Phase transition in the three dimensional Heisenberg spin glass: Finite-size scaling analysis

L. A. Fernandez, 1, 2 V. Martin-Mayor, 1, 2 S. Perez-Gaviro, 3, 2 A. Tarancon, 2, 4 and A. P. Young 5

1 Departamento de Física Teórica I, Universidad Complutense, 28040 Madrid, Spain.
2 Instituto de Biocomputación y Física de Sistemas Complejos (BIFI), Zaragoza, Spain.
3 Dipartimento di Fisica, SMC of INFN-CNR and INFN, Università di Roma La Sapienza, 00185 Roma, Italy.
4 Departamento de Física Teórica, Universidad de Zaragoza, 50009 Zaragoza, Spain.
5 Department of Physics, University of California, Santa Cruz, California 95064

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We have investigated the phase transition in the Heisenberg spin glass using massive numerical simulations to study larger sizes, $48^3$, than have been attempted before at a spin glass phase transition. A finite-size scaling analysis indicates that the data is compatible with the most economical scenario: a common transition temperature for spins and chiralities.

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I. INTRODUCTION

As a result of extensive and careful numerical studies, 1, 2, 3, 4, 5 there is now compelling evidence for a finite temperature phase transition in the Ising spin glass in three dimensions. However, the situation for the Heisenberg spin glass, in which the spins are classical 3-component vectors, is still controversial. The Heisenberg spin glass is a suitable first model to describe experimental systems with weak anisotropy, such as dilute Mn glasses, we have undertaken a massive set of simulations.

In view of this controversy over the nature of the transition in the three-dimensional Heisenberg spin glass, which is of great importance for the understanding of spin glasses, we have undertaken a massive set of simulations to study even larger sizes, $12, N = L^3$ where $L \leq 48$. Our conclusion is that the data is consistent with a common transition temperature for spins and chiralities, though, of course, numerics can never prove that they are exactly equal.

The paper is organized as follows. In Sect. II we define the model and the observables. Finite size scaling, which is central in our analysis, is recalled in Sect. III. Simulation details are in Sect. IV while our equilibration tests are addressed in Sect. V. We find that a uniform allocation of computational resources is inefficient (equilibration is much harder to achieve for some particular samples). The numerical results are in Sect. VI while our conclusions are presented in Sect. VII.

II. MODEL AND OBSERVABLES

We use the standard Edwards-Anderson spin glass model on a cubic lattice

$$\mathcal{H} = - \sum_{\langle i,j \rangle} J_{ij} \mathbf{S}_i \cdot \mathbf{S}_j,$$  \hspace{1cm} (1)

where the $\mathbf{S}_i$ are 3-component classical vectors of unit length at the sites of a simple cubic lattice, and the $J_{ij}$ are nearest neighbor interactions with a Gaussian distribution with zero mean and standard deviation unity. Periodic boundary conditions are applied.

The spin glass parameter is $\hat{q}^{\mu\nu}_{CG} = S^{(1)}_{i} S^{(2)}_{j}$, where “$(1)$” and “$(2)$” are two identical copies of the system (same interactions), and $\nu$ and $\mu$ are spin components. Its Fourier transform at wave vector $\mathbf{k}$, is denoted by $\hat{q}^{\mu\nu} (\mathbf{k})$.

For the Heisenberg spin-glass, Kawamura 5 defines the chirality from spins on a line: $\kappa^{\mu}_{i} = \mathbf{S}_{i+\mu} \cdot (\mathbf{S}_{i} \times \mathbf{S}_{i-\mu})$, where here $\mu$ refers to a direction on the lattice. The related chiral spin-glass order parameter is $\hat{q}^{\mu}_{CG,i} = \kappa^{\mu(1)}_{i} \kappa^{\mu(2)}_{i}$, its Fourier transform being $\hat{q}^{\mu}_{CG} (\mathbf{k})$.

The wave vector dependent susceptibilities are computed from the two order parameters:

$$\chi_{SG}(\mathbf{k}) = N \sum_{\mu,\nu} \langle |\hat{q}^{\mu\nu}(\mathbf{k})|^2 \rangle_{av}, \hspace{1cm} (2)$$

$$\chi_{CG}(\mathbf{k}) = N \langle |\hat{q}^{\mu}_{CG}(\mathbf{k})|^2 \rangle_{av}, \hspace{1cm} (3)$$

where $\langle \cdots \rangle$ denotes a thermal average and $[\cdots]_{av}$ denotes an average over disorder.
The susceptibilities yield the second-moment finite-lattice estimator of the correlation length \( \xi \):

\[
\xi_L = \frac{1}{2\sin(\pi/L)} \sqrt{\frac{\chi(0)}{\chi(k_{\text{min}})} - 1},
\]

with \( k_{\text{min}} = (2\pi/L, 0, 0) \). The spin and chiral correlation lengths are denoted by \( \xi_{SG,L} \) and \( \xi_{CG,L} \) respectively.

