The phase diagram of the Heisenberg antiferromagnet with four-spin interactions

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Abstract
We study the quantum phase diagram of the Heisenberg planar antiferromagnet with a subset of four-spin ring exchange interactions, using the recently proposed hierarchical mean-field approach. By identifying relevant degrees of freedom, we are able to use a single variational ansatz to map the entire phase diagram of the model and uncover the nature of its various phases. It is shown that there exists a transition between a Néel state and a quantum paramagnetic phase, characterized by broken translational invariance. The non-magnetic phase preserves the lattice rotational symmetry, and has a correlated plaquette nature. Our results also suggest that this phase transition can be properly described within the Landau paradigm.

1. Introduction
Quantum phases of matter and their transitions are of fundamental concern to modern condensed matter physics [1]. Such interest is motivated not only by potential technical applications, but also on purely scientific grounds. Research in this field may lead to a deeper understanding of the fundamental working principles behind nature’s behavior, and often original new physical theories emerge. One of them, recently proposed in [2], predicts the existence of a class of systems whose critical behavior lies outside the scope of the Landau theory of phase transitions [3]. Critical points in these systems are characterized by the deconfinement of fractionalized excitations, parameterizing the original degrees of freedom, which occurs right at the transition. It was observed that this scenario can, in principle, be realized in spin systems, which exhibit a second-order phase transition point characterized by the simultaneous breakdown of continuous (e.g. spin $SU(2)$) and discrete (e.g. lattice) symmetries, in such a way that symmetry groups on opposite sides of the transition are not group–subgroup related. Such critical points cannot be described in the framework of Landau’s theory.

According to [2], there should exist a substantial number of spin systems, which exhibit deconfined critical points. For instance, frustrated two-dimensional (2D) antiferromagnets (AFs), like the $J_{1}−J_{2}$ model, are believed to fall into this category. However, there seems to be no experimental or theoretical proof of this claim. Another class of models believed to display such a behavior includes non-frustrated AFs with multi-spin exchange interactions. One such model was studied by Sandvik [4] and other authors [5, 6] and, although seemingly artificial, it provides a playground for testing new theories. Their quantum Monte Carlo (QMC) simulations claimed numerical evidence for the deconfined quantum criticality scenario.

The model studied in [4] is a Heisenberg AF with a subset of four-spin ring exchange interactions, defined on a square lattice (named the $J−Q$ model)

$$H = J \sum_{(ij)} S_i S_j - Q \sum_{(ijkl)} (S_i S_j - \frac{1}{4})(S_k S_l - \frac{1}{4}),$$

where $Q > 0$, $i,j,...$ denote sites in a 2D square lattice and $S_i$ are spin-1/2 operators. The first summation extends over bonds (nearest neighbor sites). The second term contains two sums over plaquettes (sites of the dual lattice): first, $(ij)$ and $(kl)$ denote parallel horizontal links of the plaquette, and then $(ik)$ and $(jl)$ correspond to parallel vertical bonds. It was concluded [4] that there exists a critical point at $Q_c/J \sim 25$ separating the antiferromagnetic phase from a valence-bond solid (VBS) state, whose nature is, strictly speaking, unclear [4] but the calculations suggested a columnar (dimer) order in this paramagnetic region.

In the present paper we study the phase diagram of the $J−Q$ model, using a recently proposed hierarchical mean-field (HMF) technique [7, 8]. The main idea of the method revolves around the concept of a relevant degree of freedom (a ‘quark’)—spin cluster in this particular case—which can be used to build up the system. The initial Hamiltonian is
then rewritten in terms of these coarse-grained variables and a mean-field approximation is applied to determine properties of the system. Thus, the (generally) exponentially hard problem of determining the ground state of the system is reduced to a polynomially complex one. At the same time, essential quantum correlations, which drive the physics of the problem, are captured by the local representation. In other words, provided the quark is chosen properly, even a simple single mean-field approximation, performed on these degrees of freedom, will yield the correct phase diagram. Moreover, our HMF ansatz provides an educated nodal surface that can, in principle, be used in conjunction with fixed-node (or constrained path [9]) QMC approaches to further improve correlations, and thus energy estimates, in those cases when there is a sign (phase) problem.

