Critical behavior of the dilute antiferromagnet in a magnetic field

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We study the critical behavior of the diluted antiferromagnet in a field with the tethered Monte Carlo formalism. We compute the critical exponents (including the elusive hyperscaling violations exponent $\theta$). Our results provide a comprehensive description of the phase transition and clarify the inconsistencies between previous experimental and theoretical work. To do so, our method addresses the usual problems of numerical work (large tunneling barriers and self-averaging violations).

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Understanding collective behavior in the presence of quenched disorder has long been one of the most challenging and interesting problems in statistical mechanics. One of its simplest representatives is the random field Ising model (RFIM), which has been extensively studied both theoretically and experimentally.1 The RFIM is physically realized by a dilute antiferromagnet in an applied magnetic field (DAFF).

It is known that the $D = 3$ DAFF (or RFIM) undergoes a phase transition, but the details remain controversial, with severe inconsistencies between analytical, experimental, and numerical work. A scaling theory is generally accepted, where the dimension $D$ of the system is replaced by $D - \theta$ in the hyperscaling relation. This third independent critical exponent, believed to be $\theta \approx 1.5$, is inaccessible both to a direct experimental measurement and to traditional Monte Carlo methods.

The values of the remaining critical exponents, seemingly more straightforward, are also controversial. On the experimental front, different ansätze for the scattering line shape yield mutually incompatible estimates of the thermal critical exponent, namely, $\nu = 0.87(7)$ (Ref. 2), or $\nu = 1.20(5)$.5 Furthermore, the experimental estimate of the anomalous dimension $\eta = 0.16(6)$ (Ref. 2) violates hyperscaling bounds, if one is to believe the experimental claims of a diverging specific heat ($\alpha \geq 0$).4

On the other hand, the numerical determination of $\nu$ has steadily shifted, the most precise estimate being $1.37(9)$,5 inconsistent with the experimental values and barely compatible with $\alpha \approx 0$. The value of $\alpha$ itself is very hard to measure in a numerical simulation.6

More fundamentally, the smallness of the magnetic exponent $\beta$, combined with the numerical observation of metastability,7 has led some authors to suggest that the transition in the DAFF may be of first order.

Ultimately, the physical reasons for this confusion betray the fact that the traditional tools of statistical mechanics are ill suited to systems with rugged free-energy landscapes. Both experimentally and numerically, the system gets trapped in local minima, with escape times that grow as $\log \tau \sim \xi^\xi$ ($\xi$ is the correlation length). This not only makes it exceedingly hard to thermalize the system, but also generates a rare-events statistics, causing self-averaging violations.5

In this Rapid Communication we study the DAFF with the tethered Monte Carlo (TMC) formalism.9 Our approach restores self-averaging and is able to negotiate the free-energy barriers of the DAFF to equilibrate large systems safely. It also provides direct access to the key parameter $\theta$. We thus obtain a comprehensive picture of the phase transition, consistent both with analytical results for the RFIM and with experiments on the DAFF, and shed light on the reasons behind the previous discrepancies.

In the following we provide a brief outline of the tethered formalism applied to the DAFF (see Refs. 9 and 10 for details). We note, however, that most of our physical results are translated into the familiar canonical language. In a tethered computation, we run simulations where one (or more) order parameters of the system are (almost) constrained. In this way, we eliminate the need for exponentially slow tunneling caused by the free-energy barriers associated with these parameters. From these tethered simulations the Helmholtz effective potential is accurately reconstructed with a fluctuation-dissipation formalism.

We consider a system with $N = L^D$ spins, $s_x = \pm 1$, on the nodes of a cubic lattice with periodic boundary conditions and interacting through the Hamiltonian

$$H = \sum_{(x,y)} \epsilon_x s_x \epsilon_y s_y - h M - h_s M_s = U - h M - h_s M_s. \quad (1)$$

Here $h$ and $h_s$ are the applied fields, coupled to the magnetization and staggered magnetization,

$$M = N m = \sum_x \epsilon_x s_x, \quad M_s = \sum_x \epsilon_x s_x e^{i\pi \sum_{\mu = 1}^D \mu_x}. \quad (2)$$

We are ultimately interested in $h_s = 0$, but we will find this parameter useful. The quenched occupation variables $\epsilon_x$ are 1 with probability $p = 0.7$ and zero otherwise (this value is chosen to be far both from the percolation threshold and from the pure system). For $D = 3$, the system undergoes a paramagnetic-antiferromagnetic phase transition, where $m_s$ is the order parameter.

