Simulation of vacancies in a two-dimensional Ising antiferromagnet

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We present simulations of one- and two-vacancy states in a two-dimensional square-lattice Ising antiferromagnet. We find that the energy of the single-vacancy ground state scales as $J^{2/3}$, where $J$ is the spin-spin coupling. The $S$- and $P$-wave states of two vacancies are bound, with binding energy $\mathcal{E} \propto J^{1/3}$ and $J$, respectively, while the $D$-wave state is unbound.

I. INTRODUCTION

The discovery of copper-oxide-based high-temperature superconductors has stimulated a large number of theoretical and experimental investigations and generated interest in non-phonon-mediated pairing. Because of the high transition temperatures and the small isotope effects, it is believed that superconductivity in these new materials arises primarily from strongly interacting electrons, rather than from phonons. In addition, these materials exhibit antiferromagnetic correlations.

There have been suggestions that vacancies in the anti-ferromagnetically ordered copper-oxide planes form pairs and thus give rise to superconductivity. A fully microscopic description of this phenomenon could be found, for example, in the Hubbard model near half-filling. However, more tractable models of vacancies in spin systems (Heisenberg or Ising) could yield equivalent descriptions with reduced complexity.

There have been several studies along these latter lines. In Ref. 9, a finite basis of states is diagonalized to find one- and two-vacancy states in a square two-dimensional (2D) antiferromagnet. In Ref. 10, the approximation of no closed loops (introduced in Ref. 11 for single vacancy when there is no coupling) was extended to finite couplings for both one- and two-vacancy Ising systems.

In this paper, we employ Monte Carlo methods to study both a single vacancy and the interaction of two vacancies in a 2D square-lattice Ising antiferromagnet. Our method is, in principle, exact and in practice is subject only to statistical and discretization errors, both of which can be controlled. We find binding of two vacancies in states with spatial $S$ and $P$ symmetries.

Our presentation is organized as follows. In Sec. II we introduce the model and describe how our basis states are constructed. In Sec. III the Monte Carlo procedure used to solve the path-integral problem is discussed. Section IV will present the numerical results of our studies for some test cases, for which analytic results are available, and for one and two vacancies in Ising antiferromagnets. A discussion of our results can be found in Sec. V.

II. THE MODEL

We investigate the Ising truncation of the two-dimensional Hubbard Hamiltonian in the large-$U$ limit,

$$\mathcal{H} = -t \sum_{\langle i,j \rangle, \sigma} (c_{i \sigma}^{\dagger} c_{j \sigma} + \text{H.c.}) + J \sum_{\langle i,j \rangle} s_{3i} s_{3j}$$

$$= \mathcal{H} + \mathcal{V},$$

(1)

where the summation extends over all nearest-neighbor pairs of sites $\langle ii \rangle$ on a 2D square lattice, and $J = t^2/U$. $c_i^{\dagger}$ and $c_i$ are the usual Fock-space fermion creation and annihilation operators, and the spin operators are defined as $s_{3i} = \sum_{\sigma} c_{i \sigma}^{\dagger} \tau_{\sigma\sigma'} c_{i \sigma'}$, where $\tau^2$ is the Pauli matrix

$$\begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}.$$

In particular, we are interested in the strong-coupling limit (|J| < 1) near half-filling. Depending upon the sign of $J$, this Hamiltonian describes a ferromagnet ($J < 0$) or an antiferromagnet ($J > 0$) with holes.

We restrict our Hilbert space to states composed of unoccupied or singly occupied sites; doubly occupied sites are excluded, consistent with large $U$. A convenient set of basis states is specified by $|\sigma\rangle$, the values of the spins at every lattice site, for which we will use the symbols $+$ and $-$ for spin up and down, and 0 for empty sites. As we study here the cases in which there are zero, one, or two vacancies present in an otherwise half-filled lattice, it is convenient to label the basis states by the spatial coordinates of the vacancies. Thus, $|\{\sigma\}\rangle$ represents the case in which no vacancy is present, $|r_1,\{\sigma\}\rangle$ stands for a one-vacancy state, and $|r_1 r_2, \{\sigma\}\rangle$ is the symbol for a two-vacancy state.

For convenience, we will set $t = 1$ in the following unless it is shown explicitly. Values of the energies, $E$, and coupling constants, $J$, given below are then always expressed in units of $t$.