It is important to emphasize the simplicity of our method. In this work we concentrate on symmetries of the various phases, exhibited by the $J-Q$ model. By using a more sophisticated variational ansatz (e.g. a Jastrow-type correlated wavefunction), one can also improve numerical values of the observable quantities and phase transition points, but the physical picture will remain intact. Nevertheless, the HMF method was quite accurate to yield the quantitatively correct phase diagram [7] of the $J_1-J_2$ model, whose behavior is driven by the interplay of two gapless phases: the Néel and columnar AF states. In the $J-Q$ model the large-$Q$ phase is gapped. Due to its real-space nature, the HMF method should be appropriate for this model. Indeed, our recent studies [10] of another gapped system—the Heisenberg model on the Shastry–Sutherland lattice—support this assumption.

Our findings are summarized in figure 1. There indeed exists a non-magnetic phase, which we found to be of a correlated plaquette type, and not of a dimer character, separated from the Néel state by a second-order phase transition. Furthermore, the numerical value of the critical point, $Q_c/J \sim 2$, which we obtained, is quite different from that of [4]. Our results are consistent with data obtained from our HMF ansatz provides an educated nodal surface that can, in principle, be used in conjunction with fixed-node (or constrained path [9]) QMC approaches to further improve correlations, and thus energy estimates, in those cases when there is a sign (phase) problem.

In the next section we set up the formalism. The results are presented in section 3, followed by a discussion in section 4.

2. Coarse graining and HMF approximation

For our purposes it is convenient to separate the two- and four-spin terms in the $J-Q$ Hamiltonian:

$$H = -\frac{2NQ}{16} + \left( J + \frac{Q}{2} \right) \sum_{ij} S_i S_j - Q \sum_{ijkl} (S_i S_j)(S_k S_l).$$

A satisfactory coarse graining procedure should partition the lattice into spin clusters (quarks), containing $Q_n$ sites, that explicitly preserve symmetries of the Hamiltonian. In particular, the $J-Q$ Hamiltonian is explicitly spin-$SU(2)$ invariant. Moreover, it is invariant under transformations from the lattice rotational group $C_4$. Therefore, we will consider only symmetry preserving degrees of freedom: (i) plaquettes ($2 \times 2$ spin clusters) and (ii) $4 \times 4$ spin clusters. Each cluster state will be associated with a hard-core bosonic operator $\gamma$. These operators are Schwinger bosons of $SU(2^N)$ and must obey the local constraint: $\sum_n \gamma_n^\dagger \gamma_n = 1$. They define the hierarchical language [8] for our problem.

From the form of equation (1) it is clear [7] that the (equivalent and exact) bosonic Hamiltonian will contain not only two-body scattering processes, but also four-boson interactions. Therefore, we can write down symbolically

$$H = \sum_i (H_{2b})_{\alpha \beta} \gamma_i^\dagger \gamma_\beta + \sum_{(ij)} (H_{4b}^{\alpha \beta \gamma \delta})_{(ij)} \gamma_i^\dagger \gamma_j^\dagger \gamma_\beta \gamma_\gamma + \sum_{(ijkl)} (H_{4b}^{\alpha \beta \gamma \delta})_{(ijkl)} \gamma_i^\dagger \gamma_j^\dagger \gamma_k^\dagger \gamma_l^\dagger \gamma_\alpha \gamma_\beta \gamma_\gamma \gamma_\delta,$$

where $a, \ldots$ label states in the Hilbert space of a quark, $i, j, \ldots$ denote sites in the coarse-grained lattice, and summations are assumed over all repeated indices. The term with $H_{2b}^{2b}$ encodes two-body interactions, while the last two lines describe the correlated four-boson scattering. The superscript $h$ indicates that $i_1 i_2$ and $i_3 i_4$ are horizontal links of a plaquette, and similarly $v$ denotes the case when $i_1 i_3$ and $i_2 i_4$ are vertical links of the same plaquette.