Let us consider a single sample of the system (i.e., a fixed $\{\epsilon_x\}$). In our tethered computation, we define smooth magnetizations $\hat{m}$ and $\hat{m}_s$ by coupling $m$ and $m_s$ to Gaussian baths and work in a statistical ensemble for fixed ($\hat{m}, \hat{m}_s$) with weight9

$$\omega(\hat{m}, \hat{m}_s; \{s_x\}) \propto e^{-\beta U} \gamma(\hat{m}, m) \gamma(\hat{m}_s, m_s), \quad (3)$$

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where $\gamma(\hat{x},x) = e^{N(x-\hat{x})(\hat{x} - x)/N-2}/\Theta(\hat{x} - x)$, and $\Theta(\hat{x} - x)$ is the step function. The smoothing procedure shifts the mean value of the parameters, so $\hat{x} \simeq x + 1/2$. This ensemble is related to the canonical one through a Legendre transformation. For instance, the partition function of the system is

$$Z = \int d\hat{m} d\hat{m}_s \sum \omega(\hat{m},\hat{m}_s; \{s_x\}) e^{\beta N(\hat{m} + h,\hat{m}_s)}$$

$$= \int d\hat{m} d\hat{m}_s e^{-N[\Omega_e(\hat{m},\hat{m}_s) - \hat{m} h,\hat{m}_s]}$$

where $\Omega_e(\hat{m},\hat{m}_s)$ is the Helmholtz effective potential.

We can reconstruct $\Omega_e$ from computations at fixed $(\hat{m},\hat{m}_s)$ via the so-called tethered field $(\hat{b},\hat{b}_s)$

$$\hat{b} = 1 - 1/2 - 1/N \hat{m} - m, \quad \hat{b}_s = 1 - 1/2 - 1/N \hat{m}_s - m_s$$

In particular, the gradient $\nabla \Omega$ is

$$\langle \hat{b}, \hat{b}_s \rangle = \langle \hat{b}, \hat{b}_s \rangle$$

The notation $(\cdot \cdot \cdot )_{\hat{b}, \hat{b}_s}$ denotes tethered expectation values, computed with weight $(3)$.

A TMC computation consists of a set of independent Monte Carlo simulations at fixed $(\hat{m},\hat{m}_s)$ that are then combined to reconstruct $\Omega_e$. Note that the effective potential (as a function of the magnetizations) has all the information about the system in the tethered ensemble, just as the free energy (as a function of the applied fields) has all the information in the canonical ensemble.

The canonical averages at fixed $(h,h_s)$ can be recovered with Eq. (4). Note that, according to (6), this integral is dominated by saddle points $(\hat{m},\hat{m}_s)$ such that

$$\langle \hat{b}, \hat{b}_s \rangle = \beta h, \quad \langle \hat{b}_s, \hat{b}_s \rangle = \beta h_s$$

The notation $(\cdot \cdot \cdot )_{\hat{b}, \hat{b}_s}$ denotes tethered expectation values, computed with weight $(3)$.

So far we have summarized the application of TMC for a single sample. Since it consists of simulations at fixed $(\hat{m},\hat{m}_s)$, it eliminates the need to tunnel between coexisting phases and, hence, equilibrates the system much faster than a canonical simulation. However, we still face the serious problem of self-averaging violations. In principle, the definition of quenched disorder implies reconstructing the free energy with (4) before computing the disorder average. In this Rapid Communication, however, we sample average the Helmholz potential rather than the free energy (a similar approach was taken in Ref. 11).

In order to motivate this approach, let us consider the top panel in Fig. 1. We compare the tethered average $(\hat{b},\hat{b}_s)_{\hat{m},\hat{m}_s}$ for two individual samples with the disorder average over 1000 samples. The zeros of this latter curve separate an internal gap with chaotic fluctuations, where the field vanishes in the thermodynamical limit, from an external region where the field is actually self-averaging.

We exploit the situation by considering a small, but finite, value of $h_s$. The saddle point defined by this field will be in the self-averaging region. We can therefore solve the saddle-point equations (7) on average, rather than by sample by sample. Only afterward do we make $h_s \rightarrow 0$ in the solution (this is analogous to the mathematical definition of spontaneous symmetry breaking). The limit $h_s = 0^+$ is essentially equivalent to considering a “smeared” saddle point and averaging over all $\hat{m}_s$

$$\overline{\langle O \rangle}_{\hat{m}} = \int d\hat{m}_s \langle O \rangle_{\hat{b},\hat{b}_s} e^{-N[\Omega_e(\hat{m},\hat{m}_s) - \Omega_0]}$$

$\Omega_0$ is a normalization constant. Since we work at fixed $\hat{m}$, $\Omega_N$ is just the one-dimensional integral of $\langle \hat{b}, \hat{b}_s \rangle$.

