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High-order study of the quantum critical behavior of a frustrated spin-$\frac{1}{2}$ antiferromagnet on a stacked honeycomb bilayer

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We study a frustrated spin-$\frac{1}{2}$ $J_1$-$J_2$-$J_3$-$J_4$ Heisenberg antiferromagnet on an AA-stacked bilayer honeycomb lattice. In each layer we consider nearest-neighbor (NN), next-nearest-neighbor, and next-next-nearest-neighbor antiferromagnetic (AFM) exchange couplings $J_1$, $J_2$, and $J_3$, respectively. The two layers are coupled with an AFM NN exchange coupling $J^z_0 = \delta J_1$. The model is studied for arbitrary values of $\delta$ along the line $J_1 = J_2 = \alpha J_3$, that includes the most highly frustrated point at $\alpha = \frac{1}{2}$, where the classical ground state is macroscopically degenerate. The coupled cluster method is used at high orders of approximation to calculate the magnetic order parameter and the triplet spin gap. We are thereby able to give an accurate description of the quantum phase diagram of the model in the $\alpha\delta$ plane in the window $0 \leq \alpha \leq 1$, $0 \leq \delta \leq 1$. This includes two AFM phases with Néel and striped order, and an intermediate gapped paramagnetic phase that exhibits various forms of valence-bond crystalline order. We obtain accurate estimations of the two phase boundaries, $\delta = \delta_\alpha(0)$, or equivalently, $\alpha = \alpha_\delta(0)$, with $i = 1$ (Néel) and 2 (striped). The two boundaries exhibit an “avoided crossing” behavior with both curves being re-entrant. Thus, in this $\alpha\delta$ window, Néel order exists only for values of $\delta$ in the range $\delta_\alpha(0) < \delta < \delta_\alpha^1(0)$, with $\delta_\alpha^1(0) = 0$ for $\alpha < \alpha_\delta^1(0) \approx 0.46(2)$ and $\delta_\alpha^1(0) > 0$ for $\alpha_\delta^1(0) < \alpha < \alpha_\delta^2(0) \approx 0.49(1)$, and striped order similarly exists only for values of $\delta$ in the range $\delta_\alpha^2(0) < \delta < \delta_\alpha^2(0)$, with $\delta_\alpha^2(0) = 0$ for $\alpha > \alpha_\delta^2(0) \approx 0.600(5)$ and $\delta_\alpha^2(0) > 0$ for $\alpha_\delta^2(0) > \alpha > \alpha_\delta^2 \approx 0.56(1)$.

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I. INTRODUCTION

Quantum spin-lattice models, in which various types of interactions between pairs of spins compete to form different types of order in the system, provide a rich arena in which to study a wide variety of quantum phase transitions (QPTs) [1,2]. Such competition can occur either with or without magnetic frustration between the interaction bonds. In the latter case, the bonds are typically spatially anisotropic. Simple examples of such systems comprise models containing nearest-neighbor (NN) exchange interactions between spins $s_i$ on lattice sites $i$ of the Heisenberg type, $J_{ij} s_i \cdot s_j$, all with bond strength $J_{ij} > 0$, and thus all acting to promote antiferromagnetic (AFM) long-range order (LRO), but where the strengths $J_{ij}$ are not all equal. The so-called coupled-dimer magnets comprise a particularly simple, yet important and nontrivial, class of this type.

Dimerized quantum Heisenberg antiferromagnets (HAFMs) are obtained by placing quantum spins, with spin quantum number $s$, on a regular $d$-dimensional lattice with an even number of spins per unit cell. Each unit cell is divided into nonoverlapping pairs of spins (dimers). In the limit where the intradimer exchange constants $J_{ij}$ are much stronger than all of the corresponding interdimer constants, the zero-temperature ($T = 0$) ground-state (GS) phase of the system is a simple paramagnetic product of nonmagnetic spin singlets, which preserves the full spin-rotational invariance. This state has a nonzero energy gap to the lowest-lying spin triplet excitation, formed by breaking one of the spin-singlet dimers. As the interdimer exchange interactions are turned on these triplet excitations acquire mobility on the lattice, resulting in the appearance of a spin-1 bosonic quasiparticle, viz., the triplon [3]. In principle, of course, such bosons may undergo Bose-Einstein condensation (BEC) under suitable conditions and become superfluid. Indeed, such a superfluid state has been observed experimentally in the magnetic insulator TICuCl$_3$ [4–7], which is a physical realization of such a coupled-dimer magnet, in which pairs of Cu$^{2+}$ ions are antiferromagnetically coupled to form a crystalline network of dimers in a specific pattern. The BEC is induced by placing the compound in an external magnetic field, which thereby Zeeman splits the otherwise degenerate three magnetic triplet substates. At some critical field strength, the lowest lying triplet state intersects the GS dimer singlet and BEC into this triplet substate occurs, with the consequent appearance of the magnetic LRO corresponding to the off-diagonal LRO that characterizes BEC in the boson-mapped equivalent system. The whole area of BEC in magnetic insulators [8] has become one of considerable activity in recent years, at both the theoretical and experimental levels. The applied magnetic field here thus acts as a chemical potential that promotes dimer spin singlets (leaving a hole) to spin triplets (creating a triplon).

