has a random configuration. This implies that the selection between \( Q_0 \) and other states is due to entropy. To confirm our intuition, we computed the specific heats of the \( Q_0 \), \( Q_{\text{mix}} \), and \( Q_{\text{sp}} \) states, and found that the specific heat of \( Q_0 \) state is larger than that of \( Q_{\text{mix}} \) or \( Q_{\text{sp}} \) states below the transition temperature as shown in Fig. 3 (b). This confirms there are more low lying modes for the \( Q_0 \) state, which leads to the selection of the \( Q_0 \) state over the other states of the degenerate ground state manifold.

Now let us study the cooperative paramagnetic state above the transition temperature. Magnetic susceptibility (not shown) has been calculated within both approaches and shows good agreement until very close to the ordering temperature \( T_c \approx 0.155J \), well below the Curie-Weiss temperature \( \Theta_{\text{CW}} = 2J \), and in the zero temperature limit. Qualitative comparisons between the large-\( N \) theory and Monte Carlo results for the structure factor are shown in Fig. 7 in the cooperative paramagnetic phase at temperatures slightly above the onset of magnetic order. We see excellent qualitative agreement between these two approaches. On the distorted windmill lattice the structure factor intensity is concentrated along the degenerate lines of the large-\( N \) result. Within both approaches there is very little weight in the first Brillouin zone due to the geometric factor. In Fig. 8 a quantitative comparison along several high symmetry directions between these two approaches is presented. There is extremely good agreement at all wavevectors.

V. DISCUSSION AND SUMMARY

A magnetic material where the magnetic properties can be described by the AF Heisenberg model on the trilium lattice is CeIrSi. It shows\(^{25}\) a Curie-Weiss susceptibility above 100 K with a magnetic moment of 2.56\( \mu_B/\text{Ce} \) atom and a \( \theta_{\text{CW}} = -24 \) K. At low temperatures, \( \chi^{-1} \) shows a gradual downturn\(^{28,29}\) a characteristic common to many triangle-based frustrated magnets. X-ray scattering shows the lattice structure of this material to have a cubic \( P2_13 \) symmetry\(^{28}\), with a positional parameter \( x_{\text{Ce}} = 0.6183\).\(^{30}\) Then the Ce sites, which hold magnetic moments consistent with 4\( f^1 \) electrons, form a trilium lattice of corner-shared triangles. The (non-zeroed) in-
The relevance of our study to $\beta$-Mn and MnSi is less obvious, as both materials are metallic. However, the magnetic sites of each lattice feature one of the three-dimensional corner-shared triangle lattices structures here studied, with $\beta$-Mn being the only known material to form in the distorted windmill structure. While the origin of the unusual non-Fermi liquid resistivity seen under pressure in MnSi\textsuperscript{22} ($\Delta\rho \approx AT^2$) is not at present understood, the same temperature exponent is observed\textsuperscript{22} in $\beta$-Mn, where it is expected to result from antiferromagnetic spin fluctuations. We believe it is worthwhile to investigate the possible link between the magnetic fluctuations and the non-Fermi liquid behavior in these materials. In this vein, it is interesting to note that powder neutron scattering down to 1.4 K\textsuperscript{22} in $\beta$-Mn shows no signature of magnetic order. We are hopeful that this study might provide the motivation for single crystal neutron scattering to be carried out on $\beta$-Mn.

In summary, we have used large-$N$ theory for $O(N)$ vector spins and classical MC simulations to study the AF Heisenberg model on two three-dimensional corner-shared triangle lattices, each site of which belongs to three equilateral triangles. The large-$N$ studies suggested that the geometrical frustration present would lead to a partially ordered state on both lattices. However, through the minimization of finite size spin clusters, we found the ground state manifolds on these two lattices to be quite different, despite the local similarity between these corner sharing triangle lattice structures. In both cases, we found that there is a first order transition to a magnetically ordered state using MC methods. We further showed that the trillium lattice exhibits a unique ground state with a spiral ordering, while the distorted windmill has a macroscopic ground state degeneracy. Magnetic ordering of the classical AF Heisenberg model on the distorted windmill lattice is therefore seen to arise via an order by disorder mechanism. The degeneracy of this model on the trillium lattice is seen to be an artificial effect of the soft-constraint of the large-$N$ theory.

