A frustrated spin-1/2 Heisenberg antiferromagnet on a chevron-square lattice

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The coupled cluster method (CCM) is used to study the zero-temperature properties of a frustrated spin-half ($s = \frac{1}{2}$) $J_1$-$J_2$ Heisenberg antiferromagnet (HAF) on a two-dimensional (2D) chevron-square lattice. On an underlying square lattice each site of the model has 4 nearest-neighbor exchange bonds of strength $J_1 > 0$ and 2 frustrating next-nearest-neighbor (diagonal) bonds of strength $J_2 \equiv \kappa J_1 > 0$, such that each fundamental square plaquette has only one diagonal bond. The diagonal $J_2$ bonds are arranged in a chevron pattern such that along one of the two basic square axis directions (say, along rows) the $J_2$ bonds are parallel, while along the perpendicular axis direction (say, along columns) alternate $J_2$ bonds are perpendicular to each other, and hence form one-dimensional (1D) chevron chains in this direction. The model thus interpolates smoothly between 2D HAFs on the square ($\kappa = 0$) and triangular ($\kappa = 1$) lattices, and also extrapolates to disconnected 1D HAF chains ($\kappa \to \infty$). The classical ($s \to \infty$) version of the model has collinear Néel order for $0 < \kappa < \kappa_c$ and a form of noncollinear spiral order for $\kappa_c < \kappa < \infty$, where $\kappa_c \equiv \frac{1}{2}$. For the $s = \frac{1}{2}$ model we use both these classical states, as well as other collinear states not realized as classical ground-state (GS) phases, as CCM reference states, on top of which the multispin-flip configurations resulting from quantum fluctuations are incorporated in a systematic truncation hierarchy, which we carry out to high orders and then extrapolate to the physical limit. At each order we calculate the GS energy, GS magnetic order parameter, and the susceptibilities of the states to various forms of valence-bond crystalline (VBC) order, including plaquette and two different dimer forms. We find strong evidence that the $s = \frac{1}{2}$ model has two quantum critical points, at $\kappa_{c_1} \approx 0.72(1)$ and $\kappa_{c_2} \approx 1.5(1)$, such that the system has Néel order for $0 < \kappa < \kappa_{c_1}$, a form of spiral order for $\kappa_{c_1} < \kappa < \kappa_{c_2}$ that includes the correct three-sublattice $120^\circ$ spin ordering for the triangular-lattice HAF at $\kappa = 1$, and parallel-dimer VBC order for $\kappa_{c_2} < \kappa < \infty$.

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I. INTRODUCTION

The simultaneous presence of strong frustration and large quantum fluctuations in highly frustrated and strongly correlated quantum antiferromagnets on two-dimensional (2D) lattices makes these systems of huge theoretical interest for investigating possible novel quantum phases with exotic ordering\(^\dagger\). Particular interest has focussed on the zero-temperature ($T = 0$) phase transitions that can occur between both quasiclassical states showing various forms of magnetic order and magnetically disordered, quantum paramagnetic (QP) phases, as some control parameter characterizing the degree of frustration present in the system, is varied. The latter QP phases include both various types of valence-bond crystalline (VBC) solid and quantum spin-liquid (QSL) states.

Since quantum fluctuations tend to be the largest, other things being equal, for spins with the smallest spin quantum number $s$, spin-1/2 systems always occupy a special niche. Furthermore, novel quantum phases often emerge from the corresponding classical ($s \to \infty$) models that exhibit an infinitely degenerate family of ground states in some region of the classical $T = 0$ phase diagram. What one typically finds in such a scenario is that this (accidental) classical ground-state (GS) degeneracy may be lifted entirely (or partially), by the well-known order by disorder mechanism\(^\dagger\) to favor just one (or a few) particular member(s) of the family as the actual quantum GS phase. This is often found to be the case in the quasi-classical limit ($s \gg 1$), where one works to keep quantum corrections to leading order in the parameter $1/s$. However, what is also often then found in those situations is that for the $s = \frac{1}{2}$ system none of the infinitely degenerate set of classical states that form the GS phase in the $s \to \infty$ limit survive the quantum fluctuations to form a stable magnetically ordered GS phase over part, or even all, of the $T = 0$ quantum phase diagram. Instead, in their place, emerges one or more novel QP phases with no classical counterpart.

An example of a 2D spin system that fulfills the above scenario is the anisotropic planar pyrochlore (APP) model (also known as the crossed chain model). It comprises a frustrated $J_1$-$J_2$ Heisenberg antiferromagnet (HAF) on the 2D checkerboard lattice with nearest-neighbor (NN) and next-nearest-neighbor (NNN) exchange bonds of strength $J_1 > 0$ and $J_2 \equiv \kappa J_1 > 0$, respectively. It differs from the full $J_1$-$J_2$ model on the 2D square lattice by having half of the NNN $J_2$ bonds removed, such that alternate squares have zero or two
$J_2$ bonds, resulting in a checkerboard pattern. It may thus be regarded as a 2D analog of a three-dimensional anisotropic pyrochlore model of corner-sharing tetrahedra.

Since the $T = 0$ phase diagram of the $s = \frac{1}{2}$ APP model is thus of considerable interest, much attention has been paid to it, using a large variety of theoretical techniques.\(^\text{25–23}\) Despite this huge effort, the structure of its full phase diagram has remained unsettled, at least until very recently, particularly in the regime of large values of the frustration parameter, $\kappa > 1$, which is precisely where the GS phase of the classical ($s \to \infty$) version of the model is infinitely degenerate. For example, in the limit $\kappa \to \infty$ the APP model reduces to one of essentially decoupled one-dimensional (1D) crossed isotropic HAF chains. It hence exhibits in that limit a Luttinger QSL GS phase, with a gapless excitation spectrum of deconfined spin-1/2 spinons. It might be supposed (as was argued in Ref. 4\(^\text{1}\)) that this Luttinger-liquid behavior is robust against the gradual turning on of interchain ($J_1$) couplings, so that the 2D system continues to act as a quasi-1D Luttinger liquid for large but finite values of $\kappa$. Such a 2D QSL GS phase is an example of what has been called a sliding Luttinger liquid (SLL)\(^\text{26–28}\).

Some putative numerical evidence for such a SLL phase in the APP model at large values of $\kappa$ was claimed in experiments\(^\text{9}\) that this Luttinger-liquid behavior is altered by the effects of frustration, this is not so for either of the latter models.

The corresponding GS phase diagrams of both the spin-1/2 interpolating square-triangle and the Union Jack HAF models are very different both from each other and from that of the full $J_1$–$J_2$ square-lattice HAF model. They both also differ markedly from that of the spin-1/2 APP model, despite the fact that both the interpolating square-triangle and APP models reduce to uncoupled 1D chains in the $\kappa \to \infty$ limit. Presumably the large-$\kappa$ difference in phase structure has to do with the fact that whereas the GS phase of the classical ($s \to \infty$) version of the APP model is infinitely degenerate after Néel order has been destroyed by the effects of frustration, this is not so for either of the latter models. There is also another member of the same half-depleted $J_1$–$J_2$ class of square-lattice models, which we refer to here as the chevron-square lattice model, that has largely been ignored up to now. Once again, each square plaquette contains only one $J_2$ bond, but now in a chevron pattern such that neighboring square plaquettes along one of the two square-lattice axis directions (say, rows) have the $J_2$ bond on the same diagonal, whereas along the perpendicular direction (say, columns) neighboring plaquettes have $J_2$ bonds along opposite diagonals. In our convention the chevron lattice is henceforth drawn as in Fig. 1 such that the basic chevron stripes of alternating V-shapes and inverted V-shapes are oriented vertically. Hence the square plaquettes along each row have the diagonal $J_2$ bond oriented in the same direction, and with that direction alternating between neighboring rows. On
the CCM technique that we will apply to the triangular optical lattice. It is conceivable that this model too may be experimen-
as a different form of anisotropic triangular-lattice HAF, since, as we discuss below in Sec. II, the
Jour previous interpolating square-triangle AFM model. The recent experimental quantum simulation
of frustrated magnetism in triangular optical lattices, which exploited the motional degrees of freedom of ul-
tracold atoms, provides a clear additional impetus for such studies. In the experimental simulation a specific
modulation of the optical lattice was used to tune the NN couplings on the triangular lattice in different direc-
tions independently. By introducing a fast oscillation of the lattice the experimentalists were thus able to simulate
our previous interpolating square-triangle AFM model. Since, as we discuss below in Sec. II, the J1–J2 chevron-
square lattice model may also equivalently be regarded as a different form of anisotropic triangular-lattice HAF, it
is conceivable that this model too may be experimentally realizable with ultracold atoms trapped on a 2D
diagonal lattice.