In principle, another way to induce magnetic LRO in a coupled-dimer magnet, without the application of an external magnetic field, is to increase the relative strength of the interdimer couplings $J_{ij}$ with respect to their intradimer counterparts. For example, for all two-dimensional (2D) bipartite lattices and with all couplings $J_{ij}$ between NN pairs only, when all $J_{ij}$ are equal the system will have Néel AFM magnetic LRO. Thus, if we consider the class of so-called $J$-$J'$ models on bipartite lattices, in which the intradimer NN bonds...
all have the same strength $J^\prime > 0$ and the interdimer NN bonds all have the same strength $J > 0$, there will clearly be same critical value of the relative strength parameter, $(J^\prime/J)_c > 1$, that marks a QPT between a Néel-ordered AFM GS phase and a dimerized paramagnetic GS phase. Experimentally, both in principle and sometimes in practice, such QPTs can be driven by the application of pressure to the system.

On the 2D square lattice with NN interactions only, $J$-$J^\prime$ models with specific arrangements of the $J^\prime$ bonds that have been studied include the columnar-dimer, staggered-dimer, and herringbone-dimer models (and see, e.g., Ref. [9]). The first two, each with two sites per unit cell, both have the dimer $J^\prime$ bonds parallel (say, along the row direction of the square lattice). Whereas in the columnar arrangement each basic square plaquette contains two solv or no dimer $J^\prime$ bonds, in the staggered arrangement each basic square plaquette contains a single $J^\prime$ dimer bond. Finally, the herringbone-dimer model contains four sites per unit cell with two isolated (nontouching) dimer $J^\prime$ bonds perpendicular to one another. Each basic square plaquette also contains a single $J^\prime$ dimer bond. Interestingly, in the limit $J^\prime/J \to 0$, both the staggered-dimer and herringbone-dimer models become equivalent to the HAFM on a hexagonal lattice. Thus, both of these models interpolate between HAFMs on the hexagonal and square 2D lattices for values of $J^\prime/J$ in the range $0 < J^\prime/J \leq 1$.

So far we have considered coupled-dimer magnets that include competition between bonds without frustration. The inclusion of extra bonds can now lead us into the realm of magnetic frustration, which adds further to the complexity and inherent interest of these models. For example, increasing frustration can have the effect of enhancing the repulsive interactions between triplons. In turn this can then eventually lead to the stabilization of incompressible phases that break the translational symmetry of the lattice. If such GS phases of the system are placed in an external magnetic field, the itinerant triplons become localized in a crystalline phase.

Such phases have been observed experimentally in the spin-gap material SrCu$_2$(BO$_3$)$_2$ [10–12], which is well modeled [13] by the 2D Shastry-Sutherland model [14]. This is a spin-$1/2$ coupled-dimer model on a square lattice, with four sites per unit cell, in which all NN pairs have an AFM Heisenberg bond of equal strength $J$, and the equivalent AFM dimer bonds, all of equal strength $J^\prime$, join nonoverlapping diagonal next-nearest-neighbor (NNN) pairs in an orthogonal pattern. The unit cell thus contains two orthogonal dimers arranged across NNN diagonal pairs. Clearly, in the limit $J \to 0$ the model reduces to a Hamiltonian of decoupled dimers. This dimerized state then remains the exact GS phase [14] for all values of $(J^\prime/J)$ above a certain critical value $(J^\prime/J)_c$. In the opposite limit, when $J^\prime \to 0$, the model reduces to the pure spin-$1/2$ HAFM (i.e., with NN interactions only) on the square lattice, which has Néel AFM magnetic LRO. In between, when $J^\prime = J$, the model reduces to the pure spin-$1/2$ HAFM on another of the 11 2D Archimedean lattices, the GS phase of which is also known to have Néel order [15], which implies $(J^\prime/J)_c > 1$. Various theoretical studies (see, e.g., Refs. [16,17]) yield $(J^\prime/J)_c \approx 1.48$. In the Shastry-Sutherland material SrCu$_2$(BO$_3$)$_2$, for which $(J^\prime/J) \approx 1.6$, the triplon crystalline phases show up as a series of magnetization plateaux at unconventional filling fractions [11,12] that are stabilized by complex many-body interactions among the triplons [9,12,18,19]. Indeed, the magnetization plateaux in SrCu$_2$(BO$_3$)$_2$ at low magnetization are now quite well understood in terms of triplon bound states [20,21].

