Spin liquid ground state in a two dimensional non-frustrated spin model

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We consider an exchange model describing two isotropic spin-1/2 Heisenberg antiferromagnets coupled by a quartic term on the square lattice. The model is relevant for systems with orbital degeneracy and strong electron-vibron coupling in the large Hubbard repulsion limit, and is known to show a spin-Peierls-like dimerization in one dimension. In two dimensions we calculate energy gaps, susceptibilities, and correlation functions with a Green’s Function Monte Carlo. We find a finite spin gap and no evidence of any kind of order. We conclude that the ground state is, most likely, a spin liquid of resonating valence bonds.

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The existence of a homogeneous resonating valence bond (RVB) ground state (GS) for two-dimensional (2D) spin-1/2 systems has been a subject of intense theoretical study in the past two decades. The idea that the reduced dimensionality and the small spin value might enhance quantum fluctuations up to the point of destroying the classical Néel antiferromagnetic order was first put forward by Anderson in 1973.¹

The subject became a hot topic after Anderson suggested that the parent compounds of the high-$T_c$ copper-oxides – argued to be well described by a S=1/2 Heisenberg model on a square lattice – might have a spin liquid GS, and that superconductivity would result from doping such a spin liquid.

Since then, many studies on the S=1/2 2D Heisenberg model have shown that reduced dimensionality and low spin are not sufficient to stabilize a RVB GS. First, it is now well established that the square lattice case is Néel ordered.² More surprisingly, three-sublattice Néel order is also likely to survive on the triangular lattice, where the model is frustrated.³⁴ A stronger lattice frustration – like in the kagomé lattice –, multiple-spin exchange terms on the triangular lattice,⁵ or frustration due to the coupling constants – like in the $J_1-J_2$ Heisenberg model on the square lattice –⁶⁷ might be more effective in stabilizing a spin liquid GS. The evidences for a triplet gap and the definite assessment about the liquid nature of the GS are, however, either limited to small lattice degeneracy and strong electron-vibron coupling in the large Hubbard repulsion limit, and is known to show a spin-Peierls-like dimerization in one dimension. In two dimensions we calculate energy gaps, susceptibilities, and correlation functions with a Green’s Function Monte Carlo. We find a finite spin gap and no evidence of any kind of order. We conclude that the ground state is, most likely, a spin liquid of resonating valence bonds.

where $J > 0$, the summation runs over the nearest-neighbor (nn) sites of a square lattice, $X_{i,j}^{(S)} = (2S_i \cdot S_j - 1/2)$, and $X_{i,j}^{(T)} = (2T_i \cdot T_j - 1/2)$. $H_{ST}$, henceforth referred to as ST model, describes for $J > 0$ the low-energy physics of an insulating crystal with one electron per site in a two-fold degenerate orbital, in the limit of large on-site repulsion (Mott insulator) and in presence of Jahn-Teller (JT) effect.⁸ The derivation of Eq. (1), in the same spirit of the derivation of the Heisenberg model from the large-U Hubbard limit, is standard.⁹¹⁰¹¹ The crucial physical condition to be verified is that, among the possible two-particle states obtained upon virtual hopping, the inter-orbital singlet should turn out to be the lowest in energy, which is indeed the case when a strong dynamical JT effect is at play. A different, perhaps more common, physical situation is that the lowest two-particle intermediate state is a Hund’s triplet, in which case the exchange model leads to spin ferromagnetism.¹²¹³ In the general case, the exchange Hamiltonian contains pseudo-spin anisotropic terms, and some aspects of its phase diagram have been addressed with different techniques in Refs.¹⁴¹⁵

A class of SU(n)-invariant generalizations of the Heisenberg model has been previously proposed as candidate non-Néel antiferromagnets.¹⁶¹⁷ It turns out that the ST model is unitarily equivalent to a particular SU(4) model in that class, with $n_c = m = 1$ (notations of Ref.¹⁸). The $n \to \infty$ limit of these SU(n) models has been studied in detail, and shown to have, for $n_c = m = 1$, a spin-Peierls GS both in 2D,¹⁹ as well as in one dimension (1D).²⁰ Recently, we have verified that the spin-Peierls nature of the GS persists, in 1D, down to the ST model point, i.e., $n = 4$.²¹ The behavior in 2D is more subtle. We show in the present letter that a RVB GS is found for $n = 4$ in 2D, in marked contrast with the $n \to \infty$ limit predictions.