In Sec. II we describe the model itself further, including its classical (s → ∞) counterpart. The key elements of
the CCM technique that we will apply to the s = 1/2 model are reviewed in Sec. III before presenting our results in
Sec. IV. We end with a brief summary in Sec. V.

II. THE MODEL

In this paper we study the chevron-square lattice model whose Hamiltonian may be written as

$$H = J_1 \sum_{\langle i,j \rangle} s_i \cdot s_j + J_2 \sum_{\langle i,k \rangle'} s_i \cdot s_k$$  \hspace{1cm} (1)

where the operators $s_l \equiv (s^x_l, s^y_l, s^z_l)$ are the quantum spin operators on lattice site $l$, with $s^z_l = s(s+1)$. We
are interested here in the extreme quantum case $s = 1/2$. On the square lattice the sum over $\langle i,j \rangle$ runs over all
distinct NN bonds (with exchange coupling strength $J_1$), while the sum over $\langle\langle i,k \rangle\rangle'$ runs over only half of the
distinct NNN diagonal bonds (with exchange coupling strength $J_2$). In the latter sum only one NNN diagonal bond is retained in each square plaquette, as arranged in the (vertical) chevron pattern shown explicitly in Fig. 1.
The primitive unit cell is thus of size $1 \times 2$. In both sums in Eq. (1) each bond is counted once and once only.
In the present paper we consider the case where both sorts of bonds are AFM in nature, $J_1 > 0$ and $J_2 \equiv \kappa J_1 > 0$,
and hence act to frustrate one another. Henceforth, we put $J_1 \equiv 1$ to set the overall energy scale.

The model may clearly equivalently be defined on a triangular-lattice geometry in which every site has six
NN sites, four connected to it by $J_1$ bonds and two connected to it by $J_2$ bonds, as shown explicitly in Fig. 2.
Thus, the chevron-square lattice model may be regarded also as an anisotropic HAF on a triangular lattice, in
which every basic triangular plaquette contains two NN $J_1$ bonds and one NN $J_2$ bond in the pattern shown.
It thus clearly shares some properties with our previous interpolating square-triangle model in the sense that
both models interpolate smoothly between the HAF on the square lattice (when the NNN bonds retained on the
square lattice are removed entirely) and the HAF on the triangular lattice (when both NN and the retained NNN
bonds on the square lattice have equal strength). Both models can be regarded too as anisotropic triangular-
lattice HAF models. Whereas our former interpolating square-triangle model had the NN bonds along two of
the three equivalent triangular-lattice directions equal ($J_1$) and different along the third direction ($J_2$), in the
current chevron-lattice model the inequivalent ($J_2$) bonds are those along parallel zigzag (or chevron) chains in one of
the three equivalent directions for the triangular lattice, as shown in Fig. 2.

The current model clearly also reduces to decoupled 1D chains in the limit $\kappa \rightarrow \infty$. Thus, the case $\kappa \gg 1$
corresponds to weakly coupled 1D chains, and the model thus also interpolates between 1D and 2D limits.

Before considering the extreme quantum limit $s = 1/2$ it

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{(Color online) The $J_1$–$J_2$ chevron-square lattice model ($J_1 \equiv 1$). The solid (black) lines are $J_1$ bonds and the dashed (green) lines are $J_2$ bonds; (a) Néel state, (b) spiral state (or state s), (c) row-striped state (or state r), (d) columnar-striped state (or state c), and (e) doubled alternating striped state (or state d). The blue arrows represent the spins on the sites of the chevron lattice.}
\end{figure}
is worthwhile to examine first the classical limit, $s \to \infty$. Thus, it is easy to show that the classical chevron-square lattice model has only two GS phases. For $\kappa < \frac{1}{2}$ the stable GS phase has AFM Néel ordering on the square lattice, as shown in Fig. 1(a). At $\kappa = \kappa_{cl} = \frac{1}{2}$ the system undergoes a continuous second-order phase transition to a state with noncollinear spiral order that persists for all values $\kappa > \frac{1}{2}$. The spin direction in this spiral state, shown in Fig. 1(b), at lattice site $(i, j)$ points at an angle $\alpha_{ij} = \alpha_0 + [i + \Delta(j)]\alpha$, where $\Delta(j)$ is defined to be zero if $j$ is even and one if $j$ is odd. The GS energy per spin in this classical spiral state is thus given by

$$E/N = s^2(2\cos \alpha + \kappa \cos 2\alpha),$$

which is minimized when

$$\alpha = \alpha_{cl} = \begin{cases} \cos^{-1}(-\frac{1}{2\kappa}); & \kappa \leq \frac{1}{2} \\ \cos^{-1}(\frac{1}{2\kappa}); & \kappa > \frac{1}{2}. \end{cases} \tag{3}$$

The pitch angle $\phi \equiv \pi - \alpha$ thus measures the deviation from Néel order, and its classical value is

$$\phi_{cl} = \begin{cases} 0; & \kappa \leq \frac{1}{2} \\ \cos^{-1}(\frac{1}{2\kappa}); & \kappa > \frac{1}{2}. \end{cases} \tag{4}$$

It varies from zero for $\kappa < \kappa_{cl} = \frac{1}{2}$ to $\frac{\pi}{2}$ as $\kappa \to \infty$ (as shown later in Fig. 4). At the isotropic point, $\kappa = 1$, we regain the classical three-sublattice ordering of the isotropic triangular-lattice HAF with $\alpha_{cl} = \frac{\pi}{4}$.

The GS energy per spin of the classical model is thus given by

$$\frac{E}{Ns^2} = \begin{cases} -2 + \kappa; & \kappa \leq \frac{1}{2} \\ -(\frac{\kappa}{\kappa}); & \kappa > \frac{1}{2}. \end{cases} \tag{5}$$

Equation (5) clearly shows that the classical phase transition at $\kappa = \frac{1}{2}$ is a continuous (second-order) one, where both the GS energy and its first derivative with respect to the frustration parameter $\kappa$ are continuous.

In the limit of large $\kappa$ the GS configuration of the classical chevron-square lattice model represents a set of decoupled 1D HAF chains (along the chevrons of the model) with sites connected by $J_2$ bonds, and with a relative spin orientation approaching $90^\circ$ between sites on neighbouring chevrons. In this classical $\kappa \to \infty$ limit there is, however, also complete degeneracy between all states for which the relative ordering directions of spins on different HAF chevron chains is arbitrary. Clearly the spin-1/2 model also similarly reduces in the $\kappa \to \infty$ limit to a set of decoupled 1D HAF chevron chains, which exhibit the 1D Luttinger-liquid behavior described by the exact Bethe ansatz solution $\text{III}$.