Despite the above noted differences at low temperatures between large-$N$ and MC results, in the cooperative paramagnetic phase above the transition temperature, we find a remarkable resemblance between the respective spin-spin correlations. This leads us to ask whether the salient features of the large-$N$ theory, the angular and directional dependences of the spin-spin correlations found in the partially ordered state obtained by large-$N$ theory are present at finite temperatures above the transition temperature. As true partial order exhibits long range order along particular directions only as $T \to 0$, it is not possible to have partial order at any finite temperatures, since the spin-spin correlation decay exponentially at finite temperatures. This being said, the qualitative directional dependence characteristic of a partially ordered state survives above the transition temperature (note that the ground state is smoothly connected to the cooperative paramagnetic phase in large-$N$ theory) al-

![FIG. 8: (Color online) A quantitative comparison of the angle-dependent structure factor is shown along three high symmetry directions between the Monte Carlo (Heisenberg model) and large-$N$ results for classical spins in the cooperative paramagnetic phase on $\beta$-Mn lattice ($L = 8$). The solid line is the large-$N$ result, while the symbols are from MC simulations.](image)
With either the number of the general rules for allowed spin structures (provided below) or the present spin structures as presented in Table I, we might label -{s}^{i}_{q=0}, s^{i}_{q=0}, s^{i}_{q=2\pi/3}, s^{i}_{2\pi/3}, s^{i}_{q=2\pi/3}, s^{i}_{q=2\pi/3}. Here s^{i}_{q=0} is taken to be the spin structure as presented in Table I and s^{i}_{q} the same taking \( \alpha \rightarrow \beta \rightarrow \gamma \rightarrow \beta \). In progressing from one unit cell to the next along the (111) direction, we can follow \( s^{i}_{q=0} \) with either \( s^{i}_{q=2\pi/3} \) or \( s^{i}_{q=2\pi/3} \). Here \( s^{i}_{q=0} \) or \( s^{i}_{q=2\pi/3} \) can be followed by either \( s^{i}_{q=2\pi/3} \) or \( s^{i}_{q=2\pi/3} \). Replacing 1 \rightarrow 2 \rightarrow 3 \rightarrow 1 gives the general rules for allowed spin structures (provided the number of \( s^{i}_{q=2\pi/3} \) structures is a multiple of three).

With these rules, it becomes a problem in combinatorics to determine the ground state degeneracy.

For simplicity, let us consider a finite size lattice with \( L \times L \times L \) unit cells, with \( L \) divisible by 3. Clearly we have one state with only \( s^{i}_{q=2\pi/3} \) spin structures. By removing three \( s^{i}_{q=2\pi/3} \) spin structures, we can insert three \( s^{i}_{q=0} \) spin structures in their place, or more generally, removing \( 3m \) planes of unit cells which rotate as they progress along the (111) direction, allows the introduction of \( 3m \) planes of spin structures which need not rotate from one plane to the next. Upon such a removal there should be \( 3l - 3m \) available unit cell planes for \( 3m \) equivalent spin structures. This implies a degeneracy of approximately

\[
N_{tm} = \frac{(3l - 1)!}{(3m)! (3l - 3m - 1)!},
\]

where \( m \) can be \( \{0, 1, ..., l\} \). Summing over \( m \) then gives the total number of states:

\[
N_L \approx \sum_{m=1}^{l-1} \frac{(3l - 1)!}{(3m)! (3l - 3m - 1)!},
\]

\[
\approx \frac{1}{3} \sum_{m=0}^{3l-1} \frac{(3l - 1)!}{m! (3l - m - 1)!} = \frac{2^{3l-1}}{3} = \frac{2^L}{6},
\]

so the number of ground states grows exponentially in the linear lattice size, roughly as \( N_L \approx \frac{2^{3L}}{6} \approx \frac{2^{60L}}{6} \).