We will investigate the phase diagram of the $J-Q$ model using the HMF approximation whose variational state assumes that the hard-core bosons form an insulating state. Further, we introduce a new set of bosonic operators, related to the old ones by a real site-independent canonical transformation:

$$\gamma_\alpha = R_0^{\alpha} \Gamma_\alpha; \quad R_0^{\alpha} R_0^{\beta} = \delta_{\alpha \beta}; \quad R_0^{\alpha} R_0^{\beta} = \delta_{\alpha \beta}$$

and write the variational ground state in the form

$$|\psi_0\rangle = \prod_i \Gamma_i^{\alpha}(0),$$

where $n = 0$ denotes the lowest energy single-particle mode (we shall also denote: $R_0^{0} \equiv R_0$, and $|0\rangle$ represents the vacuum. It is important to emphasize that although the coarse graining procedure preserves the symmetries of the Hamiltonian, some of them can be spontaneously broken at the mean-field level as a result of self-consistency. In particular, the columnar dimer state is contained in the wavefunction (3) although, as we will see below, it never appears as a stable solution.

We have explicitly separated the four-boson interaction in the Hamiltonian (2) into horizontal and vertical link contributions. This distinction is important because these two terms must be properly symmetrized to fulfill bosonic statistics. In particular, the term $H_{4b}^{v}$ has to be symmetrized only with
respect to indices in the same group, and groups as a whole (groups are separated by semicolons), i.e. one needs to take into account only the following permutations: \((1 \leftrightarrow 2), (3 \leftrightarrow 4)\) and simultaneously \((1 \leftrightarrow 3, 2 \leftrightarrow 4)\). Analogously, in the term \(H_{\text{int}}^{\Delta s}\) only the permutations \((1 \leftrightarrow 3), (2 \leftrightarrow 4)\) and \((1 \leftrightarrow 2, 3 \leftrightarrow 4)\) should be accounted for.

The problem then reduces to minimization of the energy functional:

\[
\frac{NqE_0[R]}{N} = (H_{\text{int}})_{ab} R_a^b R_a + (H_{\text{int}}^{4w})_{\alpha\beta;\alpha;\beta} \prod_{i=1}^4 R_{a_i} R_{a_i}
\]

under the constraint \(R_a R_a = 1\), which leads to the self-consistent eigenvalue equation:

\[(H_{\text{HF}})_{ab} R_b = \mu R_a \quad (5a)\]

with the chemical potential \(\mu\) being the lowest eigenvalue of the Hartree–Fock Hamiltonian:

\[(H_{\text{HF}})_{ab} = (H_{\text{int}})_{ab} + 2(H_{\text{int}}^{4w})_{\alpha\beta;\alpha;\beta} R_{a_i} R_{a_i}
\]

\[+ 4(H_{\text{int}}^{4w})_{\alpha\beta;\alpha\beta;\alpha\beta} \prod_{i=1}^4 R_{a_i} R_{a_i}. \quad (5b)\]

Once the amplitude \(R_a\) is determined, the ground state energy (GSE) can be computed using equation (4).

Although we have formulated the HMF method for spin-1/2 systems, it can be straightforwardly extended to higher spins as well.

Besides the GSE we will also be interested in computing the staggered magnetization \(M_z\), and the two-component VBS ‘order parameter’ [11]:

\[
\text{Re } \Psi = \frac{1}{N} \sum_x (-1)^x S_{x+e_y} S_{x+e_y};
\]

\[
\text{Im } \Psi = \frac{1}{N} \sum_x (-1)^x S_{x+e_y} S_{x_y},
\]

which allows us to characterize lattice point group symmetries of a state.