We also consider the spin and chiral Binder ratios defined by

\[
g_{SG} = \frac{11}{2} - \frac{9}{2} \left[ \frac{\langle q^4 \rangle_{av}}{\langle q^2 \rangle_{av}} \right]_{av}, \quad g_{CG} = \frac{5}{2} - \frac{3}{2} \left[ \frac{\langle q^4_{CG} \rangle_{av}}{\langle q^2_{CG} \rangle_{av}} \right]_{av},
\]

where \( q^2 = \sum_{\mu\nu} [\tilde{q}^{\mu\nu}(k=0)]^2 \), \( q^2_{CG} = \sum_{\mu} [\tilde{q}^{\mu}_{CG}(k=0)]^2 \).

### III. Finite-size scaling

Finite size scaling is a most useful data analysis method, that exploits finite size effects where they are largest (at criticality) to gather information on the infinite system, see e.g. Ref. [15].

Finite size scaling takes the form of an asymptotic expansion on the system size, \( L \). To leading order, for a quantity \( O \) diverging in the thermodynamic limit as \( O \propto |T - T_c|^{x_0} \), it takes the form

\[
O(L,T) = L^{x_0/\nu} f \left( \frac{T^{1/\nu}(T - T_c)}{L} \right),
\]

where \( f \) is an analytic function of its argument. In particular, since \( \xi_{SG,L}/L \) is dimensionless it has the finite size scaling form

\[
\frac{\xi_{SG,L}}{L} = \frac{\xi_{SG,sL}}{sL},
\]

where \( \nu \) is the correlation length exponent. There are similar expressions for \( \xi_{CG,L}/L \), and also for the Binder ratios in Eq. (3) since these too are dimensionless.

We shall see that corrections to scaling are quite large, even for the large sizes that we study, and so we need to consider corrections to the asymptotic scaling form in Eq. (1). To investigate this we determine the intersection temperatures \( T^*_{SG}(L,sL) \), where the data for \( \xi_{SG,L}/L \) agree for sizes \( L \) and \( sL \), i.e.

\[
\frac{\xi_{SG,L}}{L} = \frac{\xi_{SG,sL}}{sL},
\]

with an analogous expression for the chiral data.

Whereas Eq. (7) predicts that all the \( T^*_{SG}(L,sL) \) are equal to the spin glass transition temperature \( T_{SG} \), when one includes the leading corrections to scaling the \( T^*_{SG}(L,sL) \) are given by

\[
T^*_{SG}(L,sL) - T_{SG} = A^{(s)}_{SG} L^{-\omega - \frac{\eta}{\nu}}.
\]

Here, \( \omega \) is the exponent for the leading correction to scaling while the amplitude is

\[
A^{(s)}_{SG} = A_{SG} \left[ \frac{1 - s^{-\omega}}{s^{1/\nu} - 1} \right],
\]

with \( A_{SG} \) a (non-universal) constant. In practice, we do not have enough information to determine the \( s \) dependence in Eq. (9), so we take the \( A^{(s)}_{SG} \) to be separate constants for each value of \( s \) that we use (\( s = 2 \) and 3/2).

In fact we may combine Eqs. (10) and (8), to obtain a modern form of Nightingale’s phenomenological renormalization\(^{16,18}\) the so called quotient method

\[
\frac{O(sL,T^*_{SG}(L,sL))}{O(L,T^*_{SG}(L,sL))} = s^{2xO/\nu} \left[ 1 + \tilde{A}_{O,s} L^{-\omega} + \ldots \right],
\]

where the dots stand for higher order scaling corrections. Were \( T_{CG} \) and \( T_{SG} \) to be different, a similar expression would hold for quantities diverging at \( T_{CG} \). In particular, one may use Eq. (11) with temperature derivatives (to obtain 1 + 1/\( \nu \)) or with the susceptibilities at zero wave number (to obtain 2 – \( \eta \), where \( \eta \) is the anomalous dimension).

### IV. Simulation details

Simulations are run in parallel on \( N_T \) processors at \( N_T \) different temperatures using the parallel tempering\(^{20}\) (PT) method to speed up equilibration at low-\( T \), see Table I for the parameters.