In general quantum fluctuations tend to favor states with collinear spin order over ones with noncollinear order, and it thus seems possible that at sufficiently large (but finite) values of $\kappa$ the classical spiral order will be destroyed by quantum fluctuations in the spin-1/2 model of interest here, to yield a state with collinear spin ordering. In this case the probable ordering is thus clearly one with all of the 1D chevron chains exhibiting Néel ordering in the same direction, with spins alternating in orientation (say, up and down). We note that there is still an infinite family of such states in which, for example, all of the spins along a given row (in our convention) can be assigned arbitrarily to be either up or down. All of these states are degenerate in energy at the classical level, with an energy per spin given $E/N = -s^2\kappa$. They include, for example, the three states shown in Figs. 1(c), 1(d), and 1(e) in which, respectively, the spins in a given row all have the same orientation (e.g., $\uparrow\uparrow\uparrow\uparrow\cdot\cdot\cdot$), alternate in orientation (e.g., $\uparrow\downarrow\downarrow\uparrow\cdot\cdot\cdot$), and alternate in orientation in a pairwise fashion (e.g., $\uparrow\downarrow\uparrow\downarrow\cdot\cdot\cdot$). Henceforth, we refer to these three states respectively as the row-striped state (or state r), the columnar-striped state (or state c), and the doubled alternating striped state (or state d).

Clearly, from Eq. (5), the classical spiral state (or state s) always has lower energy than this classical infinitely-degenerate family of collinear AFM states. Hence, unlike in the classical APP model, where the corresponding infinitely-degenerate family of crossed 1D HAF chains does form the stable GS phase for all values $\kappa > 1$, this family never becomes the stable GS phase for the classical chevron-lattice model except in the $\kappa \to \infty$ limit. Nevertheless, as we have already suggested, it does form a likely candidate for the stable GS phase of the $s = \frac{1}{2}$ version of the model considered below, as we discuss further in Sec. IV. In this case we also expect that the (accidental) classical degeneracy might again be lifted by the order by disorder mechanism.
III. THE COUPLED CLUSTER METHOD

The CCM\textsuperscript{[27–31]} that we use here has become one of the most powerful and most universally applicable techniques of modern microscopic quantum many-body theory. In recent times it has been applied with great success to a large variety of quantum spin-lattice systems (see, e.g., Refs. \textsuperscript{[25,31,49]} and references cited therein). In particular, for 2D models of quantum magnetism, it now provides one of the most accurate methods available for their study. Some of its most notable strengths are that it provides a systematic way to study various candidate GS phases and their regions of stability, and that in each case the description is systematically improvable in terms of well-defined hierarchies of approximations to incorporate more and more of the multispin configurations present in the exact, fully correlated, GS wave function.

The CCM formalism is now briefly presented. We concentrate on its key ingredients only, as needed for present purposes, and refer the interested reader to the extensive literature on the method (see, e.g., Refs. \textsuperscript{[27,31,33–35]} for further details. Any implementation of the CCM always begins with the selection of some suitable (normalized) model (or reference) state \(| \Phi \rangle \). For spin-lattice models this is often conveniently chosen as a classical (normalized) model (or reference) state

\[
\langle \Phi | = \langle \Phi | \hat{\Phi} e^{-\hat{S}} = 1.
\]

The CCM now uses the exponential parametrizations

\[
| \Psi \rangle = e^{\hat{S}} | \Phi \rangle; \quad \langle \Psi | = \langle \Phi | \hat{\Phi} e^{-\hat{S}}, \tag{6}
\]

to incorporate explicitly the multispin-flip configurations in \(| \Psi \rangle \) and \(\langle \Psi | \) above and beyond those contained in the chosen model state \(| \Phi \rangle \). The ket- and bra-state correlation operators are then expressed as

\[
S = \sum_{I \neq 0} S_I C_I^+; \quad \hat{S} = 1 + \sum_{I \neq 0} \hat{S}_I C_I^- \tag{7}
\]

where, by definition, \(C_I^+ \equiv 1\) is the identity operator, and \(C_I^- | \Phi \rangle = 0 = \langle \Phi | C_I^- \), \(\forall I \neq 0\). The set-index \(| I \rangle\) represents a multispin-flip configuration with respect to state \(| \Phi \rangle\), as we elaborate further below. The operator \(C_I^- \equiv (C_I^+)\) thus represents a multispin-flip creation operator with respect to \(| \Phi \rangle\) considered as a generalized vacuum state.

It is now convenient to choose a set of local coordinate frames in spin space defined for each model state separately, such that on each lattice site the spin aligns along the negative \(z\) axis (downwards). Such rotations obviously leave the basic SU(2) spin commutation relations unchanged. In such a basis the \(C_I^\pm\) operators now have the universal form, \(C_I^\pm \equiv s_j^+ s_{j+1}^+ ... s_n^+ \) in terms of the usual single-spin raising operators \(s_j^+ \equiv s_j^x + is_j^y\).

For a spin \(s\) the raising operator \(s_j^+\) may be applied a maximum of \(2s\) times on a given site \(j\), and hence the set-index \(| I \rangle \equiv \{ j_1, j_2, ...; j_n; n = 1, 2, ... \} \) where any given site-index \(j_k\) may appear a maximum of \(2s\) times. In the present \(s = 1/2\) case therefore, no index may be repeated.

The ket- and bra-state coefficients \(\{ S_I, \hat{S}_I \} \), which from Eqs. \textsuperscript{(6)} and \textsuperscript{(7)} completely specify the GS wave function, may now be obtained by minimizing the GS energy functional, \(H = \hat{H}(S_I, \hat{S}_I) \equiv \langle \Psi | \hat{H} | \Psi \rangle\), where \(H\) is the Hamiltonian of the system, with respect to all of the coefficients \(\{ S_I \} \) and \(\{ \hat{S}_I \} \) separately, \(\forall I \neq 0\). Use of Eqs. \textsuperscript{(6)} and \textsuperscript{(7)} then leads simply to the respective coupled sets of equations

\[
\langle \Phi | C_I^- e^{-S} \hat{H} e^{S} | \Phi \rangle = 0 \tag{8}
\]

and

\[
\langle \Phi | \hat{S}_I \hat{H} e^{S} e^{-S} | \Phi \rangle = 0; \forall I \neq 0 \tag{9}
\]

which are fully equivalent to the GS ket and bra Schrödinger equations \(\hat{H} | \Psi \rangle = E | \Psi \rangle\) and \(\langle \Psi | \hat{H} = E \langle \Psi | \). The latter set of equations may equivalently be written as

\[
\langle \Phi | \hat{S}_I e^{-S} \hat{H} e^{S} = E C_I^- | \Phi \rangle = 0; \forall I \neq 0 \tag{10}
\]

Once these sets of CCM equations have been solved for the sets of correlation coefficients \(\{ S_I \}\) and \(\{ \hat{S}_I \}\), all GS quantities may now be calculated in terms of them. The GS energy is unique in that its evaluation requires only the ket-state coefficients, \(E = \langle \Phi | e^{-S} \hat{H} e^{S} | \Phi \rangle\), whereas any other GS quantity requires the bra-state coefficients also. For present purposes, for example, we will also calculate the magnetic order parameter, defined to be the local average on-site magnetization, \(M\), in the rotated spin coordinates, \(M \equiv -\frac{1}{N} \langle \Psi | \sum_{j=1}^{N} s_j^z | \Psi \rangle\), where \(N\) is the number of lattice sites. A key feature of the CCM exponential parametrizations of Eq. \textsuperscript{(6)} is that the method automatically satisfies the Goldstone linked cluster theorem, even when truncations are made in the multispin configurations \(\{ I \}\) retained in Eq. \textsuperscript{(7)ʹ}. Thus, the method is automatically size-extensive at every level of approximation, and we may take the infinite-lattice limit, \(N \rightarrow \infty\), from the very outset. Similarly, the method also satisfies the Hellmann-Feynman theorem at all levels of truncation.