A. Zero- and one-vacancy states

A zero-vacancy state can be defined as

$$|\{\sigma\}\rangle = \prod_i c_{i \sigma}^{\dagger} |0\rangle,$$

(2)

with $2^{N^2}$ possibilities for $|\sigma\rangle$ on a lattice with $N^2$ lattice sites. The label $i$ refers to the location of the lattice site, where some standard sequential order of all lattice sites is assumed. The ground state $|\{\sigma\}\rangle$ is ferromagnetic (all spins aligned) for $J < 0$ and antiferromagnetic (Néel) for
We define the one-vacancy states as
\[
| r_i, \{ \sigma \} \rangle = (-)^{r_i} c_{r_i, \sigma_i} | \{ \sigma \} \rangle
\]
\[
= (-)^{r_i} c_{r_i, \sigma_i} \left( \prod_i c_{i, \sigma_i}^+ \right) | 0 \rangle ,
\]
where the state obtained is independent of \( \sigma_i \). The phase factor \((-)^r\) is a "checker-board" phase, which changes sign upon moving to any nearest-neighbor lattice point. The absolute phase is clearly arbitrary, and we choose it so that the phase factor for \( r=(0,0) \) is 1. A general wave function in the space of one vacancy states can then be written as
\[
| \Psi_1 \rangle = \sum_{r_i, \{ \sigma \}} \Psi(r_i, \{ \sigma \}) | r_i, \{ \sigma \} \rangle .
\]

In the ferromagnetic case, the Hamiltonian is simply diagonalized in the one-vacancy sector. The spins at all sites except the vacancy are aligned ferromagnetically, and the exact energies, relative to the no-vacancy ground state, are
\[
e_k = -4J - 2(\cos k_x + \cos k_y) ,
\]
\[
k_{x,y} = \frac{2\pi n_{x,y}}{N} ,
\]
\[
n_{x,y} = 0, \pm 1, \pm 2, \ldots (N/2-1), N/2 ,
\]
where \( N \) is the lattice size.

In the antiferromagnetic case, even in the \( J \to 0 \) limit, the one-vacancy states are nontrivial, because a vacancy can scramble the antiferromagnetic spin configuration as it moves through the lattice. Working in the \( J \to 0 \) limit, Brinkman and Rice\(^{11}\) found that the band is narrowed relative to that given by \( \delta \). In an approximation that excludes vacancy paths with closed loops, they find that the band edge is raised from its ferromagnetic value of \( E = -4 \) to \( E = -2\sqrt{3} \) and speculate on the existence of localized states below this energy.

From the anticommutation relations of the operators \( c_{i, \sigma} \), it is straightforward to derive the following matrix elements:
\[
\langle \{ \sigma' \} | \{ \sigma \} \rangle = \delta_{\sigma \sigma'} ,
\]
\[
\langle r_i', \{ \sigma' \} | r_i, \{ \sigma \} \rangle = \delta_{\sigma \sigma'} \delta_{r_{i'} r_i} ,
\]
\[
\langle r_i', \{ \sigma' \} | H | r_i, \{ \sigma \} \rangle = V(\sigma) \delta_{\sigma \sigma'} \delta_{r_{i'} r_i} ,
\]
\[
\langle r_i', \{ \sigma' \} | \mathcal{V} | r_i, \{ \sigma \} \rangle = -i \delta_{\sigma \sigma'} + \delta_{r_{i'} r_i} ,
\]
where \( \mathcal{H} \) and \( \mathcal{V} \) are the operators defined Eq. (1) and \( n \) is a lattice unit vector. The symbol \( \sigma' + n \) refers to the spin configuration which results by taking the ensemble of spins \( \{ \sigma' \} \) and exchanging the values of the spins at \( r_i \) and \( r_i' + n \) for each other. The \( \delta \) functions in \( r \) in these equations are already implicitly contained in the \( \delta \) functions in \( \sigma \), but we have included them here for clarity of notation. Thus, our basis provides a natural second-quantized language for discussing the one-vacancy sector.