In the rest of this section we will sketch the HMF calculation of \(E_0, M_z\) and \(\Psi\) for the case of plaquettes, and only present final expressions for the \(4 \times 4\) clusters. The interested reader is referred to [7], where the technique is analyzed and developed in greater detail. For simplicity we shall put \(J \equiv 1\).

(i) The plaquette degree of freedom

We start by considering the simplest way to cover the lattice—with plaquettes, as shown in figure 2. At the same time we introduce notations and concepts, which will be used in the following subsection.

The Hamiltonian for an isolated plaquette has the form

\[
H_{\square} = \left(1 + \frac{Q}{2}\right)(S_1 + S_2)(S_2 + S_3) - Q[(S_1 S_2)(S_3 S_4) + (S_1 S_3)(S_2 S_4)].
\]

\[
= H_{\text{int}}^2 + H_{\text{int}}^{4w} + H_{\text{int}}^{4w},
\]

where appropriately symmetrized individual terms are given by

\[
= \frac{1 + Q/2}{2}[(S_{11} + S_{14})(S_{22} + S_{23}) + (S_{12} + S_{13})(S_{21} + S_{24})]
\]

The interaction of this plaquette with the rest of the system can be conveniently partitioned according to (2) as

\[
H_{\text{int}} = H_{\text{int}}^2 + H_{\text{int}}^{4w} + H_{\text{int}}^{4w},
\]

Figure 1. Schematic phase diagram of the \(J–Q\) model, obtained in the present paper. The translationally invariant AF phase with broken \(SU(2)\) symmetry and the singlet paramagnetic state are separated by a quantum phase transition at \(Q = Q_c\). Results of our calculations indicate that this transition is continuous, but we cannot rigorously discard the possibility that it becomes weakly first order in the thermodynamic limit. The non-magnetic phase breaks the lattice translational symmetry and is a plaquette paramagnet.
The coarse-grained lattice obtained by choosing the $4 \times 4$ cluster as a degree of freedom is shown in figure 3. Each spin operator carries three indices: label of a cluster, label of a plaquette within this cluster, and the position within this plaquette. Writing down the cluster self-energy and the inter-cluster interactions is a straightforward, but tedious task, which can be accomplished along the lines presented in the previous subsection. Therefore, here we give only final expressions for $M_z$ and $\Psi$. 

\begin{align*}
-\frac{Q}{2}[(S_{11}S_{12})(S_{23}S_{24}) + (S_{11}S_{23})(S_{12}S_{24}) \\
+ (S_{12}S_{13})(S_{24}S_{23}) + (S_{12}S_{21})(S_{13}S_{23}) \\
+ (S_{13}S_{14})(S_{21}S_{22}) + (S_{13}S_{21})(S_{14}S_{22}) \\
+ (S_{14}S_{12})(S_{23}S_{24}) + (S_{14}S_{23})(S_{12}S_{24})] ; \\
H_{4l}^{4h} = -\frac{Q}{8} [(S_{14}S_{23} + S_{24}S_{13})(S_{32}S_{41} + S_{31}S_{42}) \\
+ (S_{14}S_{31} + S_{44}S_{33})(S_{12}S_{31} + S_{13}S_{22})] ; \\
H_{4v}^{4h} = -\frac{Q}{8} [(S_{14}S_{32} + S_{12}S_{34})(S_{23}S_{41} + S_{21}S_{43}) \\
+ (S_{24}S_{32} + S_{22}S_{44})(S_{13}S_{31} + S_{11}S_{33})] . (7a, b, c) \\
\end{align*}