Since both BEC and crystalline phases of triplons can occur in frustrated coupled-dimer magnets subjected to an external magnetic field, it is natural to wonder whether such systems might also exhibit the magnetic equivalent of supersolidity, in which a stable GS phase exhibits simultaneously both the diagonal LRO typical of a solid and the off-diagonal LRO typical of a superfluid. Such field-induced spin supersolidity has particularly been investigated for various frustrated spin-$1/2$ models on a square-lattice bilayer [8,22–27].

Such bilayer models provide other examples of coupled-dimer magnets. The simplest such bilayer models comprise two layers stacked directly on top of one another and with only NN bonds, where the intralayer bonds all have equal strength $J_1$ and the interlayer (dimer) bonds all have equal strength $J^\prime_1$. Such models on the square lattice, where the bonds compete without frustration, have been studied fairly extensively [28–32]. As the ratio $J^\prime_1/J_1$ is increased beyond a critical value $(J^\prime_1/J_1)_c$, a QPT occurs from a Néel-ordered GS phase to a paramagnetic GS phase that is approximately the product of interlayer dimer valence bonds between NN pairs coupled by $J^\prime_1$ bonds. As we have already noted above, the $T = 0$ GS phase diagram becomes appreciably richer in the additional presence of frustrating bonds of either the intralayer or interlayer type. Such frustrated square-lattice bilayer models have been much studied in recent years, using a variety of theoretical techniques, both in the absence [33,34] and presence [8,25–27,35,36] of an external magnetic field.

In the last several years attention has also been paid to analogous honeycomb-lattice bilayer models, both in the staggered Bernal AB stacking (see, e.g., Ref. [37]) relevant to bilayer graphene and in the AA stacking (see, e.g., Refs. [32,38–44]) where the two layers are stacked directly on top of one another. Since the AA stacking yields the simpler form of coupled-dimer magnets, we restrict attention here to this form of honeycomb bilayer. After the unfrustrated $J_1$-$J^\prime_1$ honeycomb bilayer was studied [32], various authors have studied the effects of both intralayer frustration [38–41] and interlayer frustration [42–44] on the system, by including NNN interactions between spins within the layers or between the layers, respectively. In the latter case the model has been studied both in the absence [42] and presence [43,44] of an external magnetic field.

There has also been experimental interest in frustrated stacked honeycomb-lattice bilayer HAFMs. For example, the Mn$^{++}$ sites in the bismuth manganese oxynitrate material Bi$_3$Mn$_4$O$_{12}$N$_3$ [45,46] form a frustrated spin-$1/2$ AA-stacked honeycomb lattice. By replacing the Mn$^{++}$ ions in this material with V$^{4+}$ ions it might also be possible to realize experimentally a spin-$1/2$ HAFM on the AA-stacked honeycomb bilayer. Ultracold atoms trapped in optical lattices formed by a periodic potential, which is created by standing waves formed from a suitable array of lasers, are now also regularly being used to simulate quantum magnets in a controllable manner [47]. For example, in the present context,
by interfering three coplanar laser beams propagating at relative angles of ±120° one may form a honeycomb lattice [48]. Even more excitingly, concrete proposals have also been given to form optical lattices representing honeycomb-lattice bilayers in both AA and AB stacking [49,50], using five lasers.

The present study has as one of its goals to investigate the effects of intralayer frustration on a particularly interesting AA-stacked bilayer honeycomb-lattice version of a spin-$\frac{1}{2}$ coupled-dimer HAFM that, to our knowledge, has not been studied before. In each layer the spins interact via NN, NNN, and NNNN intralayer bonds on each honeycomb-lattice bilayer; (c) the triangular Bravais lattice vectors $a$ and $b$, and the monolayer Néel state; and (d) one of the three equivalent monolayer striped states. Sites $(1_A, 2_B)$ and $(2_A, 1_B)$ on the two monolayer triangular sublattices are shown by filled and empty circles respectively.

The plan of the rest of the paper is as follows. In Sec. II the model

The $J_1$-$J_2$-$J_3$-$J_4^+$ model on the bilayer honeycomb lattice is described by the Hamiltonian

$$H = J_1 \sum_{\langle i,j \rangle, \alpha} s_{i,\alpha} \cdot s_{j,\alpha} + J_3 \sum_{\langle \langle i,j \rangle \rangle, \alpha} s_{i,\alpha} \cdot s_{j,\alpha}$$

$$+ J_3 \sum_{\langle \langle \langle i,j \rangle \rangle \rangle, \alpha} s_{i,\alpha} \cdot s_{j,\alpha} + J_4^+ \sum_{i} s_{i,A} \cdot s_{i,B},$$