### B. Two-vacancy states

A suitable basis for the two-vacancy sector are states of the form
\[
| r_{i, j}, \{ \sigma \} \rangle = (-)^{r_{i, j}^1 + r_{i, j}^2} c_{r_{i, j}^1, \sigma_{r_{i, j}^1}} c_{r_{i, j}^2, \sigma_{r_{i, j}^2}} | \{ \sigma \} \rangle
\]
\[
= (-)^{r_{i, j}^1 + r_{i, j}^2} c_{r_{i, j}^1, \sigma_{r_{i, j}^1}} c_{r_{i, j}^2, \sigma_{r_{i, j}^2}} \left( \prod_i c_{i, \sigma}^+ \right) | 0 \rangle ,
\]
which is antisymmetric under exchange of the two vacancies
\[
\langle r_{i, j} | r_{i, j} \{ \sigma \} \rangle = - \langle r_{i, j} | r_{i, j} \{ \sigma \} \rangle .
\]
As in the one-vacancy case, these states are independent of the "dummy" spins at the vacancy sites. A general two-vacancy wave function can be written as
\[
| \Psi_2 \rangle = \sum_{r_{i, j}, \{ \sigma \}} \Psi(r_{i, j}, \{ \sigma \}) | r_{i, j}, \{ \sigma \} \rangle ,
\]
where the antisymmetry of the basis states implies the antisymmetry of the general two-vacancy wave function \( | \Psi_2 \rangle \).

The matrix elements of operators within these two-vacancy basis states can be calculated in analogy to Eq. (6). However, the fermion nature of the electrons results in additional exchange terms. For example, the scalar product of two basis states is given by
\[
\langle r_{i, j}, \{ \sigma' \} | r_{i, j}, \{ \sigma \} \rangle = \delta_{\sigma \sigma'} (\delta_{i, i'} \delta_{j, j'} - \delta_{i, j'} \delta_{i', j}) .
\]
Thus, our basis states provide a second-quantized language for treating the two-vacancy problem. Their generalization to multivacancy situations is straightforward.

### III. MONTE CARLO PROCEDURE

The thermodynamic average of an observable \( \Omega \) is given in the canonical ensemble by
\[
\langle \Omega \rangle = \frac{\text{tr} \left( \frac{\Omega e^{-\beta \mathcal{H}}}{\text{tr}(e^{-\beta \mathcal{H}})} \right)}{\text{tr}(e^{-\beta \mathcal{H}})} ,
\]
where \( \mathcal{H} \) is the Hamiltonian of the system as defined in Eq. (1), \( \beta = 1/kT \) is the inverse temperature, and the trace extends over all states with a given number of vacancies (spin configurations and vacancy locations). However, as we are interested only in the ground states for various numbers of vacancies, we replace the trace in Eq. (10) by an expectation value in a suitable trial state \( | \Psi_T \rangle \),
\[
\langle \Omega \rangle = \frac{\langle \Psi_T | e^{-\beta \mathcal{H}} / \Omega e^{-\beta \mathcal{H}} / | \Psi_T \rangle}{\langle \Psi_T | e^{-\beta \mathcal{H}} | \Psi_T \rangle} ,
\]
in which we specify the initial locations of the vacancies and the spin configuration of the lattice. As \( \beta \) becomes large, \( \langle \Omega \rangle \) approaches the ground-state expectation value, as long as \( | \Psi_T \rangle \) is not orthogonal to the true ground state. The choice of these trial states in the one- and two-vacancy cases will be discussed below.
A. General method

We evaluate the expectation values in Eq. (11) by discretizing the imaginary time interval \([0, \beta]\) with small slices of width \(\Delta \beta\). Upon introducing complete sets of states at each time slice, the sums over intermediate states are carried out via Monte Carlo techniques. We found that a 20x20 lattice with periodic boundary conditions was sufficient for the studies presented here, and that up to 640 time slices were required for adequate accuracy.

To evaluate the propagation of the system over the small time interval \(\Delta \beta\), we break up the Hamiltonian into operators whose matrix elements are convenient to evaluate. Such a breakup of \(\mathcal{H}\) into plaquettes is well known.\(^{12}\) We have chosen instead to split the Hamiltonian into

\[
\mathcal{H} = \mathcal{H}_x + \frac{1}{2} \mathcal{V} + (\mathcal{H}_y + \frac{1}{2} \mathcal{V}).
\]

(12)

Since we use basis states in which \(\mathcal{V}\) is diagonal, we consider only the kinetic energy operator for the moment and note that

\[
e^{-\Delta \beta \Omega_1 + \Omega_2} = e^{-\Delta \beta \Omega_1} e^{-\Delta \beta \Omega_2} [1 + O(\Delta \beta^2)],
\]

(13)

which is valid for general operators \(\Omega_1\) and \(\Omega_2\) with \([\Omega_1, \Omega_2] \neq 0\).