It is important to note too that while the CCM equations are intrinsically nonlinear, due to the presence of the basic correlation operator \(S\) in the exponentiated form \(e^S\), it only ever appears in the equations to be solved in the similarity transform, \(e^{-S} \hat{H} e^S\), of the Hamiltonian. By making use of the well-known nested commutator expansion for \(e^{-S} \hat{H} e^S\), it is readily seen that the basic SU(2) spin commutation relations imply that the otherwise infinite series of nested commutators actually terminates exactly at second-order terms in \(S\) for Hamiltonians of the form of Eq. \textsuperscript{(6)} used here (and see, e.g., Refs. \textsuperscript{[31,32]} for further details). A similar exact termination generally also applies to the evaluation of the GS expectation value of any operator, such as \(M\) above, that
we calculate.

Thus, the CCM formalism is exact if all multispin-flip configurations are included in the index-set \( \{ I \} \), and the equations that we need to solve in practice are coupled sets of multinomial equations for the coefficients \( \{ S_I \} \) and linear equations for the coefficients \( \{ \tilde{S}_I \} \), in which the solutions for \( \{ S_I \} \) are needed as input. Of course, in practice we need to make finite-size truncations in the multispin-flip configurations retained in the indices \( \{ I \} \). We employ here the localized (lattice-animal-based subsystem) LSUBn scheme \(^{23,31,33}\) which has been well-studied and greatly utilized for a wide variety of spin-1/2 lattice systems \(^{23,31,33,34,49}\). At the \( n \)th level of approximation in the LSUBn scheme all possible multispin-flip configurations are retained in the index-set \( \{ I \} \) over different locales on the lattice defined by \( n \) or fewer contiguous sites. In other words, all lattice animals of size up to \( n \) sites are populated with flipped spins (with respect to the model state \( | \Phi \rangle \) as reference state) in all possible ways. Such animals (or contiguous clusters) are defined to be contiguous in this sense if every site in the cluster is adjacent (in the NN sense) to at least one other site in the cluster. For our present model we define NN pairs on the triangular-lattice geometry (as shown in Fig. \( 2a \)), rather than on the square-lattice geometry, since we wish to treat all pairs connected by both \( J_1 \) and \( J_2 \) bonds on an equal footing.

The number \( N_f \) of such fundamental configurations, which are distinct under the space- and point-group symmetries of the lattice and of the model state being used, increases very rapidly with the truncation index \( n \) of the LSUBn hierarchy. In the present study, by making use of a highly efficient parallelized CCM code \(^{52}\) and supercomputer resources we have been able to perform LSUBn calculations up to the LSUB10 level for each of the four collinear model states of Figs. \( 1a, c, d, e \), and up to the LSUB8 level for the spiral state of Fig. \( 1b \). For example, using the triangular-lattice geometry, \( N_f = 541578 \) for the LSUB10 approximation using the row-striped state of Fig. \( 2c \) as CCM model state.

Although we never need to perform any finite-size scaling of our results, since we work from the outset in the thermodynamic limit, \( N \to \infty \), we do need to extrapolate our approximate LSUBn sequence for any physical quantity to the exact \( n \to \infty \) limit. For example, for the GS energy per spin, \( E/N \), we use the well-tested extrapolation scheme \(^{23,34,41,43,45,49}\).

\[
E(n)/N = a_0 + a_1 n^{-2} + a_2 n^{-4}. \tag{8}
\]

As is to be expected other physical quantities do not converge as rapidly as the energy. For example, for the magnetic order parameter, \( M \), we generally find that a scaling law with leading power \( 1/n \), i.e.,

\[
M(n) = b_0 + b_1 n^{-1} + b_2 n^{-2}, \tag{9}
\]

works well for most systems with even moderate amounts of frustration \(^{32,37}\). However, for systems very close to a QCP or for which the magnetic order parameter of the phase under study is very small or zero, the scaling law of Eq. \( 9 \) has been found to overestimate the magnetic order and to predict a too large value for the critical value of the frustration parameter that is driving the transition. In such cases we use the well-studied extrapolation scheme \(^{23,34,41,43,45,49}\).

\[
M(n) = c_0 + c_1 n^{-1/2} + c_2 n^{-3/2}. \tag{10}
\]

Naturally, for any physical quantity \( Q \) we may always test for the correct leading power in the corresponding LSUBn extrapolation scheme by first fitting to a form

\[
Q(n) = q_0 + q_1 n^{-\nu}, \tag{11}
\]

where the exponent \( \nu \) is also a fitting parameter. For the GS energy we generally find a value \( \nu \approx 2 \) for a wide variety of both unfrustrated and (strongly) frustrated systems, as is also the case here, as we discuss in Sec. \( IV \) in more detail. On the other hand, for the magnetic order parameter, \( M \) we find \( \nu \approx 1 \) for many systems, even moderately frustrated systems such as the triangular-lattice HAF, which is the special case \( \kappa = 1 \) of the present model. However, in situations where we are either close to a transition or where \( M \) is (close to) zero for the phase under study we generally find \( \nu \approx 0.5 \). Both cases are present in the current model, as we discuss more fully in Sec. \( IV \). Once such values for the leading exponent have been found, they provide the subsequent rationale for the use of Eqs. \( 9 \)–\( 10 \), for example.

### IV. RESULTS

We report here on our CCM calculations for the present spin-1/2 J\(_1\)–J\(_2\) chevron-square lattice model of Eq. \( 1 \) based in turn on each of the Néel, spiral, row-striped, columnar-striped, and doubled alternating striped states shown, respectively in Figs. \( 1a, b, c, d, e \). The computational power at our disposal is such that we can perform LSUBn calculations for each of the collinear model states with \( n \leq 10 \), but for the more exacting spiral model state only for \( n \leq 8 \).

In the first place we discuss our results obtained using the spiral model state. Whereas the classical \( (s \to \infty) \) version of the model has a second-order phase transition from Néel order (for \( \kappa < \kappa_{c1} \)) to spiral order (for \( \kappa > \kappa_{c1} \)) at the value \( \kappa = \kappa_{c1} = 0.5 \) of the frustration parameter, our CCM results show a shift of this critical point to a value \( \kappa = \kappa_{c1} \approx 0.72 \) for the present \( s = \frac{1}{2} \) quantum case. This finding is in complete agreement with the general observation, which is found to be true for a wide variety of different spin-lattice models, that quantum fluctuations generally favor collinear over noncollinear forms of spin ordering, as discussed previously in Sec. \( III \).

Curves such as those shown in Fig. \( 3 \) show clearly that the CCM results based on the Néel model state (with \( \phi = 0 \)) give the minimum GS energy for all values of \( \kappa < \kappa_{c1} \), where our estimate for \( \kappa_{c1} \) is itself dependent on the
or more of the higher derivatives $d^2E/d\phi^2$, with probably also of one or more of the higher derivatives $d^nE/d\phi^n$ with $n > 2$ (as well as the first derivative $dE/d\phi$). Similar curves occur for other LSUB$n$ approximations. By contrast, in the classical case, at $\kappa = \kappa_{cl}$, $d^nE/d\phi^n$ vanishes only for $n \leq 2$. For values $\kappa > \kappa_{cl}$ the classical curves now have a maximum at $\phi = 0$ and a minimum at the value $\phi_{cl}$ given by Eq. (4). This behavior is broadly echoed in the LSUB4 $s = \frac{J_1}{J_2}$ curves shown in Fig. 3 for $\kappa > \kappa_{cl}$.

However, we note from Fig. 4 (and see especially the inset) that the crossover from one minimum ($\phi = 0$, Néel) solution to the other ($\phi \neq 0$, spiral) one appears to be rather abrupt, particularly for the LSUB$n$ approximation with $n = 4, 6$, with the pitch angle rising very sharply from a zero value on the Néel side to a nonzero value on the spiral side. This very sharp rise is itself just a reflection of the extreme flatness of the corresponding GS energy curves in Fig. 2 near the crossover point $\kappa = \kappa_{cl}$. This very flatness makes it difficult for us to distinguish between the two scenarios of a discontinuous jump in pitch angle versus a continuous but very steep rise in pitch angle as we traverse the transition in the frustration parameter $\kappa$ from the Néel phase into the spiral one. These two scenarios would correspond, respectively, to a weakly first-order transition versus a (continuous) second-order one. More compelling evidence for both the numerical value $\kappa_{cl}$ and the nature of the Néel-spiral phase transition comes from our results, presented below, for the local on-site magnetization (which provides the relevant order parameter for the transition).