As discussed in other work\(^{8,10}\) three types of moves are performed: (i) “overrelaxation” (OR) sweeps (which do not change the energy), (ii) “heat-bath” (HB) sweeps (which do change the energy), and (iii) parallel tempering (PT) sweeps in which the spin configurations at neighboring temperatures are swapped with a probability which satisfies the detailed balance condition. It is important to include OR sweeps because not only is the code for them much simpler (and hence faster) than that for the HB sweeps, but also because OR moves are very efficient\(^{8,9,21}\) in equilibrating the system, so many fewer sweeps are required than in a simulation with only HB and PT moves. Nonetheless, a fraction of the moves must be HB in order to change the energy. Because of the PT moves, the temperature of a set of spins (a “copy”) is not fixed but does a random walk between the minimum and maximum temperature in the set. In this way, each copy of spins can visit different “valleys” in configuration space with the correct statistical weight, even at low temperature.

The \( N_T \) temperatures were arranged in a geometric progression between \( T_{\text{min}} = 0.12 \) and \( T_{\text{max}} = 0.19 \). We do 1 HB sweep followed by 5L/4 OR sweeps and then 100 PT sweeps. We found a net CPU gain by doing a number of OR sweeps between HB sweeps which is somewhat greater than \( L \), perhaps because this transfers a fluctuation right across the system. We do a large number (100) of PT sweeps following right after each other, because the PT sweeps are very inexpensive in CPU time.
TABLE I: Parameters of the simulations. \( N_T \) is the number of temperatures. For the larger sizes, the number of sweeps varied from sample to sample. We show values for \( N_{\text{sweep}}^{\text{min}} \) and \( N_{\text{sweep}}^{\text{max}} \), the minimum and maximum number of overrelaxation sweeps.

| \( L \) | \( N_{\text{sweep}}^{\text{min}} \) | \( N_{\text{sweep}}^{\text{max}} \) | \( N_T \) | \( N_{\text{samp}} \) |
|------|-----------------|-----------------|-----|----|
| 8    | 5.0 \( \times \) 10^7 | 5.0 \( \times \) 10^7 | 5   | 984 |
| 12   | 7.5 \( \times \) 10^7 | 7.5 \( \times \) 10^7 | 9   | 984 |
| 16   | 1.0 \( \times \) 10^8 | 1.0 \( \times \) 10^8 | 15  | 984 |
| 24   | 1.5 \( \times \) 10^8 | 1.2 \( \times \) 10^7 | 27  | 984 |
| 32   | 4.0 \( \times \) 10^8 | 1.2 \( \times \) 10^8 | 43  | 984 |
| 48   | 6.0 \( \times \) 10^7 | 6.0 \( \times \) 10^8 | 79  | 164 |

FIG. 1: (color online) Top: Parallel tempering autocorrelation function as computed for three representative \( L = 48 \) samples. Here easy means that after \( N_{\text{sweep}}^{\text{min}} \) MC steps, see Table I the equilibration criterion was met (42% of samples), while medium samples (34%) required up to \( 2 N_{\text{sweep}}^{\text{min}} \) MC steps. Bottom: The quantity \( \Delta \) in Eq. (12) as a function of MC time, both in units of \( N_{\text{sweep}}^{\text{min}} \) (red triangles) and in units of the maximum number of sweeps for each sample (blue circles). For the latter, note that the data is computed at different times for different samples.

V. EQUILIBRATION

We do several tests to ensure equilibration. Firstly, we require that data satisfy the relation

\[
\Delta = \left[ \frac{q_s - q_t}{T} + \frac{2}{z} U \right]_{\text{av}} = 0, \tag{12}
\]

which is valid for a Gaussian bond distribution. Here

\[
U = -\sum_{\langle i,j \rangle} J_{ij} \langle S_i \cdot S_j \rangle, \tag{13}
\]

\[
q_l = \langle \frac{1}{N_b} \sum_{\langle i,j \rangle} (S_i \cdot S_j)^2 \rangle, \tag{14}
\]

\[
q_s = \langle \frac{1}{N_b} \sum_{\langle i,j \rangle} \langle (S_i \cdot S_j)^2 \rangle \rangle, \tag{15}
\]

in which \( U \) is the thermally averaged energy per spin, \( q_l \) is called the “link overlap”, \( N_b = (z/2)N \) is the number of nearest neighbor bonds, and \( z \) (= 6) here is the lattice coordination number. Both \( U \) and \( q_s \), being a single thermal average, come close to equilibrium relatively quickly as the number of MC sweeps increases. However, \( q_l \) involves a double thermal average, which is determined from two separate copies initialized with random spin configurations, and hence is initially very small. As the simulation proceeds, \( q_l \) increases towards its equilibrium value, so \( \Delta \) in Eq. (12) will initially be positive but will become zero (and stay zero) when equilibrium is reached. Data is shown in Fig. 1 for \( L = 48 \) (the largest size) at \( T = 0.12 \) (the lowest temperature).