If all spins on the lattice were aligned (ferromagnetic configuration), the kinetic energy operators simply would move vacancies around in a straightforward way. Then \(\mathcal{H}_x\) and \(\mathcal{H}_y\) commute, the factorization of the exponential is exact, and we obtain

\[
e^{-\Delta \beta \mathcal{H}_x + \mathcal{H}_y} = e^{-\Delta \beta \mathcal{H}_x} e^{-\Delta \beta \mathcal{H}_y}.
\]

(14)

However, in general (and, in particular, for an antiferromagnetic lattice) the kinetic energy operators must be “protected” by projectors ensuring that no-spin world lines are inadvertently disconnected as a vacancy moves. The general Eq. (13) then applies, and we will have to extrapolate our results to \(\Delta \beta = 0\).

We now insert \(2l - 1\) (\(l \Delta \beta = \beta\)) complete sets of basis states into the expectation values of Eq. (11) and obtain

\[
\langle \Omega \rangle = \sum_{i_1, \ldots, i_{2l-1}} \langle \Psi_T | e^{-\Delta \beta \mathcal{H}_x} | i_1 \rangle \cdots \langle \Psi_T | e^{-\Delta \beta \mathcal{H}_y} | i_{2l-1} \rangle e^{-\Delta \beta \mathcal{H}_y} | \Psi_T \rangle
\]

\[
\sum_{i_1, \ldots, i_{2l-1}} \langle i_1 | e^{-\Delta \beta \mathcal{H}_x} | i_{2l-1} \rangle \cdot P(i_1, \ldots, i_{2l-1}),
\]

(15)

where \(P(i_1, \ldots, i_{2l-1})\) is the probability for the configuration \(\{i_1, \ldots, i_{2l-1}\}\),

\[
P(i_1, \ldots, i_{2l-1}) = \sum_{i_1, \ldots, i_{2l-1}} \langle \Psi_T | e^{-\Delta \beta \mathcal{H}_x} | i_1 \rangle \cdots \langle \Psi_T | e^{-\Delta \beta \mathcal{H}_y} | \Psi_T \rangle.
\]

(16)

We sample histories of the system according to this probability distribution in our Monte Carlo procedure.

B. One vacancy

Consider two one-vacancy states \(| \sigma \rangle\) and \(| \sigma' \rangle\). We will omit reference to the spin configurations from now on, but note that as we need consider, for each time slice, only vacancies moving in one lattice direction, the spin configuration changes in a “natural” way. Then

\[
\langle x' | e^{-\Delta \beta \mathcal{H}_x} | x \rangle = \delta_{x'x} e^{-\Delta \beta \mathcal{H}_x}.
\]

(17)

Inserting a complete set of momentum eigenstates of \(\mathcal{H}_x\), we obtain

\[
\langle x' | e^{-\Delta \beta \mathcal{H}_x} | x \rangle = \sum_{k_x} e^{ik_x(x-x')} e^{\Delta \beta k_x x}.
\]

(18)

Consider now the product in the numerator of Eq. (16). In accord with the usual Metropolis sampling strategy, we propose a trial change in the path, say \(| \sigma' \rangle\), to \(| \rho + \mu \rangle\) where \(\rho\) is a lattice unit vector (compare Fig. 1).

\[
\langle r_{l-1} | e^{-\Delta \beta \mathcal{H}_x} | r_{l-1} \rangle \langle r_{l} | e^{-\Delta \beta \mathcal{H}_y} | r_{l+1} \rangle \langle r_{l+1} | e^{-\Delta \beta \mathcal{H}_y} | r_{l+2} \rangle
\]

\[
\rightarrow \langle r_{l-1} | e^{-\Delta \beta \mathcal{H}_x} | r_{l} + \mu \rangle \langle r_{l} + \mu | e^{-\Delta \beta \mathcal{H}_y} | r_{l+1} + \mu \rangle \langle r_{l+1} + \mu | e^{-\Delta \beta \mathcal{H}_y} | r_{l+2} \rangle.
\]

(19)