However, before doing so, we comment further on Figs. 2 and 4. Firstly, we see from Fig. 2 that all of the CCM LSUB$n$ approximation give exactly the classical value $\phi = \frac{1}{2}\pi$ for the pitch angle at the value $\kappa = 1$ corresponding to the spin-1/2 HAF on the triangular lattice. This corresponds to the correct $120^\circ$ three-sublattice ordering demanded by symmetry in this case. The fact that every LSUB$n$ approximation preserves this symmetry is a simple consequence of us defining the fundamental LSUB$n$ configurations on the triangular-lattice geometry shown in Fig. 2 (in which sites connected by both $J_1$ and $J_2$ bonds are defined to be NN pairs). By contrast, had we defined the fundamental LSUB$n$ configurations on the underlying square-lattice geometry (in which only sites connected by $J_1$ bonds are NN pairs, and where sites connected by $J_2$ bonds are NNN pairs), the exact symmetry at $\kappa = 1$ would only be approximately preserved in any LSUB$n$ approximation with a finite value of $n$. 

FIG. 3: (Color online) Ground-state energy per spin, $E/N$, as a function of the spiral angle $\phi$, of the spin-1/2 $J_1-J_2$ chevron-square lattice model ($J_1 = 1, J_2 > 0$). We show results using the LSUB4 approximation of the CCM based on the spiral model state for some illustrative values of $J_2$ in the range $0 \leq J_2 \leq 1.5$. For $J_2 \lesssim 0.752$ the minimum is at $\phi = 0$ (Néel order), whereas for $J_2 \gtrsim 0.752$ the minimum occurs at $\phi \neq \phi_{LSUB4} \neq 0$, signalling a phase transition at $J_2 \approx 0.752$ in this approximation.

FIG. 4: The angle $\phi_{LSUBn}$ that minimizes the energy per spin, $E_{LSUBn}(\phi)/N$, of the spin-1/2 $J_1-J_2$ chevron-square lattice model (with $J_1 > 0$) versus the frustration parameter $\kappa \equiv J_2/J_1$, in the LSUB$n$ approximations with $n = \{2, 4, 6, 8\}$ based on the spiral model state. The corresponding classical result, $\phi_{cl}$, of Eq. (4) is also shown for comparison.
We also note from Fig. 3 that when $\kappa > 1$ the pitch angle $\phi$ approaches the asymptotic ($\kappa \to \infty$) limiting value $\frac{1}{2}\pi$ much faster for the quantum $s = \frac{1}{2}$ model than for its classical ($s \to \infty$) counterpart. This is a first direct piece of evidence that quantum fluctuations favor, in this limit, collinear ordering along the weakly coupled 1D chevron chains. Figure 3 also shows that the CCM LSUB8 approximation based on the spiral model state appears slightly anomalous in the region $\kappa > 1$, with the solution even becoming unstable for values $\kappa \gtrsim 2.2$. We take this as a first indication that the quantum spiral state itself loses its stability as the actual GS phase in this regime, as we discuss much more fully below.

Before doing so, however, we note first from Fig. 3 that our CCM solutions at a given LSUB$n$ level of implementation (viz., LSUB4 in Fig. 3) exist only for certain ranges of the spiral pitch angle $\phi$ for some values of $\kappa$. Thus, for example, for the case $\kappa = 0$ pertaining to the pure square-lattice HAF (with NN interactions only), the CCM LSUB4 solution based on the spiral model state of Fig. 3(b) only exists for values of the frustration parameter in the range $0 \leq \kappa \lesssim 0.29\pi$. For this value $\kappa = 0$, where the Néel state is the physical GS phase, it seems that any attempt to drive the system too far away from collinearity leads to the CCM solutions themselves becoming unstable in the sense that a real solution simply ceases to exist. Similarly, for example, when $\kappa = 1.5$, Fig. 3 shows that the CCM LSUB4 solution exists only for values $0.15\pi \lesssim \kappa \lesssim 0.5\pi$. In this case the spiral state provides the stable GS phase, and any attempt now to move too close to collinearity leads again to instability.

Similar instabilities, which manifest themselves as termination points, occur in all LSUB$n$ approximation (with $n > 2$). Indeed, such terminations of CCM solutions are both very common and well understood (see, e.g., Refs. 31,12). They are always reflections of the actual quantum phase transitions in the system under study. In the present case this is simply the transition between the Néel and spiral phases.

In view of the fact that we have indications or suggestions that the classical spiral phase might become unstable (for larger values of the frustration parameter $\kappa$) against quantum fluctuations, we have also performed CCM calculations based on the three other collinear states shown in Fig. 1(c), 1(d), and 1(e). All of these share the feature that the magnetic ordering along each chevron chain (i.e., sites joined by $J_2$ bonds) is that of a 1D Néel HAF. As explained in Sec. II there is actually an infinite family of such states, all of which are degenerate in energy at the classical level. As expected, Fig. 4 now shows that this accidental classical degeneracy is lifted by quantum fluctuations. As a parenthetical note we add that in Fig. 4 the results using state d as CCM model state are shown only in the range $\kappa \geq 3$, since real solutions for all of the corresponding LSUB$n$ approximations used (viz., $n = \{4,6,8,10\}$) exist only in this range for this state. Corresponding terminations for states c and r occur only for values $\kappa < 1.5$, as we discuss more fully below for state r. Although the differences in the energies between the three states are small, it is clear that the row-striped state of Fig. 1(c) has the lowest GS energy for all values of the frustration parameter, $\kappa$, shown. Henceforth, therefore, we only present results comparing CCM solutions based on the Néel, spiral, and row-striped states as model states.

Thus, firstly, in Fig. 5 we show our CCM results for the GS energy per spin, $e \equiv E/N$, based on each of these states as model states. We show both raw LSUB$n$ results and the corresponding extrapolated LSUB$\infty$ results using the extrapolation formula of Eq. (5). It is clear that the LSUB$n$ results based on each model state converge extremely rapidly as $n \to \infty$. Figure 5(a) shows that there is no real evidence for a discontinuity in slope at the critical values $\kappa_{c1}$ (which themselves depend only weakly upon the order $n$ of the LSUB$n$ approximation used). If present at all, either in the raw LSUB$n$ results or the LSUB$\infty$ extrapolation, any such discontinuity can only be very weak indeed.

The overall accuracy of our results may also be ascertained by examining the special case of $\kappa = 0$ (corresponding to the Néel order of the square-lattice HAF) and $\kappa = 1$ (corresponding to the spiral, $\phi = \frac{1}{4}\pi$, order of the triangular-lattice HAF). Thus, for $\kappa = 0$, the extrapolated GS energy per spin, using our LSUB$n$ results with $n = \{4,6,8,10\}$ is $e(\kappa = 0) \approx -0.6697$, which may be compared with the results $e(\kappa = 0) = -0.6693(1)$ from a linked-cluster series expansion (SE) technique 25 and $e(\kappa = 0) = -0.669437(5)$ from a large-scale quantum Monte Carlo (QMC) simulation 24, free of the usual “minus-sign-problems” for this unfrustrated limiting case where the Marshall–Peierls sign rule 25 applies. Similarly,

![Figure 5: (Color online) Various GS energy differences, $\Delta e^{ab} \equiv e^{\text{state } a} - e^{\text{state } b}$, with $e \equiv E/N$, between states a and b, versus the frustration parameter $\kappa \equiv J_2/J_1$, for the spin-1/2 chevron-square lattice model (with $J_1 > 0$), where states a and b are chosen to be the states r (row-striped), c (columnar-striped), and d (doubled alternating striped of Figs. 1(c)–(b), respectively. In each case the energy per spin, $e$, is the LSUB$n$ result obtained from Eq. (5), using CCM LSUB$n$ results with $n = \{4,6,8,10\}$.](https://example.com/figure5.png)
for \( \kappa = 1 \), the extrapolated value using our LSUBn results with \( n = \{2, 4, 6, 8\} \) is \( \epsilon(\kappa = 1) \approx -0.5511 \), which may again be compared with the results \( \epsilon(\kappa = 1) = -0.5502(4) \) from a SE calculation \(^{55} \) and \( \epsilon(\kappa = 1) = -0.5458(1) \) from a QMC simulation \(^{57} \). It is worth noting that, unlike for the square-lattice HAF, the nodal structure of the exact GS wave function for the triangular-lattice HAF is unknown. Hence, in this latter \( \kappa = 1 \) case the QMC minus-sign problem cannot be avoided. Instead, a fixed-node approximation was made in Ref. \(^{57} \), which was then relaxed using a stochastic reconfiguration technique. The resulting approximate result for the GS is now only an upper bound, and our own CCM result is almost certainly more accurate in this case.