It is convenient to work in the basis which diagonalizes the $Q$-independent part of $H_{4l}$, equation (6). Such is, for instance, the basis of eigenstates of the total angular momentum of the plaquette:

$$|\alpha\rangle = |l_1 l_2 LM\rangle ,$$

$$l_1 = S_l + S_i ; \quad l_2 = S_2 + S_j ; \quad L = l_1 + l_2. \quad (8)$$

The matrix elements, which appear in equation (4)

\begin{align*}
&H_{4l}^{4h} = |a_1 a_2 \rangle \langle a_1 a_2| H_{4l}^{4h} |a_1 a_2 \rangle \\
&H_{4v}^{4h} = |a_1 a_2 a_3 a_4 \rangle \langle a_1 a_2 a_3 a_4| H_{4v}^{4h} |a_1 a_2 a_3 a_4 \rangle
\end{align*}

can now be computed using the angular momentum addition theorems.

The staggered magnetization (along the $z$-axis) within a plaquette is given by

$$M_z = \frac{1}{4} (S_i^z + S_j^z - S_{i+\hat{x}}^z - S_{j+\hat{x}}^z)_{ab} R_a R_b, \quad (9)$$

while the function $\Psi$ can be written in the plaquette representation as

\begin{align*}
\text{Re} \Psi &= \frac{1}{N} \sum_i [S_i S_{i+\hat{x}} + S_{i+\hat{y}} S_{i+\hat{y}}] \\
- \frac{1}{N} \sum_i [S_i S_{i+\hat{x}} + S_{i+\hat{y}} S_{i+\hat{y}}] ; \\
\text{Im} \Psi &= \frac{1}{N} \sum_i [S_i S_{i+\hat{x}} + S_{i+\hat{y}} S_{i+\hat{y}}] \\
- \frac{1}{N} \sum_i [S_i S_{i+\hat{x}} + S_{i+\hat{y}} S_{i+\hat{y}}] . \quad (10)
\end{align*}

In these equations the indices $i$ and $\hat{x}$ denote sites and basis vectors of the plaquette lattice.

(ii) $4 \times 4$ spin clusters

The coarse-grained lattice obtained by choosing the $4 \times 4$ cluster as a degree of freedom is shown in figure 3. Each spin operator carries three indices: label of a cluster, label of a plaquette within this cluster, and the position within this plaquette. Writing down the cluster self-energy and the inter-cluster interactions is a straightforward, but tedious task, which can be accomplished along the lines presented in the previous subsection. Therefore, here we give only final expressions for $M_z$ and $\Psi$. 

\begin{align*}
The staggered magnetization of a cluster is given by an equation analogous to (9):

$$M_z = \sum_{i=1}^{4} (S_{i1}^z + S_{i4}^z - S_{i2}^z - S_{i3}^z)_{A} R_A R_{\bar{A}} , \quad (11)$$

where the summation extends over plaquettes within a cluster, and we used capital indices $A$ to label states of the cluster. The function $\Psi$ can be written as

\begin{align*}
\text{Re} \Psi &= \frac{1}{N} \sum_i [(S_{i1} S_{i2} + S_{i3} S_{i4}) \\
+ S_{i33} S_{i3} + S_{i21} S_{i22} + S_{i23} S_{i24} \\
+ S_{i41} S_{i42} + S_{i43} S_{i44} - S_{i12} S_{i21} \\
- S_{i14} S_{i23} - S_{i32} S_{i41} - S_{i34} S_{i43}] \\
- \frac{1}{N} \sum_i [(S_{i22} S_{i21} + S_{i12} S_{i1,13}} \\
+ S_{i42} S_{i41} + S_{i44} S_{i41,33}] ; \\
\text{Im} \Psi &= \frac{1}{N} \sum_i [(S_{i11} S_{i13} + S_{i12} S_{i14} + S_{i21} S_{i23}) \\
+ S_{i22} S_{i24} + S_{i31} S_{i33} + S_{i32} S_{i34} \\
+ S_{i41} S_{i43} + S_{i42} S_{i44} - S_{i13} S_{i31} \\
- S_{i14} S_{i23} - S_{i23} S_{i41} - S_{i24} S_{i42}] \\
- \frac{1}{N} \sum_i [(S_{i33} S_{i31} + S_{i34} S_{i31,12}] \\
+ S_{i43} S_{i41,21} + S_{i44} S_{i41,22}]. \quad (12)
\end{align*}

\end{align*}
3. Results

We can now proceed with the solution of the mean-field equation (5), supplemented by equations (6), (7) for case (i) and analogous expressions in case (ii).