Upon comparing the first part of this product to Eq. (17), we see that \(y_{l-1} = y_{l}\) and so \(\mu = \mu' \mathcal{X} (\mu = \pm 1)\). Because the
second part of the product requires $\delta_{x_i, x_{i+1}}$ we have to move $|x_{i+1}\rangle$ by the same vector $\mu$. We then compute the ratio

$$w = \frac{\langle r_{i-1} | e^{-\Delta \beta H_x} | r_i + \mu \rangle \langle r_i + \mu | e^{-\Delta \beta H_x} | r_{i+1} + \mu \rangle \langle r_{i+1} + \mu | e^{-\Delta \beta H_x} | r_{i+2} \rangle}{\langle r_{i-1} | e^{-\Delta \beta H_x} | r_i \rangle \langle r_i | e^{-\Delta \beta H_x} | r_{i+1} \rangle \langle r_{i+1} | e^{-\Delta \beta H_x} | r_{i+2} \rangle}$$

$$= \frac{g(\Delta \beta, x_i - x_{i-1} + \mu) g(\Delta \beta, x_{i+1} - x_i + \mu)}{g(\Delta \beta, x_i - x_{i-1}) g(\Delta \beta, x_{i+2} - x_{i+1})}.$$  

(20)

We then use the heat-bath algorithm and accept the proposed configuration with probability

$$p = \frac{w}{1 + w}.$$  

(21)

In the case of a trial state involving vacancies embedded in an antiferromagnetic background, we must exclude vacancy paths that scramble the final spin configuration of the lattice. A simple example of such a forbidden path cited by Brinkman and Rice is when a vacancy travels once around a lattice plaquette. By simply checking that no-spin world lines are disconnected by the trial paths we propose, we can also exclude these paths, as demonstrated in Fig. 2.

Figure 2(a) shows the "once-around-the-plaquette" path must be excluded as it scrambles the spin configuration of the lattice. Indeed, the spin configuration of the lowest time slice differs from that of the highest. In our method of evolving the paths of vacancies the lowest and the highest time slice always retain their spin configurations. In Fig. 2(b) the transition from the first column to the second column represents an allowed move. However, the transition from the second to the third column represents a forbidden move, which is equivalent to the vacancy traveling once around the square plaquette as depicted in Fig. 2(a). Such a forbidden path carries the signature that at least one spin world line is disconnected. This disconnection can be seen in going from the lower right corner of the second time slice from the bottom in column three to the same location in the third time slice from the bottom. Here the spin changes sign from + to −, and we have indicated the disconnected spin world line by a dotted line in Fig. 2.

The inclusion of the potential of Eq. (1) is easy using the breakup of the Hamiltonian given in Eq. (12). Since we work in a basis in which $\mathcal{V}$ is diagonal, we always have

$$\exp \left[ -\frac{\Delta \beta \mathcal{V}}{2} \right] |\tau\rangle = \exp \left[ -\frac{\Delta \beta V(\sigma(\tau))}{2} \right] |\tau\rangle,$$  

(22)

where $\exp[-\Delta \beta V(\sigma(\tau))/2]$ is a $c$ number. This implies that Eq. (20) must be modified to

$$w = \frac{g(\Delta \beta, x_i - x_{i-1} + \mu) g(\Delta \beta, x_{i+1} - x_i + \mu)}{g(\Delta \beta, x_i - x_{i-1}) g(\Delta \beta, x_{i+2} - x_{i+1})} \times \exp \left[ -\frac{\Delta \beta}{2} [V(\sigma(\tau_i + \mu)) + V(\sigma(\tau_{i+1} + \mu)) - V(\sigma(\tau_i)) - V(\sigma(\tau_{i+1}))] \right].$$  

(23)

![FIG. 1. Illustration of the Monte Carlo method used to solve the path integral problem. The solid line represents a path of a vacancy, where the position of the vacancy at each time slice $x_i$ is given by the squares. Also indicated is a proposed new path resulting from moving the vacancy at time slice $x_i$ by $\mu$ (dashed line).](image1)