where the index $\alpha = A, B$ labels the two layers. Each site $i$ on each of the two honeycomb layers carries a spin-$s$ particle denoted by the usual SU(2) spin operators $s_{i,\alpha} \equiv (s_{i,\alpha}^x, s_{i,\alpha}^y, s_{i,\alpha}^z)$, with $s_{i,\alpha}^2 = s(s + 1)$, and where for present purposes we restrict attention to the case $s = \frac{1}{2}$. In Eq. (1) the sums over $\langle i,j \rangle$, $\langle \langle i,j \rangle \rangle$, and $\langle \langle \langle i,j \rangle \rangle \rangle$ run respectively over all NN, NNN, and NNNN intralayer bonds on each honeycomb-lattice monolayer, counting each Heisenberg bond once and only once in each sum. The last sum in Eq. (1) describes the interlayer Heisenberg bonds between NN pairs of spins across the two vertically stacked layers. The pattern of bonds is shown in Figs. 1(a) and 1(b). In the present paper we will be interested in the case when all four bonds are AFM in nature (i.e., $J_i > 0$, $i = 1, 2, 3$), and $J_4^+ > 0$. We will also restrict attention, as discussed in Sec. I, to the particularly interesting case when $J_3 = J_2$. Since we may regard the exchange coupling constant $J_1$ as simply setting the overall energy scale, the relevant parameters are thus $(J_3/J_1 = )J_2/J_1 \equiv \alpha$, and $J_4^+/J_1 \equiv \delta$.

The honeycomb lattice itself is non-Braavais. It has a two-site unit cell, with two triangular sublattices 1 and 2, and triangular Bravais lattice vectors $a$ and $b$, as shown in Fig. 1(c). In terms of unit vectors $\hat{x}$ and $\hat{y}$ that define the $xz$ plane of the monolayers, we have $a = \sqrt{3}d\hat{x}$ and $b = \frac{1}{2}d(-\sqrt{3}\hat{x} + 3\hat{y})$, where $d$ is the NN spacing on the hexagonal lattice. Sites on sublattice 1 are at positions $R_i = ma + nb$, where $m,n \in Z$. Each unit cell of position vector $R_i$ on each layer comprises two sites, one at $R_i$ on sublattice 1 and the other at $R_i + d\hat{z}$ on sublattice 2. In Fig. 1(a), we also show the corresponding four sites of the AA-stacked bilayer unit cell.
In position space the Wigner-Seitz cell for the monolayer is simply the parallelogram formed by the trigonal Bravais lattice vectors \( \mathbf{a} \) and \( \mathbf{b} \). Equivalently, it may be chosen to be one of the primitive real-space hexagons of the lattice with side length \( 2\pi/(3\sqrt{3}d) \). Its first three corners are thus at positions \( \mathbf{K}^{(1)} = 2\pi/(3\sqrt{3}d) \hat{x} \), \( \mathbf{K}^{(2)} = 2\pi/(3\sqrt{3}d) \hat{x} + 2\pi/(3d) \hat{y} \), and \( \mathbf{K}^{(3)} = -2\pi/(3\sqrt{3}d) \hat{x} + 2\pi/(3d) \hat{y} \). The remaining three corners are at positions \( \mathbf{K}^{(i+3)} = -\mathbf{K}^{(i)} \), \( i = 1,2,3 \).

Let us first briefly review the situation for the model under consideration when \( \delta = 0 \) (i.e., for the monolayer). Along the line \( J_1 = J_2 = \alpha J_1 \) in the classical case there is a direct transition between the two collinear AFM phases at \( \alpha_{cl} = \frac{1}{2} \). These are the Néel phase shown in Fig. 1(c), which is the GS phase for \( \alpha < \alpha_{cl} \), and the striped phase shown in Fig. 1(d), which is the GS phase for \( \alpha > \alpha_{cl} \). Whereas the Néel phase on the honeycomb-lattice monolayer has all three NN pairs of spins antiparallel to one another at each site, the striped phase has only one NN pair antiparallel and the other two parallel to one another. Equivalently, the striped phase is composed of parallel ferromagnetic zigzag (or sawtooth) chains along one of the three equivalent honeycomb directions, with neighboring chains coupled antiferromagnetically. The striped state shown in Fig. 1(d) thus has two other equivalent states rotated with respect to it by \( \pm 120^\circ \) in the lattice plane.

As usual, the classical phases may generically be described by an ordering wave vector \( \mathbf{Q} \), together with a parameter \( \theta \) that measures the angle between the two spins in each monolayer unit cell \( i \) at position vector \( \mathbf{R}_i \). The classical spins, of length \( s \), are thus written as

\[
s_{i,i} = s \left[ \cos(\mathbf{Q} \cdot \mathbf{R}_i + \theta) \hat{x}_s + \sin(\mathbf{Q} \cdot \mathbf{R}_i + \theta) \hat{z}_s \right],
\]