The relationship between the ST model and the usual Heisenberg model – $H_{Heis} = (J/2) \sum_{i,j} X_{i,j}^{(S)}$ – is made
more clear by working in the singlet subspace with the overcomplete basis set of the valence bond (VB) configurations, i.e., products of non-overlapping singlet dimers \((i,j)\) connecting pairs of sites on opposite sublattices and covering the lattice. More precisely, if \((i,j)_{ST} = (\uparrow\downarrow - \downarrow\uparrow)_{ST}\) denotes a singlet bond between sites \(i\) and \(j\) for the \(S\) (\(T\)) variables, we restrict the VB configurations considered to the invariant subspace – containing the GS, by positivity arguments – in which every singlet dimer \((i,j)\) is a product of an \(S\) and a \(T\)-singlet, \((i,j) = (i,j)_S(i,j)_T\). Since \((i,j)_{ST} = -(j,i)_{ST}\), we assume the sign convention that the leftmost index in the singlet pair always belongs to sublattice A. It is possible to show that the GS of both the ST and the Heisenberg model can be written as linear combinations with positive coefficients of VB configurations, with the previous restrictions and sign conventions. The construction of excellent variational wavefunctions based on combinations of VB configurations has been shown to be possible even for the Heisenberg case provided sufficiently long-ranged bonds are allowed. On the other hand, it is quite clear that whenever short-ranged bonds are the dominant ones, the GS will have a characteristic length \(\xi\) and there will be a finite gap, in the thermodynamic limit, to the lowest triplet excitations.

The bond operators \(X^{(S)}_{ij}, X^{(T)}_{ij}\), have simple properties when acting on VB configurations, because they affect at most two singlet pairs. Indeed, it is known that \(X^{(S)}_{ij}(i,j)_S = 2(i,j)_S\), and \(X^{(S)}_{ij}(i,j)_S(k,l)_S = -(k,j)_S(i,l)_S\). Using these rules for \(X^{(S)}_{ij}\) and \(X^{(T)}_{ij}\), it is straightforward to show that the bond operators \(X_{ij} = -X^{(S)}_{ij}X^{(T)}_{ij}\) appearing in the ST model, Eq. (1), obey the following rules:

\[
X_{ij}(i,j) = -n (i,j), \quad X_{ij}(k,l)(k,l) = -(k,j)(i,l),
\]

with \(n = 4\). Notice that these relationships are formally identical to those relevant to the Heisenberg case, except for a coefficient \(n = 4\), in place of \(n = 2\), when a \(nn\) Hamiltonian bond \((ij)\) acts on a single dimer \((i,j)\). This enhanced coefficient favors the formation of short-ranged bonds in the GS of the ST model, making the suppression of Néel long range order (LRO) more likely. In the limit \(n \rightarrow \infty\) we recover, from Eq. (2), the known results both in 1D and 2D. Indeed, for \(n \rightarrow \infty\) the only surviving VB configurations are those with \(nn\) dimers only. In 1D this leads to a doubly-degenerate spin-Peierls GS. In 2D the model maps onto the purely kinetic limit of the quantum dimer model (QDM) of Rokhsar and Kivelson. (This purely kinetic point of the QDM is believed to be characterized by a plaquette resonating VB state, breaking translational invariance.)