![FIG. 2. (a) Example of a forbidden path of a vacancy in the approximation of Brinkman and Rice. The spin configuration of the plaquette in the lowest time slice is scrambled with respect to the highest. (b) The same path (column 3) in our numerical simulation. Here the signature of a forbidden path is the disconnected spin world line between the second and third time slice from the bottom in this column (indicated by the dotted line). The transition from the path of the vacancy in column one to the path in column two is an allowed one, whereas the transition from column two to three is forbidden.](image2)
Using the probabilities $P$ from Eq. (16) we can now compute observables. In general this will involve commutators $[\Omega, \exp(-\Delta \beta \mathcal{H})]$ that vanish like $O(\Delta \beta)$ in the case that $[\Omega, \mathcal{H}]$ is nonzero. We will therefore have to extrapolate linearly to $\Delta \beta = 0$ for cases where these commutators do not vanish.

\[
\langle r_i | \mathcal{H}_x e^{-\Delta \beta \mathcal{H}_x} | r_{i+1} \rangle = -t [\langle r_i+1 | e^{-\Delta \beta \mathcal{H}_x} | r_{i+1} \rangle + \langle r_i-1 | e^{-\Delta \beta \mathcal{H}_x} | r_{i+1} \rangle] \\
= -t [g(\Delta \beta, x_i - x_{i+1} + 1) + g(\Delta \beta, x_i - x_{i+1} - 1)] \delta_{r_{i+1}.} \tag{24}
\]

C. Two vacancies

Our algorithm for sampling histories of a two-vacancy system is analogous to that we have described for the one-vacancy system. Minor complications arise from the facts that two vacancies cannot be at the same site at the same time and that we have excluded doubly occupied sites from our Hilbert space. The latter result in correction terms to the kinetic energy

\[
\langle r_i | \mathcal{H}_x e^{-\Delta \beta \mathcal{H}_x} | r_{i+1} \rangle = -t [\langle r_i+1 | e^{-\Delta \beta \mathcal{H}_x} | r_{i+1} \rangle + \langle r_i-1 | e^{-\Delta \beta \mathcal{H}_x} | r_{i+1} \rangle] \\
= -t [g(\Delta \beta, x_i - x_{i+1} + 1)(1-\delta_+) + g(\Delta \beta, x_i - x_{i+1} - 1)(1-\delta_-)] \delta_{r_{i+1}.} \tag{25}
\]

where $\delta_+$ is 1 if there is another vacancy at the lattice site $(x_i+1, y_j)$ and 0 otherwise, and $\delta_-$ is 1 if there is another vacancy at the lattice site $(x_i-1, y_j)$ and 0 otherwise. An additional complication is that imposed by the antisymmetry constraint discussed in connection with Eq. (9); we deal with this in the course of discussing our results below.

IV. RESULTS

A. Numerical tests

One simple check of our algorithm is to simulate one vacancy on a ferromagnetic lattice. Here the single-particle energies are given by Eq. (5). The expectation value of the energy is

\[
E = \langle \mathcal{H} \rangle = \sum_k e^{-\beta e_k} e_k \\
= -4J - 2 \sum_k e^{2i\beta \cos k} \cos k \tag{26}
\]

In Fig. 3 we display the exact result of Eq. (26) as a function of $\beta$ (solid line). The plot symbols represent the result of our Monte Carlo calculations. For every data point three runs of the program using 10,000 updates of the path were performed. The results of the Monte Carlo simulation agree completely with the exact calculation within the statistical errors; this is hardly a surprise. Our factorization is exact for all values of $\Delta \beta$ in the case of a ferromagnetic lattice; i.e., Eq. (14) holds. To generate

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig3}
\caption{Energy of the one-vacancy state in a ferromagnetic lattice as a function of $\beta$. The solid line represents the exact analytic result, whereas the results of our Monte Carlo calculation are shown by the plot symbols.}
\end{figure}
culated $E$ for $\beta=8$ and different values of $\Delta \beta$. Since the
density of states varies like $\rho(\omega) \propto \sqrt{\omega - \omega_0}$ near the band
edge, we would expect a value of

$$E(\beta=8) \approx -2\sqrt{3} + \frac{3}{2\beta} \approx -3.28 .$$  (27)

This value is represented by the dashed line in Fig. 4, while the points are the results of our calculations. For
every value of $\Delta \beta$ we performed five independent runs
with different seeds of the random number generator and
50,000 updates of each time slice of the path. Our results
approach the exact value for $\Delta \beta \to 0$. A linear $\chi^2$ fit and
extrapolation to $\Delta \beta=0$ yields the value $E_{\text{calc}}(\beta=8) = -3.25 \pm 0.07$, accurate within the uncertainty. The
addition of more data points with smaller $\Delta \beta$ would not
improve the accuracy of the final result since the statistical
errors associated with the individual data points increase
with decreasing $\Delta \beta$. But even the value for $\Delta \beta=0.1$ is
wrong by only 0.1, which will turn out to be acceptable for
most applications we deal with in this work. We have
therefore used this value of $\Delta \beta$ for the following calculations with $J \neq 0$.