The other saddle-point equation $(\hat{b},\hat{b}_s) = \beta h_s$ defines a one-to-one relation $\hat{m}_s(h)$ so that $\overline{\langle O \rangle}_{\hat{m}(h)}$ and the canonical $\langle O \rangle(h)$ both tend to the same thermodynamical limit (ensemble equivalent). Furthermore, for finite lattices $\overline{\langle O \rangle}_{\hat{m}}$ is better behaved statistically and arguably more faithful to the physics of an experimental sample. Therefore, we shall identify $\overline{\langle O \rangle}(h) = \overline{\langle O \rangle}_{\hat{m}(h)}$ and use the more familiar canonical notation. See Refs. 9 and 10 for a more detailed study of this ensemble equivalence.

We have used the above outlined procedure to thermalize the DAFF for temperatures down to $T = 1.6$ and sizes up to $L = 32$ (1000 samples for $L = 8, 12, 16, 24,$ and 700 samples for $L = 32$). For each size we simulate a grid of $\approx 150$ points in the $(\hat{m},\hat{m}_s)$ plane (five values of $\hat{m}_s$, and $\approx 30$ values of $\hat{m}_s$ on each). We also use temperature parallel tempering. This is only necessary to thermalize $L > 24$, but it is convenient for smaller lattices because we are also interested in the $T$ dependence. Thermalization is ensured using the methods described in Ref. 12. We provide more technical details in Ref. 10.

The first interesting physical result is the effective potential itself. Some authors have found metastable behavior in the DAFF, interpreted as a sign of a first-order transition.7 This should manifest as the coexistence of antiferromagnetic and paramagnetic minima in $\Omega$. However (see the bottom panel in Fig. 1), our results exhibit only two antiferromagnetic minima, separated by a very large free-energy barrier. In a canonical simulation, the system tunnels back and forth between the two, with an escape time $\tau \sim \exp[N \Delta \Omega]$. This explains the metastable behavior observed in previous work.
scaling (we use corrections, but not eliminated them completely. and using the quotients method, we have minimized the scaling phase: the free-energy barrier between the ordered and the disordered numerical estimates of $\xi$ experimental work.

We can compute this barrier simply by evaluating the line integral of $(\langle b \rangle_{\tilde{m},\tilde{m}_0} - \beta \tilde{h}_{\tilde{m}}$, $(\tilde{b}_s)_{\tilde{m},\tilde{m}_0})$ along a path joining the two saddle points. We know that one of them will lie on the line $\tilde{m}_0 = 0.5 (m_0 \approx 0)$. Therefore, we first integrate from the antiferromagnetic saddle point to $\tilde{m}_0 = 0.5$ at fixed $\tilde{m}_0$. We then integrate at fixed $\tilde{m}_0 = 0.5$ until we reach the disordered saddle point. We give the resulting values of $\Delta \Theta_N = \Delta F/N$ in Table II. Our final estimate is $\theta = 1.469(20)$, incompatible with the $\theta = D - 1$ of a first-order phase transition.

Notice that the hyperscaling relation $2 - \alpha = \nu (D - \theta)$, coupled with our values for $\nu$ and $\theta$, predicts not only a divergence of the specific heat, as observed in experiments, but also a positive $\alpha$. We could test this result directly by computing $C = \partial h \xi$. Unfortunately, the quotients method is ill suited to this quantity, whose scaling is more aptly described as $C \simeq A + B L^{\eta/\nu}$. Therefore, one needs extremely large values of $L$ to reach the asymptotic regime $C \sim L^{\eta/\nu}$. The behavior of the quotients in Table I is consistent with this expectation.

It has been proposed that $\theta$ is not independent, but given by $\theta = D/2 - \beta / \nu$. Combining Tables I and II, we see that our numerical results are indeed compatible with this two-exponent scenario.

We can use our results to comment on the experimental situation. In an experimental study, the critical exponents are computed from fits to the scattering line shape $S(k) = S_0(k) + S_1(k)$, where the two terms distinguish connected and disconnected contributions. In the two-exponent scenario, strongly supported by our data, the most singular term in $S_0$ is the square of $S_c$. This ansatz was applied in Ref. 2, yielding $\nu = 0.87(7)$ and $\eta = 0.16(6)$. Since $\eta = \theta - 1 + 2 \beta / \nu$, however, this last value violates hyperscaling bounds and is also

![Graph showing correlation length $\xi$ as a function of the applied magnetic field $h$ for $T = 1.6$. The curves intersect, marking a second-order phase transition. Bottom: Scaling plot of $\xi$ as a function of $T$ for $h = -2.13$, showing large corrections to leading scaling (we use $\nu = 1.05$).](image)