where the index \( \rho \) labels the two sites in the unit cell, and \( \hat{x}_s \) and \( \hat{z}_s \) are two orthogonal unit vectors that define the plane of the spins. The angles \( \theta_{\rho} \) may be chosen with no loss of generality so that \( \theta_1 = 0 \) for spins on triangular sublattice 1 and \( \theta_2 = \theta \) for spins on triangular sublattice 2. In this description of the classical phases, the Néel phase shown in Fig. 1(c) has wave vector \( \mathbf{Q} = 0 \) and \( \theta = \pi \). Similarly, the striped phase shown in Fig. 1(d) has wave vector \( \mathbf{Q} = \mathbf{M}^{(2)} \) and \( \theta = \pi \), where \( \mathbf{M}^{(2)} = 2\pi/(3d) \hat{z} \) is the vector of the midpoint of the edge joining the corners of the hexagonal first Brillouin zone at positions \( \mathbf{K}^{(2)} \) and \( \mathbf{K}^{(3)} \). The two other inequivalent striped states are hence described by the wave vectors of the remaining inequivalent midpoints of the other two edges of the first Brillouin zone, and in each case now with \( \theta = 0 \). Thus, the other two striped states have wave vectors \( \mathbf{Q} = \mathbf{M}^{(1)} = \pi/(\sqrt{3}d) \hat{x} + \pi/(3d) \hat{y} \) (that is the midpoint of the edge joining corners \( \mathbf{K}^{(1)} \) and \( \mathbf{K}^{(2)} \)), and \( \mathbf{Q} = \mathbf{M}^{(3)} = -\pi/(\sqrt{3}d) \hat{x} + \pi/(3d) \hat{y} \) (that is the midpoint of the edge joining corners \( \mathbf{K}^{(3)} \) and \( \mathbf{K}^{(4)} \)).

Let us now compare this classical result for the monolayer (\( \delta = 0 \)) version of our model with the corresponding case when \( s = \frac{1}{2} \). In the spin-\( \frac{1}{2} \) case the classical transition point is split into two quantum critical points (QCPs) at \( \alpha_{cl} < \alpha_{cl} \) and \( \alpha_{cl} > \alpha_{cl} \), with a magnetically disordered paramagnetic GS phase in the intermediate region [53,54]. Lowest-order spin-wave theory, for example, provides the estimates [53] \( \alpha_{cl} \approx 0.29 \) and \( \alpha_{cl} \approx 0.55 \). By contrast, the more powerful, and potentially more accurate, method of Schwinger-boson mean-field theory (SBMFT) yields the estimates [53] \( \alpha_{cl} \approx 0.41 \) and \( \alpha_{cl} \approx 0.6 \). SBMFT also predicts a quantum disordered phase in the intermediate regime \( \alpha_{cl} < \alpha < \alpha_{cl} \), where a gap opens up in the bosonic dispersion and the spin-spin correlation function displays traces of Néel short-range order. These results are broadly confirmed by high-order CCM calculations [54], which yield the values \( \alpha_{cl} \approx 0.47 \) and \( \alpha_{cl} \approx 0.6 \).

Furthermore, CCM calculations of the plaquette valence-bond crystalline (PVBC) susceptibility [54] provide strong evidence for the intermediate paramagnetic phase to be a gapped state with PVBC order over the entire region.

Of special interest for the spin-\( \frac{1}{2} \) monolayer, the QPT at \( \alpha_{cl} \) between the Néel phase (that is the stable GS phase for \( \alpha < \alpha_{cl} \)) and the paramagnetic phase (that is the stable GS phase for \( \alpha_{cl} < \alpha < \alpha_{cl} \)) appears to be continuous, while that at \( \alpha_{cl} \) appears to be of first-order type [54]. Since the Néel and intermediate phases break different symmetries, the QPT at \( \alpha_{cl} \) is thus favored to be described by the scenario of deconfined quantum criticality [57]. In view of this rich scenario it seems of considerable interest to study the comparable spin-\( \frac{1}{2} \) \( J_1-J_2-J_3-J_1^\perp \) model on the AA-stacked honeycomb bilayer, where we now include AFM interlayer NN bonds of strength \( J_{1^\perp} > 0 \). Once again we will study the model here along the line \( J_1 = J_2 = \alpha J_1 \) with \( J_{1^\perp} = \delta J_1 \). Specific goals will be to study how the QCPs \( \alpha_{cl} \) and \( \alpha_{cl} \) now depend on the interlayer coupling parameter \( \delta \). To that end we will use the same theoretical technique, viz., the coupled cluster method (CCM), as has been used previously to describe accurately the corresponding monolayer case (\( \delta = 0 \)) [54].