The physical quantities that we want to compute in the first place are the GSE and the staggered magnetization. These are given by equation (4) and equation (9) for case (i), and (11) for case (ii). In figure 4 we present GSEs for both degrees of freedom. All energies monotonically decrease with increasing $Q/J$ as a consequence of the negative sign in front of the last term in equation (1). At some critical value of $Q = Q_c$, the system undergoes a phase transition from the Néel state at small $Q$ to a spin-disordered state at $Q > Q_c$. This transition can be seen either from the second derivative of the GSE, $d^2E_0/N\,dQ^2$, shown in figure 5, or from the staggered magnetization as a function of $Q/J$, presented in figure 6. Using these plots one obtains the numerical values $Q_c/J = 1.61$ for plaquettes and $Q_c/J = 2.00$ for $4 \times 4$ clusters. Although the jump $g = J\,d^2E_0/N\,dQ^2|_{Q_c=0}$ is numerically small, it remains finite: $g \to 0.016$, if extrapolated to the thermodynamic limit, based on these two points (see the inset to figure 5). The finite-size scaling of the critical point itself, presented in the inset to figure 6, shows that $\lim_{N_q\to\infty} Q_c/J = 2.13$. In order to demonstrate that our results are reliable, we compute limiting values of the GSE and the magnetization at $Q = 0$: $\lim_{N_q\to\infty} E_0/NJ = -0.64$ and $\lim_{N_q\to\infty} M_z = 0.39$. These numbers should be compared to the accepted QMC results [12]: $E_0/NJ = -0.67$ and $M_z = 0.31$.

We note, finally, that due to few data points, the finite-size scalings presented here are qualitative, and are intended to provide only an estimate for the extrapolated quantities in the thermodynamic limit.

Let us now discuss the symmetries of the various phases. The antiferromagnetic state, which occurs for $Q < Q_c$, is known to preserve the lattice rotational symmetry $C_4$, and spontaneously breaks the spin $SU(2)$ symmetry. The nature of the paramagnetic phase, stablized for $Q \geq Q_c$, can be unveiled by computing expectation values of the function $\Psi$ given by equations (10) and (12) for cases (i) and (ii), respectively. Although $\Psi$ is an integral quantity, it is sufficient for the purpose of discriminating between plaquetted and dimerized ground states. Namely, a plaquette phase preserves the four-fold lattice rotational symmetry, implying

$$\text{Re} \, \Psi = \text{Im} \, \Psi,$$

while in a dimerized state this equality does not hold. In figure 7 we show $\text{Re} \, \Psi$ and $\text{Im} \, \Psi$. The equality (13) is satisfied throughout the phase diagram. This fact is not surprising in the antiferromagnetic phase, but in the paramagnetic region it presents strong evidence against any type of dimerized ground states. Although such states were allowed in the process of minimization, the $C_4$-symmetric states always had lower energy. In fact, the ground state in the non-magnetic region is a plaquette paramagnet, with each plaquette being in its singlet state.
ground state. However, due to the tensor nature of interactions in (1) these plaquettes are interacting.

As already mentioned in section 1, the coarse graining procedure explicitly breaks the lattice translation invariance, which should be restored in the thermodynamic limit. Extrapolation to \( Nq \to \infty \) shows that \( \text{Re} \Psi, \text{Im} \Psi \to -0.04 \), suggesting that the translational invariance is indeed being recovered.