<table>
<thead>
<tr>
<th>$h^*(L)$</th>
<th>$\beta / \nu_h$</th>
<th>$\nu_h$</th>
<th>$\alpha / \nu_h$</th>
<th>$\nu_T$</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>-2.178(4)</td>
<td>0.0125(7)</td>
<td>0.887(5)</td>
<td>0.075(25)</td>
</tr>
<tr>
<td>12</td>
<td>-2.140(5)</td>
<td>0.0104(5)</td>
<td>0.790(9)</td>
<td>0.0781(27)</td>
</tr>
<tr>
<td>16</td>
<td>-2.123(3)</td>
<td>0.0119(4)</td>
<td>0.742(7)</td>
<td>0.224(4)</td>
</tr>
</tbody>
</table>

We need an additional critical exponent in order fully to characterize the critical behavior of the DAPF. This is the hyperscaling violations exponent $\theta$, which can be related to the free-energy barrier between the ordered and the disordered phase: $\Delta F \propto L^\theta$. The computation of these barriers is very difficult with traditional methods, but straightforward with TMC. Indeed, we can identify $\Delta F$ with the $\Delta \Theta_N$ between the two saddle points (disordered and antiferromagnetic) defined by the critical $h_c$.

![Graph showing scaling plot of $\xi$ as a function of $L^{1/\nu}$ for different $L$.](image)

**Table I.** Computation of the critical exponents using the quotients method. We extract our estimates from ratios of physical observables for sizes ($L, 2L$), computed at the intersection point of $\xi/L$. The first four columns give results for fixed $T = 1.6$ and the last one at fixed $h = -2.13$.

<table>
<thead>
<tr>
<th>$L$</th>
<th>$h^*(L)$</th>
<th>$\beta / \nu_h$</th>
<th>$\nu_h$</th>
<th>$\alpha / \nu_h$</th>
<th>$\nu_T$</th>
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<td>1.07(9)</td>
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<td>1.01(4)</td>
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<tr>
<td>16</td>
<td>-2.123(3)</td>
<td>0.0119(4)</td>
<td>0.742(7)</td>
<td>0.224(4)</td>
<td>1.10(15)</td>
</tr>
</tbody>
</table>

**Table II.** Computation of the hyperscaling violations exponent $\theta$ from the free-energy barriers $\Delta F$. We report fits to $\Delta F = A L^\theta$, for different ranges, giving the $\chi^2$ and the degrees of freedom of each fit. Our preferred final estimate is $\theta = 1.469(20)$, taking the central value of the fit for $L \geq 12$ and the more conservative error of the fit for $L \geq 16$.

<table>
<thead>
<tr>
<th>$L$</th>
<th>$\Delta F / N$</th>
<th>Fit range</th>
<th>$\theta$</th>
<th>$\chi^2/\text{d.o.f.}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>0.03382(29)</td>
<td>$L \geq 8$</td>
<td>1.448(9)</td>
<td>5.56/3</td>
</tr>
<tr>
<td>12</td>
<td>0.01756(15)</td>
<td>$L \geq 12$</td>
<td>1.469(13)</td>
<td>0.44/2</td>
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<tr>
<td>16</td>
<td>0.01138(9)</td>
<td>$L \geq 16$</td>
<td>1.461(20)</td>
<td>0.16/1</td>
</tr>
<tr>
<td>24</td>
<td>0.00608(5)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>32</td>
<td>0.00392(5)</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>
incompatible with our results. Perhaps taking $S_\eta = (S_\nu)^2$ for the whole function, not just its singularity, is an excessive simplification. Clearly a better theoretical determination of $S(k)$ is needed. Our methods are well suited to a direct numerical approach to this question.

We have used the tethered formalism to obtain a comprehensive picture of the critical behavior of the DAFF, resolving the inconsistencies in previous work. This method restores self-averaging to the problem and is capable of handling rugged free-energy landscapes to equilibrate much larger systems than canonical parallel tempering. Our simulations show clear signs of a second-order phase transition and are consistent both with experiments on the DAFF and with analytical results for the RFIM. The critical exponents $\theta$ and $\beta/\nu$ (equivalently, $\eta$ and $\tilde{\eta}$) are computed with a high precision, although our simulations were not optimized for the computation of $\nu$ (equivalently, of $\alpha$). We obtain $\nu = 0.90(15)$, consistent with a positive $\alpha$.

The tethered approach demonstrated in this Rapid Communication has a very broad scope and we believe it can be fruitfully applied to many systems featuring large free-energy barriers. Indeed, it has already been successfully implemented for hard-spheres crystallization.17 Other promising avenues are the study of Goldstone bosons and the equation of state for the $D = 3$ spin glass,18 or equilibrium and aging relaxation in a metastable phase (e.g., to prevent crystallization of supercooled liquids, see Ref. 19).

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