In both Figs. 6(a) and 6(b) the extrapolated results are shown only for those values of \( \kappa \) in each case for which real solutions exist for all of the LSUBn approximations used in the extrapolation. In the case of the spiral state used as CCM model states, shown in Fig. 6(a), the extrapolation is thus shown for the range \( 0 < \kappa < 2.2 \), since the LSUB8 solution shows an instability for \( \kappa \geq 2.2 \) as discussed previously. Similarly, in Fig. 6(b), we can see that each of the LSUBn results with \( n > 2 \) based on the row-striped state as model state, terminate at some lower value, \( \kappa_t(n) \), which itself depends on the level \( n \) of approximation. Thus, we find \( \kappa_t(4) \approx 0.95 \pm 0.05, \kappa_t(6) \approx 1.15 \pm 0.05, \kappa_t(8) \approx 1.35 \pm 0.05, \text{and } \kappa_t(10) \approx 1.35 \pm 0.05. \)

As we have noted above, such termination points of our CCM LSUBn calculations are always reflections of the true quantum phase transitions in the system. \(^{31,32,33} \) They arise due to the solution to the CCM LSUBn equations becoming complex there, such that beyond any such termination point two unphysical branches of complex conjugate solutions exist. Conversely, in the region where the true physical solution is real, there actually exists another (unstable) real solution, which is both unphysical and generally very difficult to determine numerically. The physical branch is also usually easily identifiable in practice as the one which becomes exact in some limiting case. At a termination point the two real branches of solution (physical and unphysical) meet, and then diverge again as wholly unphysical complex conjugate pairs beyond the termination point for real solutions. The values of \( \kappa_t(n) \) may themselves be used to estimate the corresponding quantum critical point (QCP) \( \kappa_c = \lim_{n \to \infty} \kappa_t(n) \). We do not attempt to do so here since it is difficult to obtain values of \( \kappa_t(n) \) with high precision, since the number of iterations required to solve the CCM LSUBn equations to a given level of accuracy generally increases significantly as \( \kappa \to \kappa_t(n) \). Furthermore, we have other more accurate criteria for determining the QCPs, as we discuss below. Nevertheless, even the crude values for \( \kappa_t(n) \) above for \( n = \{4, 6, 8, 10\} \) indicate that the row-striped state itself can only be a possible candidate for the GS phase of our system for values of \( \kappa \gtrsim 1.5 \).

We show in Fig. 7 our CCM LSUB\( \infty \) extrapolated results for the GS energy per spin, \( \epsilon \equiv E/N \), based on the Néel, spiral and row-striped model states, using Eq. 9 and our corresponding LSUBn results for the two data sets \( n = \{2, 4, 6, 8\} \) and \( n = \{4, 6, 8, 10\} \). We note in passing that if our LSUBn results are instead fitted to a form \( \epsilon(n) = \epsilon_0 + \epsilon_1 n^{\nu} \), the fitted value for the exponent \( \nu \) turns out to be very close to 2 for all values of \( \kappa \) except those close to the termination points of the data set used, for each of our model states. The use of Eq. 8 is thereby again justified for this model. We see firstly from Fig. 7 that the results are almost identical using both data sets for the row-striped case illustrated. Similar insensitivity of our extrapolated results for the GS energy against the LSUBn data set used holds for the other model states too. Secondly, we note how our extrapolated results based on both the spiral and row-striped states are very close to one another, although not identical, as we discuss further below. Thirdly, we note that in the \( \kappa \to \infty \) (decoupled
spin-1/2 1D HAF chevron chains) limit, the results shown in Fig. 4 for the row-striped state yield an extrapolated value of the energy per spin which approaches the value \( E/N = -0.4431\kappa \), which agrees precisely with the exact result, \( E/N = \left( \frac{1}{4} - \ln 2 \right)\kappa \), from the Bethe ansatz solution.51,58

In Fig. 5 we show directly the GS energy difference, \( e \equiv E/N \), between the spiral and row-striped states. We display both raw results at various LSUBn levels of approximation and extrapolated results where we use Eq. (5) with various LSUBn data sets for each state separately before taking the difference. In each case the extrapolated results are shown for values between the respective termination points for each of the two states. Although the energy difference is small, it is well within our level of accuracy.

It is interesting to note that while the spiral state has lower energy than the row-striped state for all values (\( \kappa < 5 \)) of the frustration parameter shown at the lowest LSUB2 level of approximation, there is a crossover point above which the row-striped state lies lower in energy than the spiral state for each of the higher LSUB4 and LSUB6 levels of approximation. This is also true at the LSUB8 level, but the crossover point is now rather close to the point (\( \kappa \approx 2.2 \)) at which the corresponding LSUB8 solution for the spiral state becomes unstable. The crossover point for the extrapolated results clearly shows some (relatively small) sensitivity with respect to which set of LSUBn results is used to perform the extrapolation. In view of the instability of the spiral (LSUB8) results in the relevant range, our best estimate undoubt-
GS phase. In Fig. 10 we present various extrapolations of the exponent \( \nu \) more detail later.

Our best estimate yields to the noncollinear spiral order. Our best estimate \( \kappa_1 \approx 0.307(1) \) from a fixed-node QMC simulation, and \( \kappa_2 \approx 1.7(5) \) from a linked-cluster SE calculation. Corresponding CCM LSUBn results for the magnetic order parameter, \( M \), for the row-striped state are shown in Fig. 9(b) with \( n = \{2, 4, 6, 8\} \), and \( M(\kappa = 1) \approx 0.2219 \) using Eq. (6) with LSUBn results for \( n = \{2, 4, 6, 8\} \) and \( M(\kappa = 1) = 0.1893 \) from \( n = \{4, 6, 8\} \). Again, we are in excellent agreement with other values \( M(\kappa = 1) = 0.205(10) \) from a fixed-node QMC simulation, and \( M(\kappa = 1) = 0.19(2) \) from a linked-cluster SE calculation.

Figure 9(a) is completely consistent with the result that the magnetic order parameter \( M \to 0 \) at the critical point \( \kappa = \kappa_{c1} \), where the collinear Néel order for \( \kappa < \kappa_{c1} \) yields to the noncollinear spiral order. Our best estimate for \( \kappa_{c1} \), based on the results for \( M \), is \( \kappa_{c1} \approx 0.72(1) \), where the error is estimated from a sensitivity analysis with respect to which LSUBn results are used in the extrapolation. There is also definite evidence from Fig. 9(a) that in the spiral regime \( (\kappa > \kappa_{c1}) \), the spiral order parameter \( M \) again becomes zero (or very close to zero) at a second critical value, \( \kappa_{c2} \approx 1.4 \), which we discuss in more detail later.