B. One vacancy, $J \neq 0$

Figure 5 shows our results for $E$ as a function of $\beta$ for
different values of $J$. From top to bottom the $J$ values
used are 0.6, 0.2, 0.06, and 0. In all cases the statistical
errors are smaller than the plot symbols used. In all four
cases calculated the curves approach a constant value for
large $\beta$. As expected, we find that $E$ increases with in-
creasing $J$, as it becomes energetically more costly for the
vacancy to hop to another site; the vacancy is bound to the
origin by a string of overturned bonds.

The functional dependence of $E$ on $J$ is of some
interest. Shraiman and Siggia\textsuperscript{10} have recently proposed that
$E(J')=E(0)+2.74J'^{1/3}$, where their coupling constant,
which we call $J'$, is $4J$. We have calculated $E(J)$ at

![FIG. 5. Energy, $E$, as a function of $\beta$ for different values of $J$ in the case of the one-vacancy state.](image)

a finite value of $\beta=8$. The saturation of the curves in
Fig. 5 shows that this is a reasonable approximation to
$E(J, \beta \to \infty)$. The results of our calculations are
displayed in Fig. 6. The data points are represented by
the plot symbols, and the solid straight line corresponds
to a linear regression fit

$$E(J)-E(0)=aJ^{2/3}+b .$$  (28)

We extract from our data the values $a=3.15$ and
$b=0.06$ with a correlation coefficient $\kappa=0.992$. If this is
expressed in terms of $J'$, we obtain a coefficient of $J'^{2/3}$ of
1.25, or less than half of that found by Shraiman and Siggia.
We attribute this reduction to self-intersections and
juxtapositions of the vacancy path, as already hypothesized in Ref. 10.
C. Two vacancies

We now turn to the two-vacancy sector for antiferromagnetic couplings. Because of possible connections with the newly discovered high-temperature superconductors, the binding of two vacancies is the most interesting question we can address in this sector. A simple picture leads to the expectation that two vacancies will indeed bind. As one of the vacancies travels through an antiferromagnetic lattice, it leaves behind it a string of energetically unfavorable ferromagnetic bonds, which binds it to the origin. If the second vacancy follows behind the first one, it will "heal" the bonds. The pair would thus be bound together and would not be localized. However, naive expectations that this scenario automatically leads to binding can be countered with a one-dimensional example. It is therefore necessary to perform careful calculations to address this issue.

To investigate the binding of two vacancies, we compare the expectation values of the energy, Eq. (11) with $\Omega=\mathcal{F}$, calculated with different trial states. One of these is our "reference state," consisting of two widely separated vacancies in a Néel background of spins. As long as their separation is large enough and $\beta$ is finite, it is clear that these two vacancies will never "find each other" to bind, so that the energy found for this state (relative to the no-vacancy Néel state), will simply be twice the energy of a one-vacancy state. We have verified that this is so in our calculations.

The other trial states we use in our calculations have the two vacancies neighboring each other in a Néel background of spins. If one of the vacancies is at the origin [lattice coordinates (0,0)], then the other vacancy is at the neighboring site (1,0) in our "s-wave" trial state (denoted by $S$), and we impose no restrictions on the paths, other than those described in the previous section. Our first "p-wave" trial state (denoted by $P_1$) corresponds to this same initial configuration, but we impose the additional requirement that there be a node in the many body wave function [Eq. (8)] along the line $x_1=x_2$. Practically, this is done by simply forbidding Monte Carlo moves that would cause the vacancy paths to have the same x coordinate at any particular time. Our second "p-wave" trial state ($P_2$) has the second vacancy at the site (1,1), and we impose a nodal surface along the line $x_1-x_2=y_1-y_2$. Finally, our "d-wave" trial state ($D$) has the same initial configuration of vacancies as $P_2$, with nodes imposed both along $x_1=x_2$ and along $y_1=y_2$.