4. Discussion

Our calculations, presented in the previous section, demonstrate that the Hamiltonian (1) exhibits a phase transition point separating the Néel ordered state from a paramagnetic phase with broken translational invariance, in agreement with conclusions of previous works [4–6]. Most importantly, besides establishing the existence of a phase transition, we were also able to unveil the nature of the paramagnetic phase and show that a correlated plaquette state is favored over a columnar dimer state which, although not conclusively, seems to be preferred according to previous calculations [4].

However, despite qualitative agreement, there is a quantitative discrepancy in the numerical value of \( Q_c \). Namely, the value obtained in the present paper is much smaller than the one presented in [4]. Although we cannot provide a rigorous explanation for this discrepancy, we would like to make some qualitative remarks in the following.

First of all, it is clear that the variational wavefunction (3), being a low-density ansatz, generally leads to an underestimation of the four-boson scattering terms in equation (2). In order to understand how significant this error is and check that the results, presented in figures 4–6, are reasonable, we used data from the exact diagonalization of \( 4 \times 4 \) spin clusters to compare magnitudes of the two terms: the ones, proportional to \( J \) and \( Q \), in equation (1). On physical grounds one would expect a phase transition to occur when these terms become comparable. Figure 8 presents the two contributions and their dependence on \( Q/J \). Of course, the crossing point at \( Q/J \sim 1 \) does not determine the critical value \( Q_c \), but it provides a clue on where the phase transition may occur. Since the system is gapped in the paramagnetic phase, one can argue that the size \( 4 \times 4 \) is large enough to describe the thermodynamic limit. Indeed, QMC data for \( Q/J = 10 \) indicates that the GSE converges very rapidly with increasing system size [13]. Also, calculations analogous to that shown in figure 8, performed [13] for systems up to \( 16 \times 16 \) sites, indicate that the magnitude of the crossing point stays of order unity.

Second, we would like to emphasize that, although there is no question about the correctness of the QMC studies of [4–6, 14], the procedure used to extract physical quantities, like \( Q_c \), from the raw statistical data, is not straightforward and requires certain assumptions [4]. Therefore, it is desirable to have another independent determination of the phase transition point, for example, from the data on staggered magnetization, computed in the entire \( Q \) range. While such calculations would definitely help to resolve this issue, surprisingly, they have never been performed. QMC computations of finite lattices does not suffer from the infamous sign problem in this case, thus it yields better energy and magnetization values than the ones obtained here. Our approach, on the other hand, focuses on establishing symmetry properties of different phases, rather than improving numerical values for observable quantities. It is this fact, which enables us to detect phase transition points within a simple framework.

Our conclusions raise another important question regarding the nature of the phase transition. We find it to be of Landau type. Although the finite-size scaling of the second-order derivative of the GSE, presented in the previous section, displays a finite jump as \( Nq \to \infty \), there is no way to rigorously prove it. Thus, the possibility of a weakly first-order transition at \( Q_c \) cannot be completely excluded. Indeed, in [14] it was argued that this phase transition, which was claimed to occur at the same point as in [4], is of the first order. As any
real-space method our approach explicitly breaks translational invariance, and although the finite-size scaling for $\Psi$ implies that this property is restored with increasing cluster size, we cannot provide a rigorous symmetry-based analysis.

In summary, we determined the phase diagram of the $J-Q$ model (1), by using the recently proposed hierarchical mean-field approach [7, 8]. It was shown that there exists a single (i.e. universal) mean-field framework (variational ansatz for the ground state), which gives the complete phase diagram of the model. In particular, we found that there exists a critical point at $Q_c \approx 2.13 J$, which separates the antiferromagnetic phase from the non-magnetic state. The latter breaks lattice translational invariance and was shown to represent a correlated plaquette paramagnetic phase. Our results suggest that the phase transition at $Q_c$ is of a Landau second-order type, even in the thermodynamic limit $N_q \to \infty$, although we cannot rigorously exclude the possibility for it to become weakly first order.

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