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**APPENDIX A: DEGENERACY OF THE DISTORTED WINDMILL LATTICE GROUND STATE**

To estimate the degeneracy of the ground state, we realize that fixing \( \{\alpha, \beta, \gamma\} \) in Table I leaves us with six tilings of the spins in the unit cell, which we might label \( \{s^{i}_{q=0}, s^{i}_{q=0}, s^{i}_{q=2\pi/3}, s^{i}_{q=2\pi/3}, s^{i}_{q=2\pi/3}, s^{i}_{q=2\pi/3}\} \). Here \( s^{i}_{q=0} \) is taken to be the spin structure as presented in Table I and \( s^{i}_{q} \) the same taking \( \alpha \rightarrow \beta \rightarrow \gamma \rightarrow \beta \). In progressing from one unit cell to the next along the (111) direction, we can follow \( s^{i}_{q=0} \) with either \( s^{i}_{q=2\pi/3} \) or \( s^{i}_{q=2\pi/3} \). Here \( s^{i}_{q=0} \) or \( s^{i}_{q=2\pi/3} \) can be followed by either \( s^{i}_{q=2\pi/3} \) or \( s^{i}_{q=2\pi/3} \). Replacing 1 \rightarrow 2 \rightarrow 3 \rightarrow 1 gives the general rules for allowed spin structures (provided the number of \( s^{i}_{q=2\pi/3} \) structures is a multiple of three).

\[
\text{FIG. 9: Contour plots of the intensity of the large-}\ N \text{theory structure factor in the } hhk \text{ plane for different values of } u \text{. The vertical axis is } k(001), \text{ and the horizontal axis is } h(110), \text{ where } k \text{ and } h \text{ both run from } (\pm 4\pi, 4\pi). \text{ Varying the parameter } u, \text{ does not change the energy within either the large-}\ N \text{ theory (here } N = 3, L = 24 \text{ and } T = 1/4.5J\text{), nor in classical Monte Carlo simulations. It does, however, change the geometric contribution to the structure factor quite dramatically. While near } u = 0.125 \text{ (center) one appears to see few vestiges of the degenerate spheres of the treatment, for both large (right) and small (left) values of } u, \text{ one sees that these are quite well captured. Curiously, most real systems seem to lie closer to the middle graph.}
\]

\[\text{A. M. Reimer, Phys. Rev. B 45, 7287, (1992).}\]

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\[\text{J. M. Hopkinson and H.-Y. Kee, Phys. Rev. B 74, 224441 (2006).}\]

\[\text{B. Canals and C. Lacroix, Phys. Rev. B 61, 11251 (2000).}\]
As the structure factor geometrically vanishes at the origin, we define the order parameter in terms of the structure factors at equivalent wavevectors \( Q = (\pi, 0, 0) \) or \((0, 0, \pm \pi)\).

In the graphs here we have used \( u = 0.138 \) which is the value relevant for the Mn sites on the MnSi lattice. This corresponds to the distance \( \sqrt{\frac{1}{2}^2 + \frac{3}{2}^2} \) for the trillium and distorted windmill lattices respectively. The positional parameter corresponds to the location within the first unit cell with coordinates \((x, x, x)\). The other three Ce atoms within the first unit cell should then be given by \( \{(x + \frac{1}{2}, \frac{1}{2}, 1 - x), \{1 - x, x + \frac{1}{2}, \frac{3}{2} - x\}\) and \(\{(\frac{1}{2} - x, -1 - x, x + \frac{1}{2})\}\).

In the thermodynamic limit corrections resulting from this assumption will be small.

Some of these states may be equivalent, but we expect this to contribute at most a factor of \( \frac{1}{M_{\text{even}}} \), which should merely be a logarithmic correction to our result.