Let us now turn to the corresponding case of the \( J_1-J_2-J_3-J_1^\perp \) model (with \( J_3 = J_2 \)), on the AA-stacked honeycomb-lattice bilayer, which we aim to study here for the spin-\( \frac{1}{2} \) case. At the classical (\( s \rightarrow \infty \)) level the inclusion of an NN interlayer coupling with strength \( J_{1^\perp} > 0 \) introduces no extra frustration, and its effect is essentially trivial. Simply the NN interlayer pairs of spins antialign, with each monolayer having the same Néel order for \( \alpha < \alpha_{cl} \) or ordered for \( \alpha > \alpha_{cl} \) as in the absence of interlayer pairing. By contrast, as we have discussed in detail in Sec. I, the spin-\( \frac{1}{2} \) case is expected to be of greater interest and subtlety, due to the expected formation of NN interlayer spin-singlet dimers in the large-\( \delta \) limit, where the GS phase will thus be a valence-bond crystalline (VBC) phase formed from interlayer dimers. This interlayer dimer VBC (IDVBC) phase will be gapped, by contrast to the gapless nature of the quasiclassical Néel and striped phases with magnetic LRO, where the magnon excitations are massless Goldstone modes. In the complete IDVBC phase (i.e., in the limit \( \delta \rightarrow \infty \)), the lowest-lying excited state is a spin-1 state formed by breaking a single NN interlayer dimer from a spin-singlet to a spin-triplet state. Thus, we expect the scaled triplet spin gap to be given, in the limiting case, by

\[
\frac{\Delta}{J_1} \overset{\delta \rightarrow \infty}{\longrightarrow} \delta.
\]

As we have noted above, the phase diagram of the model along the line \( \delta = 0 \) (i.e., for the monolayer) is already a rich
one, with two QCPs in the range $0 \leq \alpha \leq 1$ of the frustration
parameter, with one being continuous (and hence probably of a deconfined quantum critical nature) and the other being
of first-order type. Clearly, the enlargement of the model by
the addition of the extra parameter $\delta$ can only increase our
understanding of the quantum phase diagram, even for the
case of the monolayer. In particular, the two QCPs for the
limiting case $\delta = 0$ now become quantum critical lines (or
quantum phase boundary lines) in the $\alpha \delta$ plane. As a foretaste
of our overall results, one of our most important findings is
that these two phase boundary lines display a marked “avoided
crossing” behavior, with both curves consequently exhibiting
a distinct re-entrant nature. This finding by itself clearly throws
new light on the phase diagram of the monolayer ($\delta = 0$), as
discussed more fully in Secs. IV and V.

In order to examine the effect of interlayer coupling on the
QCPs $\alpha_c$, and $\alpha'_c$ of the spin-$\frac{1}{2}$ honeycomb-lattice monolayer
we will therefore present in Sec. IV results for both the
magnetic order parameter (i.e., the average local on-site GS
magnetization) $M$ and the triplet spin gap $\Delta$, for each of the
Néel and striped quasiclassical phases, as functions of both
parameters $\alpha$ and $\delta$. We utilize both perfectly ordered states as
model wave functions, on top of which we include quantum
fluctuations in a fully systematic formalism (viz., the CCM), as
we first demonstrate in Sec. III. In particular we show
how the CCM can be implemented in very high orders in a
well-defined and fully systematic approximation hierarchy, to
yield sequences of approximants for both $M$ and $\Delta$. We show
further how these sequences can then also be extrapolated,
in a controlled and stable manner, to the limit where the
corresponding wave functions are exact in principle. These
extrapolations are the only approximations made. In Sec. IV
we will show explicitly how the results for both $M$ and $\Delta$ yield
accurate and consistent estimates of the phase boundaries,
$\alpha_c(\delta)$ and $\alpha'_c(\delta)$, of the Néel and striped GS phases.

III. THE COUPLED CLUSTER METHOD

The CCM [58–69] is one of the very few size-extensive
and size-consistent techniques of quantum many-body theory. It
thereby provides results in the $N \to \infty$ limit (where $N$ is
the number of particles, i.e., lattice spins in our case) from
the outset, at all levels of approximation. Hence, no finite-
size scaling is ever required. Particularly apposite to the CCM
also is the fact that both the Goldstone linked-cluster theorem
and the very important Hellmann-Feynman theorem are also
preserved at every level of approximate implementation of
the formalism. The latter plays a large part in ensuring that
the method yields accurate, self-consistent, and robust results
for a variety of physical parameters for a given system. The
method has been applied very widely, yielding results of great
(and often unsurpassed) accuracy to systems as diverse as
finite nuclei [58], the electron gas (or jellium) [59,60], atoms
and molecules of interest in quantum chemistry [64], and a
broad spectrum of spin-lattice problems of interest in quantum
magnetism [15,41,68–94].

To initiate the CCM in practice one needs to choose a
suitable model (or reference) state to act as a generalized
vacuum state. The quantum correlations present in the exact
GS or excited-state (ES) wave function of the system are then
systematically incorporated on top of the model state in a
hierarchical scheme that becomes exact in some limit, which
is usually unattainable at particular levels of computational
implementation. Appropriate conditions for a state to be a
suitable CCM model state have been discussed extensively in
the literature [61,63,66–69]. For spin-lattice models, however,
all (quasi)classical states with perfect magnetic LRO are
suitable CCM model states. Hence, we use here both the Néel
and striped states shown in Figs. 1(c) and 1(d) respectively,
for each honeycomb-lattice monolayer, in each case with NN
interlayer spins on the $A$-stacked bilayer aligned antiparallel
to each other, as our two choices for CCM model state. We
will present independent sets of results in Sec. IV based on
both model states taken separately.