By working with the VB basis, the action of the Hamiltonian \(H_\psi\) on any basis element \(\psi_i\) defines a non-symmetric matrix, \(H_\psi = \sum h_{ji} \psi_i\) with all elements \(h_{ji}\) non-positive, as implied by Eq. (3). The right eigenvector of \(h_{ji}\) with minimum eigenvalue, corresponding for \(h_{ji} \leq 0\) to the GS of \(H\), can be computed by applying the power method, as implemented stochastically by means of the Green’s Function Monte Carlo (GFMC) method. The GFMC is in fact not limited to symmetric matrices, and there is no sign problem when all \(h_{ji}\) are non-positive. Using the VB basis, the GFMC turns out to have extremely small statistical errors, compared to the more conventional algorithm employing an Ising basis. In this formulation, the GFMC does not require the calculation of the overlaps \(\langle \psi_i | \psi_j \rangle\) between VB configurations. Details of the method are given elsewhere.

This new and simple GFMC allows us to obtain a very accurate determination of the triplet gap by performing two independent simulations of the singlet GS and the triplet lowest excited state. In the latter case, the basis employed is slightly modified, by allowing for the presence of a single triplet bond \((i,j)^T = (\uparrow\downarrow + \downarrow\uparrow)_{ST}(i,j)_{ST}\) in each element of the VB basis. The corresponding rules for the application of \(X_{ij}\) are: \(X_{ij}(i,j)^T = 0\), and \(X_{jk}(i,j)^T(k,l) = -(k,j)(i,l)^T\). Notice that this implies the absence of sign problem in the triplet subspace as well.

Fig. 1 shows the results obtained for the triplet gap for \(L \times L\) square lattices with \(L\) up to 24, as a function of the inverse volume. The corresponding data for the Heisenberg case, also shown for comparison, are consistent with a finite size gap \(\Delta_L\) scaling to zero as \(a/L^2 + b/L^3 + \cdots\). The dashed lines (see also left inset in Fig. 1) show our best two-parameter fit to the ST data obtained by imposing the same gap behavior as in the Heisenberg case: such a fit is clearly unsatisfactory. Instead, the solid line through the ST data is the result of a three parameter fit of the form \(\Delta_L = \Delta + a/L^2 + b/L^3 + \cdots\) giving a clear evidence of a finite gap \(\Delta\) in the thermodynamic limit. Clearly, the detailed finite-size behavior of the ST gap in 2D is non-trivial, requiring the simulation of quite large lattices (\(L = 24\)) to pin-down the presence of a gap. This suggests the presence of a length scale \(\xi\) of the order of \(10/20\) lattice spacings. This behavior should be contrasted to that of the triplet gap for the ST model in 1D, shown in the right inset of Fig. 1, where the size scaling of the gap is straightforward (\(\xi \approx 1\)).

In principle, either a VB crystal with some broken spatial symmetry, as in 1D, or a homogeneous spin liquid is compatible with the existence of a spin gap. In order to investigate the possible kinds of LRO which might characterize the GS of the ST model in 2D, we have calculated the expectation value of several spin-spin correlation functions by the forward walking technique.
tor $S(\pi, \pi) = \sum_x (-1)^x \langle S_0 \cdot S_x \rangle$, which should diverge with the system volume $L^2$, when the Néel order parameter $m$ is non-vanishing, $S(\pi, \pi) \sim (L m)^2$. The comparison between the Heisenberg and the ST model results for $S(\pi, \pi)$ strongly suggests the absence of Néel order in the ST case, in agreement with the presence of a triplet gap. More interestingly, the inset in Fig. 2 shows the results obtained, in the ST case, for the dimer-dimer structure factors with nn bonds in the $x$-direction, $S^{d-d}(q) = \sum_x e^{i x \cdot r} \langle S_0 \cdot S_{x+r} \rangle$. The $q = (\pi, \pi)$ and $q = (0, \pi)$ dimer structure factors are clearly finite in the $L \to \infty$ limit; the $q = (\pi, 0)$ also does not seem to diverge linearly with the volume, but the present data are limited to too small sizes ($L \leq 10$) to be conclusive.