Eq. (12) also provides a control variate to reduce statistical errors in \( \xi_{SG,L} \) and \( \xi_{CG,L} \). The key is in the strong statistical correlations between the Monte Carlo estimator for \( \Delta \) and those for the susceptibilities. Since we happen to know that \( \Delta = 0 \), reduced-variance estimators for the susceptibilities are obtained straightforwardly (see Ref. 22 for details). In practice, this method halves the errors for \( \xi_{CG,SG} \) at \( T = 0.12 \) (however, for \( T \gtrsim 0.14 \) the gain is less than 10%).

Some samples are harder to equilibrate than others so, ideally, we should spend more MC sweeps on the “hard” ones than on “easy” ones. The key to classifying samples in a PT simulation is to consider the dynamics of the temperature random-walk. In “hard” samples the \( T \) random-walk is slower (a copy trapped in a deep valley needs a longer time to wander to a \( T \) high enough to escape). We use correlation functions and autocorrelation times to formalize this idea, see Fig. 1 and the comments in Ref. 22.

For each sample, we impose a minimum number of sweeps, Table I then keep simulating until the total number of MC iterations exceed 9 autocorrelation times. For \( L = 48 \), the average number of MC iterations per sample was 1.8 times the minimum. Figure 1 shows that the data equilibrates more convincingly by running the “hard” samples for longer than the “easy” samples.

VI. RESULTS

We now present our results. Figure 2 shows data for the spin glass and chiral glass correlation lengths divided by \( L \). The resulting intersection temperatures, obtained from a jackknife analysis, are shown in Table II.
and $2 - \eta$ for the case of $s = 2$, using the quotient method \cite{15,16} see Eq. (11). The operators used are $\partial_T \xi_{SG, L}$, $\partial_T \xi_{CG, L}$, $\chi_{SG}$ and $\chi_{CG}$ which have scaling exponents $1 + 1/\nu_{SG}$, $1 + 1/\nu_{CG}$, $2 - \eta_{SG}$ and $2 - \eta_{CG}$ respectively. Spin (chiral) exponents are computed from data at $T_{SG}(L, sL)$ ($T_{CG}(L, sL)$).

\begin{table}[h]
\centering
\begin{tabular}{cccccccc}
$L$ & $sL$ & $T_{SG}$ & $T_{CG}$ & $\nu_{SG}$ & $\nu_{CG}$ & $2 - \eta_{SG}$ & $2 - \eta_{CG}$ \\
8  & 16  & 0.158(1)  & 0.156(1)  & 1.01(2)  & 1.34(5)  & 1.99(1)  & 0.72(2) \\
12  & 24  & 0.142(2)  & 0.150(1)  & 1.35(5)  & 1.51(6)  & 2.08(1)  & 0.96(3) \\
16  & 32  & 0.136(1)  & 0.147(1)  & 1.50(7)  & 1.46(6)  & 2.14(1)  & 1.11(3) \\
24  & 48  & 0.133(2)  & 0.142(1)  & 1.49(13) & 1.30(8)  & 2.19(2)  & 1.44(4) \\
8  & 12  & 0.164(2)  & 0.157(2)  &           &           &           &           \\
12  & 24  & 0.135(2)  & 0.147(2)  &           &           &           &           \\
32  & 48  & 0.130(3)  & 0.138(2)  &           &           &           &           \\
\end{tabular}
\caption{Table of intersection temperatures $T^*(L, sL)$ for the spin glass and chiral glass correlation length data presented in Fig. 2. Also shown are estimates for the exponents $\nu$ and $2 - \eta$ see Eq. (11). The operators used are $\partial_T \xi_{SG, L}$, $\partial_T \xi_{CG, L}$, $\chi_{SG}$ and $\chi_{CG}$ which have scaling exponents $1 + 1/\nu_{SG}$, $1 + 1/\nu_{CG}$, $2 - \eta_{SG}$ and $2 - \eta_{CG}$ respectively. Spin (chiral) exponents are computed from data at $T_{SG}(L, sL)$ ($T_{CG}(L, sL)$).}
\end{table}