We shall only briefly review here some of the principal features and most important elements of the CCM, and refer
the reader to Refs. [41,58–69] for a fuller description. It is
very convenient to be able to treat each lattice spin in each
model state as being equivalent to one another. A simple way
to do so is to perform a separate passive rotation of each
such spin so that they all point in the same direction, say
downward (i.e., along the local negative $z_\parallel$ axis), in its own
local spin-coordinate frame. Accordingly, after such a choice
of local spin-coordinate frames has been made, each model
state takes the universal form $|\Phi \rangle = | \downarrow \downarrow \downarrow \cdots \downarrow \rangle$. Naturally,
the Hamiltonian also needs to be rewritten appropriately for
each such choice.

For a completely general quantum many-body system,
it exact GS energy eigenket $|\Psi \rangle$, where $H|\Psi \rangle = E|\Psi \rangle$, is
expressed in the exponentiated form,

$$|\Psi \rangle = e^S|\Phi \rangle ; \quad S = \sum_{I \neq 0} S_I C^+_I ; \quad (4)$$

that is a hallmark of the CCM. Its GS energy eigenbra counterpart $\langle \Psi |$, where $\langle \Psi |H = E\langle \Psi |$, is correspondingly
expressed in the CCM parametrization,

$$\langle \Psi | = \langle \Phi | S e^{-S} ; \quad S = 1 + \sum_{I \neq 0} S_I C^-_I , \quad (5)$$

where $C^-_I \equiv (C^+_I)^\dagger$ and $C^+_I \equiv 1$, the identity operator. The set
index $I$ here represents a multiparticle configuration, such that the set of states $\{C^+_I|\Phi \rangle\}$ completely spans the many-body
Hilbert space. The model state $|\Phi \rangle$ and the complete set of
mutually commuting, multiconfigurational creation operators
$\{C^+_I\}$ must be chosen so that the former is a fiducial vector
(or generalized vacuum state) with regard to the latter, in the
sense that they obey the conditions

$$\langle \Phi |C^-_J = 0 = C^+_I |\Phi \rangle , \quad \forall I \neq 0 , \quad (6)$$
as well as

$$[C^-_I, C^+_J] = 0 \quad \forall I, J . \quad (7)$$

The model state $|\Phi \rangle$ is chosen to be normalized, $\langle \Phi |\Phi \rangle \equiv 1$, and
the CCM parametrization of Eq. (4) ensures that the exact
GS wave function $|\Psi \rangle$ obeys the intermediate normalization
condition, $\langle \Psi |\Psi \rangle = 1$. Similarly, the CCM parametrization
of Eq. (5) for $\langle \Psi |$ ensures the automatic fulfillment of the
normalization condition $\langle \Psi |\Psi \rangle = 1$. In practice, it is also
convenient to orthonormalize the set of states $\{C^+_I|\Phi \rangle\}$, i.e., so
that they obey the relations
\[ \langle \Phi | C^+_I C^-_J |\Phi \rangle = \delta_{I,J}, \quad \forall I, J \neq 0, \] (8)
where \( \delta_{I,J} \) is a suitably generalized Kronecker symbol.

We note that Hermiticity clearly ensures that the destruction correlation operator \( \tilde{S} \) may formally be expressed in terms of its creation counterpart \( \tilde{S} \) via the relation
\[ \langle \Phi |\tilde{S} = \frac{\langle \Phi | e^S e^\tilde{S} |\Phi \rangle}{\langle \Phi | e^S e^\tilde{S} |\Phi \rangle}. \] (9)

A key feature of the CCM is that the constraint of Eq. (9) is not imposed explicitly. Rather, the operator \( \tilde{S} \) is formally decomposed independently of \( S \), as in Eq. (5). Naturally, in the exact limit, when all multiconfigurational clusters specified by the complete set \( I \) are retained, Eq. (9) would be exactly fulfilled. In practice, when approximations are made to restrict the set \( I \) to some manageable subset, Eq. (9) may only be approximately fulfilled. This manifest loss of Hermiticity, however, is more than compensated by the gain that the Hellmann-Feynman theorem is itself exactly fulfilled by the CCM parametrizations of Eqs. (4) and (5), even when the sums over the multiconfigurational indices \( I \) are restricted.