In order to have further evidence about the existence of a true homogeneous spin liquid GS, we have directly calculated, using the new more efficient GFMC algorithm described previously, the response of the system to symmetry-breaking operators. It is practically impossible to exclude all possible types of crystalline order numerically; in the following we restrict our consideration to the most plausible types of order, involving either a broken translation symmetry $T$ with momenta compatible with a real GS ($q = (\pi, 0), (0, \pi), (\pi, \pi)$), or/and a broken $\pi/2$ rotation symmetry $R$. This includes all types of crystalline dimer and plaquette order proposed so far. More precisely, we have perturbed the ST Hamiltonian by adding a term $\alpha T$, with $T$ an operator which breaks one of the symmetries above: $T = \sum_x \sum_y e^{i x \cdot r} X_{x,y}$ with the appropriate $q$ for the translation case, and $T = \sum_x (X_{x,y} - X_{x,\pi+y})$ for the $\pi/2$ rotation. On finite size, the GS expectation value of $T$ vanishes by symmetry, and the GS energy per site has corrections proportional to $\alpha^2$, $\epsilon_0 = \epsilon_0 - \chi_0$ being the generalized susceptibility associated to the symmetry-breaking operator $T$. On the other hand, if symmetry breaking occurs in the thermodynamic limit, it is possible to have $\lim_{L \to \infty} \lim_{\alpha \to 0} \langle T/L^2 \rangle = 0$. More precisely, it is possible to show that $\chi_0$ is bounded from below by the order parameter times the system volume squared, $\chi_0 > \text{const} \cdot p^2 L^4$. These susceptibilities are therefore a very sensitive tool – much more than the usual square of the order parameter – for detecting LRO. For instance, as shown in Fig. 3 for the ST model in 1D, the presence of dimerization in the thermodynamic limit is readily inferred from the behavior of $\chi_0$, with $\chi_0 = \sum_x (-1)^x \langle X_{x+1} \rangle$, even for very small system sizes ($L \leq 12$). Fig. 3 shows the results obtained for the ST model in 2D. For all the symmetry breaking operators considered, the associated susceptibilities are found to be finite. The largest susceptibility is found to be the one associated to columnar order ($T(\pi, 0)$ in Fig. 3), and is only weakly increasing with size, eventually saturating to a constant, in marked contrast to the strong divergence of the 1D analog (see Fig. 3).

In conclusion, we have studied a non-frustrated exchange Hamiltonian, Eq. (4), which describes the low-energy physics of a Mott insulator with orbital degeneracy in the regime in which the inter-orbital singlet is the lowest-energy intermediate state available to virtual hopping. We find a clear evidence for a spin gap without crystalline VB order in 2D. A homogeneous liquid of resonating valence bonds appears as the natural candidate GS for this model. An important question about the nature of the low-lying excitations remains to be addressed: is there a branch of gapless excitations in the singlet sector, or is the system is gapped to all excitations? The answer to this question will require future work.

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FIG. 1. Finite size gap between the GS and the first excited triplet state for the ST model and for the Heisenberg model [20] on the square lattice. The ST data are obtained by the new GFMC method working with the VB basis. The solid and dashed lines are different fits described in the text. The data for the ST model clearly indicate the existence of a spin gap in the thermodynamic limit. Left inset: The ST data plotted versus \( 1/L \). Right inset: The triplet gap for the ST in 1D.

FIG. 2. Finite size structure factors obtained by GFMC with the forward walking technique. From the spin-spin \( S(\pi, \pi) \) we find no clear sign of Néel LRO. The size scaling of the Néel order parameter for the Heisenberg model [20] is shown for comparison. Inset: The important components of the Fourier transform of the dimer-dimer correlation functions appear to be finite.

FIG. 3. The susceptibility to breaking translation symmetry for the ST model in 1D, which is known to have a spin-Peierls-like dimerization. Data are obtained by exact diagonalizations. Inset: the extrapolated \( \alpha = 0 \) value of \( \chi_O \) divided by the square of the system volume \( L^2 \).

FIG. 4. Susceptibilities to breaking the most important crystal symmetries for the ST in 2D. The finite-size, finite-\( \alpha \) data for \( \chi_O \) can be readily extrapolated, by quadratic fits, to \( \alpha = 0 \) values: notice the distinctly different behavior for the corresponding 1D results of Fig. 3. The insets summarize the results obtained for the extrapolated \( \chi_O \).