Since Eq. (3) holds only asymptotically, for large $L$, it is necessary to decide on the smallest size $L_{min}$ to be included in the analysis. We consider first the five pairs of sizes with $L_{min} = 12$, see Table II. Fitting spin and chiral data separately there are 4 parameters for each: $T_{CG, SG}$, the exponent $\omega + 1/\nu$, and amplitudes $A_{(2)}$ and $A_{(3/2)}$. For the $s = 2$ and $s = 3/2$ size ratios. We determine the best fit parameters, and estimate the quality of the fit from the value of $\chi^2$. Fitting the spin data to Eq. (3), gives $T_{SG} = 0.129^{+0.016}_{-0.016}$, which is compatible with Viet and Kawamura’s result of 0.120(6). However, in the chiral sector $\chi^2$ as a function of $T_{CG}$ does not have a local minimum with $T_{CG} > 0$, so subleading scaling corrections are sizable for chiralities and $L_{min} = 12$.

Hence we have also performed an analysis with a larger value, $L_{min} = 16$. Unfortunately, we only have data for four pairs of sizes, and still four parameters to be fitted if we fit the spin and chiral data separately. Since the number of points is equal to the number of parameters we do not gain useful information. However, if we assume a common transition temperature and do a joint fit we have 8 data points, and 6 parameters (1 transition temperature, one exponent, and 4 amplitudes). The resulting fit gives $T_c (= T_{SG} = T_{CG}) = 0.120^{+0.010}_{-0.100}$ with a $\chi^2$ per degree of freedom of 0.029 so the fit is good. The error bar on $T_c$ is very large on the low-$T$ side but if we assume that the exponent $1/\nu + \omega$ in Eq. (6) is greater than 0.5, plausible given the values for $\nu$ in Table II we find $T_c = 0.120^{+0.010}_{-0.004}$, much more tightly constrained.

In Fig. 3 we show data for the spin glass and chiral glass Binder ratios defined in Eq. (5) of the text. Our definition of $g_{CG}$ differs from that of Kawamura and Viet \cite{10,11} and our results have an even more pronounced negative dip. Interestingly, we find that the results for $g_{SG}$ also become negative at the largest sizes. Hence, the apparent vanishing of $g_{SG}$ near $T_{CG}$, a strong argument for spin-chirality decoupling\cite{15} is an artifact caused by the lattice sizes being too small and the temperatures too high. Our interpretation of the Binder parameter data is that there is negative dip in both channels, and the minimum of this dip approaches the transition temperature as $L$ grows. The chiral-dip approaches $T_{CG}$ from high temperatures, while the spin-dip approaches $T_{SG}$ from low temperatures, where plausibly $T_{SG} = T_{CG}$. However, much larger sizes would be needed to confirm this hypothesis.

By studying sizes $L \leq 32$ Viet and Kawamura\cite{10,11} find $T_{SG} = 0.120(6)$ and $T_{CG} = 0.145(4)$. Since the difference is about 3.5 times the errors they argue that $T_{CG} > T_{SG}$. However, their value for $T_{CG}$ is actually
higher than our intersection temperatures for $L = 48$ shown in Table II and so seems to us to be too high. Also they estimate the transition temperatures from $T^*(L, sL) - T_{CG} = \text{const.}/L^\nu$, where $L^\nu$ is the average of $L$ and $sL$, rather than Eq. (9). In other words they replace the exponent $\omega + 1/\nu$ by 1, and the $s$ dependence in Eq. (9) by $2/(1+s)$. At the very least, we argue that these replacements lead to an underestimate of the error bars. Hence, we do not feel that the results of Viet and Kawamura contradict our conclusion that the data is consistent with the spin and chiral glass transition temperatures being equal.

VII. CONCLUSIONS

In summary, our low-temperature simulations for the Heisenberg spin glass are unprecedented in system size. To achieve thermalization, we have needed not only a huge amount of CPU ($7 \times 10^6$ hours) but a careful sample-by-sample thermalization check that allowed us to concentrate efforts on the “hard” samples. The results for the spin-glass sector can be accounted for using only leading-order scaling corrections, but subleading corrections are sizable for the chiral glass sector. This is the reason for the overestimate of $T_{CG}$ in previous work.\textsuperscript{10,11} Data for $L \geq 16$ support the most economic scenario, $T_{SG} = T_{CG}$. We also see that the spin Binder parameter is not trivial at $T_{CG}$.

Acknowledgments

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