Corresponding CCM LSUBn results for the magnetic order parameter, \( M \), for the row-striped state are shown in Fig. 9(b) with \( n = \{2, 4, 6, 8, 10\} \). In this highly frustrated regime, if we fit our results to a form \( M(n) = m_0 + m_1 n^{-\nu} \), as in Eq. (11), we find that the fitted value of the exponent \( \nu \) turns out to be very close to 0.5 over almost the entire range of values of \( \kappa \), independent of which LSUBn data set is used for the fit. Hence, in this case it is appropriate to use the extrapolation scheme of Eq. (10), and in Fig. 9(b) we use the data set \( n = \{2, 4, 6, 8\} \) for illustration purposes and compatibility with Fig. 9(a).

What is most striking about the extrapolated result is that \( M \) appears to be essentially zero (or less than zero, and hence unphysical) over the entire regime in which the row-striped state exists as a candidate for the stable GS phase. In Fig. 10 we present various extrapolations of our CCM LSUBn data, from which we observe their relative insensitivity to the choice of data set.

All of our results so far are consistent with the interpretation that the spiral state exists as the stable GS phase only in a regime \( \kappa_{c1} < \kappa < \kappa_{c2} \), where \( \kappa_{c2} \approx 1.7(5) \). However, the estimates so far for \( \kappa_{c2} \) are based on the

FIG. 9: (Color online) CCM LSUBn results for the GS magnetic order parameter, \( M \), versus the frustration parameter \( \kappa \). In (a) and (b) respectively as model states with \( n = \{2, 4, 6, 8\} \), and \( M(\kappa = 1) \approx 0.2219 \) using Eq. (6) with LSUBn results for \( n = \{2, 4, 6, 8\} \) and \( M(\kappa = 1) = 0.1893 \) from \( n = \{4, 6, 8\} \). Again, we are in excellent agreement with other values \( M(\kappa = 1) = 0.205(10) \) from a fixed-node QMC simulation, and \( M(\kappa = 1) = 0.19(2) \) from a linked-cluster SE calculation.

FIG. 10: (Color online) Various extrapolated CCM results for the GS magnetic order parameter, \( M \), versus the frustration parameter \( \kappa \), based on the Néel (left curves) and spiral (central curves) model states and the row-striped (right curves) model state, for the spin-1/2 \( J_1-J_2 \) chevron-square lattice model (with \( J_1 > 0 \)). The curves LSUB\( \infty \)(k) with \( k = 1, 2, \ldots, 5 \) use the extrapolation scheme of Eq. (10) and LSUBn data sets respectively: \( k = 1, n = \{2, 4, 6, 8\} \); \( k = 2, n = \{4, 6, 8, 10\} \); \( k = 3, n = \{6, 8, 10\} \); \( k = 4, n = \{4, 6, 8\} \), and \( k = 5, n = \{2, 4, 6\} \). The curves LSUB\( \infty \)(k) with \( k = 6, 7, 8 \) use the extrapolation scheme of Eq. (10) and LSUBn data sets, respectively: \( k = 6, n = \{4, 6, 8, 10\} \), for the Néel model state; and \( k = 7, n = \{2, 4, 6, 8\} \), and \( k = 8, n = \{2, 4, 6\} \) for the spiral model state.
energy crossing point between the spiral and row-striped states, the termination points for the LSUBn row-striped results, and the magnetic order parameter results for the row-striped state. Nevertheless, the latter results also indicate that, since $M \approx 0$ everywhere that the row-striped solution exists as a real solution, the row-striped state it-

dicate that, since $\chi_p$ is a real solution, the row-striped state. Nevertheless, the latter results also in-

taken implicitly in the caption. For each form of VBC order when the corresponding extrapolated

termination points for the LSUBn row-striped states, the termination points for the LSUBn row-striped

energy crossed. The most direct and most unbiased way to extrapolate the results is to use the LSUBn scheme for the perturbed

ers for the perturbed energy

where the exponent $\nu$ is also a fitting parameter. Generally, as we discuss below, the fitted value of $\nu$ is close to the value 2, as expected from our standard energy extrapolation scheme of Eq. (8), over most of the range of values of $\kappa$ pertaining to the model state used, except very near any corresponding termination point for the given state, where $\nu$ then falls sharply. Alternatively, we have also found previousf that our LSUBn values $\chi(n)$ may of-
ten themselves be directly extrapolated very accurately using the same scheme as in Eq. (8) for the GS energy, viz., $\chi(n) = d_0 + d_1 n^{-2} + d_2 n^{-4}$, at least in regions not close to a divergence of $\chi$. We saw similarly that a corresponding extrapolation of the inverse susceptibility,

also often gives consistent results that agree closely with those from the corresponding above extrapolation of $\chi$, except again in regions close to where $\chi^{-1} \to 0$. However, since we shall specifically be interested in some cases here in precisely such regions over an extended range of values of the frustration parameter $\kappa$, we also use the fitting function

with an exponent $\nu$ that is itself a fitting parameter, on appropriate occasions.

We first present results in Fig. 12 for the inverse pla-

tude susceptibility, $\chi_p^{-1}$, pertaining to the PVBC order-

ing illustrated in Fig. 11(a). We show explicitly the LSUBn results with $n = \{4, 6, 8, 10\}$ based on both the Néel and row-striped states as CCM model states. The corresponding susceptibility,

is then calculated. Clearly the phase corresponding to the model state used becomes unstable against the specified form of VBC order when the corresponding extrapolated inverse susceptibility, $\chi_d^{-1}$, goes to zero.

The most direct and most unbiased way to extrapolate the results is to use the LSUBn scheme for the perturbed energy

where

energy crossing point between the spiral and row-striped states, the termination points for the LSUBn row-striped

FIG. 11: (Color online) The perturbations (fields), $F = \delta \hat{O}$, for the various forms of valence-bond crystal (VBC) susceptibility considered, namely: (a) plaquette (PVBC), $\chi_p$; (b) parallel-dimer (||DVBC), $\chi_{d||}$; and (c) perpendicular-dimer (⊥DVBC), $\chi_{d\perp}$. In case (a) the thick (red) and thin (black) solid lines correspond respectively to strengthened and weakened $J_1$ exchange couplings, where $\hat{O}_p = \sum_{\langle i,j \rangle} a_{ij} \mathbf{s}_i \cdot \mathbf{s}_j$, and the sum runs over all NN bonds of the square lattice, with $a_{ij} = +1$ and $-1$ for thick (red) and thin (black) solid lines respectively, as shown. In cases (b) and (c) the thick (red) and thin (green) dashed lines correspond respectively to strengthened and weakened $J_2$ exchange couplings, and $\hat{O}_d = \sum_{\langle i,j,k \rangle} a_{ik} \mathbf{s}_i \cdot \mathbf{s}_k$, and the sum runs over the diagonal bonds of the chevron-square lattice, with $a_{ik} = +1$ and $-1$ for thick (red) and thin (green) dashed lines respectively, as shown in each case.
a single value which is very close to our previous best estimate for $\kappa_{c_1}$. While the row-striped state, on the other hand, is clearly much less susceptible to PVBC ordering, the corresponding LSUB\infty estimate for $\chi_p^{-1}$ again becomes very close to zero at a value $\kappa_{c_2} \approx 1.4$, which is again compatible with our previous estimates for $\kappa_{c_2}$. Presumably, if we were also to perform calculations for $\chi_p^{-1}$ for the spiral phase, it too would be nonzero (and positive) everywhere except at the values $\kappa_{c_1}$ and $\kappa_{c_2}$. However, such calculations for the spiral phase are computationally very expensive, and as noted previously can be performed only for relatively lower LSUBn orders of approximation. Even for the collinear row-striped state the number $N_f$ of fundamental spin configurations included in our CCM calculation at the LSUB10 level is 2160167.