In view of the requirement that the two-vacancy wave function be antisymmetric under the interchange of the two vacancies, we regard only the $P_1$ and $P_2$ trial states as physical. However, as we do not allow for the most general nodal structure of the wave function, our results for the energies of these state, and for the $D$ state as well, can only be regarded as variational upper bound; i.e., the binding could be greater than what we calculate.

All of our two-vacancy calculations have been performed with $\Delta \beta=0.1$. To check the validity of this approximation, we performed a series of calculations analogous to that used in Fig. 4. For a constant value of $\beta=8$ and a constant value of $J=0.6$ we varied $\Delta \beta$. The upper panel of Fig. 7 shows the results of these tests. The upper curve represents the energy calculated with our reference trial state ($R$), while the lower curve shows the results obtained with the $S$ trial state. The difference of the two expectation values, which we identify as the binding energy, $\epsilon$, is displayed in the lower panel of Fig. 7. From these results, it appears reasonable to fix $\Delta \beta=0.1$, since the binding energy we find for this value of $\Delta \beta$ lies com-
one vacancy states when $J=0$, we find an energy in agreement with the analytic approximation of Brinkmann and Rice,\textsuperscript{11} while for $0 < J \leq 1$, we find an energy varying as $J^{2/3}$. This latter is in qualitative agreement with the results of Siggia and Shraiman,\textsuperscript{10} although our numerical coefficient is less than half of theirs, presumably because of their neglect of self-intersecting and juxtaposed paths.

For two vacancies, we find bound states with both $S$ and $P$ spatial symmetries, but no binding in states with a $D$ symmetry. The binding we find is "strong" for the $S$ state in the sense that the binding energy scales as $J^{2/3}$. For the $P$ spatial symmetry we find binding proportional to $J$. These results are in qualitative agreement with those of Ref. 10 and in contrast to those of Ref. 9. The former work finds strong binding in $S$ states (although again with a larger numerical coefficient—our $S$ state is to be identified with the boson state of Ref. 10), and weak binding ($\propto J$) in $P$ and $D$ states. The later work finds no binding at all.

As neither of Refs. 10 or 9 is sufficiently detailed for us to make a thorough analysis of their calculations, we can only speculate as to the sources of our discrepancies with them and their discrepancies with each other. The limited basis of Ref. 9 allows the vacancy to be at most three hops away from its initial site, although it does allow the vacancy to travel by translations of the computational unit cell. This implies that the vacancy travels through the lattice with a very localized disturbance of the Néel background. In contrast, our calculations show the vacancy often five or more hops away from its starting site with a substantial scrambling of the Néel background. The method of Ref. 10 allows paths with an arbitrary number of hops, but ignores self-intersections and juxtapositions of the paths, which is probably the reason why these authors obtain numerical coefficients for their binding energies different from ours.

Our calculations presented here can be extended in two interesting ways. One is to consider multi-vacancy states. As the fermion nature of only the vacancies need be treated explicitly, existing Monte Carlo methods tractable for few-fermion systems might be applied; a calculation on a $20 \times 20$ lattice with 5% vacancies appears quite feasible.

A second extension is to include spin dynamics. It is likely that the ability of the spins to relax would weaken or destroy the "string" confining a single vacancy and so increase its mobility. The interaction of two vacancies might similarly weaken, although spin fluctuations might also bind two vacancies together.\textsuperscript{14} The ideal simulation addressing these ideas would be a spin-$\frac{1}{2}$ Heisenberg system with vacancies. Although the 2D antiferromagnetic Heisenberg model without vacancies can be simulated,\textsuperscript{15} we know of no scheme that allows vacancies to be incorporated without encountering the "negative probabilities" that often plague Monte Carlo calculations. However, a tractable alternative is to promote our static Ising spins to the dynamical ones of the $O(3)$ nonlinear $\sigma$ model. Such a simulation would reveal the influence of spin waves on our present results and also allow study of the importance of topologically nontrivial spin textures and their interaction with vacancies.\textsuperscript{16}

V. DISCUSSION

We have presented Monte Carlo simulations of one and two vacancies in a 2D Ising antiferromagnet. For
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13Below the value of $E$ given here, Brinkman and Rice find "localized" states which extend in energy to $E = -4$. However, these states are only accessible through at least triple loops around a plaquette. For the parameters considered here, we did not find these states in our Monte Carlo samples.