While the results shown in Fig. 12 corroborate our previous estimates for the QCPs at $\kappa_{c_1}$ (at which Néel order yields to spiral order) and $\kappa_{c_2}$ (at which spiral order ceases), they essentially add nothing new. Thus, it is clear that the PVBC state cannot be the stable GS phase for any range of values $\kappa > \kappa_{c_2}$, since $\chi_p^{-1}$ does not vanish over any finite range in this regime. The fact that $\chi_p^{-1}$ vanishes at individual points (viz., $\kappa_{c_1}$ and $\kappa_{c_2}$ here) is simply a reflection of these being actual QCPs. Thus, at a QCP we expect that the system will indeed become infinitely susceptible to all forms of ordering (or, at least, those which are compatible with the symmetries of the actual and trial states).

We now turn our attention to the two DVBC states of Figs. 13(b) and (c) as possible GS phases for $\kappa > \kappa_{c_2}$.

Firstly, since our results for the inverse susceptibility, $1/\chi_{d_{\parallel}}$, for $\parallel$DVBC ordering are qualitatively very similar to those shown in Fig. 12 for PVBC ordering, we do not show them. Again, we find that using the Néel model state $\chi_{d_{\perp}}^{-1}$ vanishes at a single point, which is itself completely compatible with our previous estimates for $\kappa_{c_1}$, and using the row-striped model state $\chi_{d_{\parallel}}^{-1}$ vanishes at a single point compatible with our previous estimates for $\kappa_{c_2}$. Thus, again, the $\parallel$DVBC state is excluded as a stable GS phase in any range $\kappa > \kappa_{c_2}$.

By contrast, our results for the inverse susceptibility, $1/\chi_{d_{\perp}}$, for $\parallel$DVBC ordering, shown in Fig. 13(a) are qualitatively quite different in the regime $\kappa > \kappa_{c_2}$. In the inset to Fig. 13(a) we show the raw LSUBn results for $n = \{2, 4, 6, 8, 10\}$ using both the Néel and row-striped states as CCM model states, while in the main figure we show several LSUB\infty extrapolations based on both Eqs. (13) and (14), and using various LSUBn data sets for the fitting procedure. In Fig. 13(b) we also show the fitted values for the exponent $\nu$ in the corresponding extrapolation schemes.

In the first place we find that on the Néel side all of the extrapolations yield almost identical results, with $\chi_{d_{\parallel}}^{-1}$ going to zero at values $\kappa \rightarrow \kappa_{c_1}$, which are in agreement with all previous estimates. Secondly, however, we now see from Fig. 13(a) that using the row-striped state as CCM model state, all of our extrapolations yield the result that $1/\chi_{d_{\perp}}$ is zero (or very close to zero) for all values $\kappa > \kappa_{c_2} \approx 1.5$. We thus now have very strong evidence indeed that for $\kappa > \kappa_{c_2}$, the GS spiral ordering present for $\kappa_{c_2} < \kappa < \kappa_{c_{1}}$ gives way to $\parallel$DVBC ordering as the stable GS configuration.

Figure 13(b) shows that when $\chi_{d_{\perp}}$ is calculated using Eq. (13) as the extrapolation scheme the fitted exponent $\nu$ is very close to the expected value 2 for a very wide range of values of $\kappa$, as in our standard GS energy scheme of Eq. (4), except for values of $\kappa$ close to $\kappa_{c_2}$, where it drops. In practice we calculate the second derivative of $\epsilon(\delta)$ in Eq. (12) numerically using values $\delta = 0, \pm d$, typically with $d = 0.001$, and the value of $\nu$ plotted in Fig. 13(b) is essentially identical for these three values of $\delta$ used. By contrast, when $1/\chi_{d_{\parallel}}$ is calculated using Eq. (14), the fitted exponent $\nu$ is seen to lie very close to the value 0.75 for a very wide range of values of $\kappa$, except for values close to $\kappa_{c_2}$ where it rises rapidly. Interestingly, this same value of 0.75 was also observed previously for the spin-1/2 $J_1$–$J_2$ Heisenberg antiferromagnet on the 2D checkerboard lattice (viz., the APP model) when using a similar state as CCM model state in a region where the order parameter for that state was zero everywhere, and where a corresponding DVBC state also provided the stable GS configuration.

V. SUMMARY

We have investigated the $T = 0$ GS phase diagram of an $s = \frac{1}{2}$ $J_1$–$J_2$ HAF on a chevron-square lattice,
the classical ($s \rightarrow \infty$) version of which has only two GS phases, viz., a Néel phase for $J_2/J_1 \equiv \kappa < \kappa_{c1} = \frac{1}{3}$, and a spirally-ordered phase for $\kappa > \kappa_{c1}$. The model interpolates continuously between 2D HAF's on the square ($\kappa = 0$) and triangular ($\kappa = 1$) lattices, and also extrapolates in the $\kappa \rightarrow \infty$ limit to uncoupled 1D HAF chains.

We found, as expected, that the quantum fluctuations present in the $s = \frac{1}{2}$ model tend to stabilize the collinear Néel-ordered state, which is well established as the GS phase for the square-lattice HAF, to larger values of the frustration parameter $\kappa$ than in the classical case. We found a first QCP at $\kappa = \kappa_{c1} \approx 0.72(1)$ at which the Néel order gives way to spiral order. The pitch angle $\phi$ of the spiral phase (that measures the deviation from Néel order corresponding to $\phi = 0$) was found to increase markedly more rapidly as $\kappa$ is increased beyond $\kappa_{c1}$, than in the classical version as $\kappa$ is increased beyond $\kappa_{c1}$.

Although the transition at $\kappa = \kappa_{c1}$ appears still to be a continuous (second-order) one, just as for the classical model at $\kappa = \kappa_{c1}$, we cannot from our results rule out a weakly first-order transition at which the pitch angle $\phi$ undergoes a discontinuous jump from a value of zero (Néel order) for $\kappa < \kappa_{c1}$ to a nonzero value (spiral order) infinitesimally above $\kappa_{c1}$. The angle $\phi$ becomes exactly $60^\circ$ at all levels of CCM LSUBn approximation at $\kappa = 1$, corresponding to the exact three-sublattice relative $120^\circ$ spin ordering that is well established as the GS phase for the spin-1/2 triangular-lattice HAF.

At still higher values of the frustration parameter, $\kappa > 1$, we found that the pitch angle $\phi$ of the $s = \frac{1}{2}$ model approaches the limiting value of $90^\circ$ (which corresponds to Néel AFM ordering along the 1D chevron chains of $J_2$ bonds) much faster than in the classical ($s \rightarrow \infty$) model. In turn this led us to consider other possible collinear phases in this regime, which never form the stable GS phase in the classical case, except in the precise $\kappa \rightarrow \infty$ limit, but which comprise an infinitely degenerate family of states corresponding to Néel ordering along the 1D chevron chains.

We found that the order by disorder mechanism distinguishes one such state from the rest, so that quantum fluctuations in the $s = \frac{1}{2}$ case favor it to lie lowest in energy. Furthermore, above some value of the frustration parameter, $\kappa > \kappa_{c2}$, this state was seen to become lower in energy too than the spiral state for the $s = \frac{1}{2}$ model. However, when we then calculated its magnetic order parameter, we found it to be zero, within very small numerical error bars, everywhere.

We were thus led to the conclusion that the spiral order in the $s = \frac{1}{2}$ model does not persist for all values $\kappa > \kappa_{c1}$, but only for a finite range $\kappa_{c1} < \kappa < \kappa_{c2}$, unlike in the classical model where it persists for all values $\kappa > \kappa_{c1}$. Instead, we found again that the magnetic order parameter of the spiral phase seems to vanish for values of $\kappa$ above a certain critical value, which is itself consistent within small errors bars with the energy crossing point of the spiral state with the nonclassical collinear state mentioned.

Since this latter collinear state itself has a vanishing magnetic order parameter in the same region, as already noted, it must itself yield to a state with different order-
(2004). It would certainly be interesting, therefore, both to confirm our results using other theoretical methods and to examine the corresponding \( s = 1 \) version of the model.

\[ \kappa > \kappa_{c_2} \]

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