Turning now to the specific case of a quantum spin-lattice system, in the local spin-coordinate frames discussed above, where the CCM model state takes the universal form \( |\Phi\rangle = |\downarrow\downarrow\downarrow\cdots\downarrow\rangle \), the operator \( C^+_I \) can also now be chosen to have the universal form of a product of single-spin raising operators, \( \delta^+_I \equiv s^+_k + is^\dagger_k \). Thus, the set index \( I \) now takes the form of a set of site indices,
\[ I \equiv \{k_1, k_2, \ldots, k_n\}; \quad n = 1, 2, \ldots, 2sN \], (10)
in which no given site index \( k_i \) may appear more than \( 2s \) times. Correspondingly, the operator \( C^+_I \) creates a multispin configuration cluster,
\[ C^+_I \equiv s^+_k s^+_l \cdots s^+_m, \quad n = 1, 2, \ldots, 2sN \]. (11)

Clearly, all GS quantities may now be expressed wholly in terms of the CCM correlation coefficients \( \{S_I, \tilde{S}_I\} \). For example, the GS magnetic order parameter, which is simply the average local onsite magnetization, takes the form
\[ M = -\frac{1}{N} \langle \Phi | \tilde{S} \sum_{k=1}^N e^{-S_k} e^\tilde{S}_k |\Phi \rangle, \] (12)
where \( s^+_k \) is expressed in the local rotated spin axes described above. Thus, all that remains for the GS calculations is to calculate the coefficients \( \{S_I, \tilde{S}_I\} \).

Formally, this is done by minimizing the GS energy expectation functional,
\[ \tilde{H} = \tilde{H}[S_I, \tilde{S}_I] \equiv \langle \Phi |\tilde{S} e^{-S} He^S |\Phi \rangle, \] (13)
with respect to all coefficients \( \{S_I, \tilde{S}_I; \forall I \neq 0\} \). Extremization with respect to \( \tilde{S}_I \), using Eq. (5), trivially yields the relations
\[ \langle \Phi | C^+_I e^{-S} He^S |\Phi \rangle = 0, \quad \forall I \neq 0, \] (14)
which are a coupled set of nonlinear equations for the coefficients \( \{S_I\} \). Similarly, use of Eq. (4) and extremization of \( \tilde{H} \) with respect to \( S_I \) yield the relations
\[ \langle \Phi |\tilde{S} (e^{-S} He^S - E) C^+_I |\Phi \rangle = 0, \quad \forall I \neq 0, \] (15)
where we have also used the simple relation \( \{S, C^+_I\} = 0 \), which follows from the explicit CCM parametrization of Eq. (4) together with Eq. (7). Equation (15) is just a set of linear equations for the coefficients \( \{\tilde{S}_I\} \), once the coefficients \( \{S_I\} \) are input, having first been obtained by solving Eq. (14).

An ES energy eigenket \( |\Psi_e\rangle \), where \( H |\Psi_e\rangle = E_e |\Psi_e\rangle \) is similarly expressed in the CCM formalism in terms of a linear excitation operator, \( X^e \), as
\[ |\Psi_e\rangle = X^e e^S |\Phi \rangle; \quad X^e = \sum_{I \neq 0} \lambda^+_I C^+_I, \] (16)
as the analog of Eq. (4) for the GS counterpart. Using the obvious commutativity relation, \( [X^e, S] = 0 \), which follows from Eqs. (4), (7), and (16), it is easy to combine the GS and ES Schrödinger equations to yield the equation
\[ e^{-S} [H, X^e] e^S |\Phi \rangle = \Delta_e X^e |\Phi \rangle, \] (17)
where \( \Delta_e \) is the excitation energy,
\[ \Delta_e \equiv E_e - E. \] (18)
By taking the inner product of Eq. (17) with \( \langle \Phi | C^+_I \) and making use of Eq. (8), one may readily derive the set of equations
\[ \langle \Phi | C^+_I (e^{-S} He^S - E) X^e |\Phi \rangle = \Delta_e \lambda^+_I, \quad \forall I \neq 0. \] (19)
Once the operator \( (e^{-S} He^S - E) \) has been input into Eq. (19) from the solution to Eq. (14), Eq. (19) is simply a set of generalized linear eigenvalue equations for the ES kete-state CCM correlation coefficients \( \{\lambda^+_I\} \) and the excitation energy (eigenvalue) \( \Delta_e \).

Everything so far has been formally exact, and we turn now to where approximations may be needed for computational implementation of the CCM formalism. One possible source of approximation could involve the evaluation of the exponentiated operators \( e^{\pm S} \) that lie at the heart of the CCM parametrizations of Eqs. (4), (5), and (16). However, we note that in each of Eqs. (14), (15), and (19) that need to be solved for the CCM correlation coefficients \( \{S_I, \tilde{S}_I\} \), and \( \{\lambda^+_I\} \), these appear only in the combination \( e^{-S} He^S \) of a similarity transform of the system Hamiltonian. We have described in detail elsewhere (and see, e.g., Refs. [69,90,93] and references cited therein) how the otherwise infinite-order nested commutator expansion
\[ e^{-S} He^S = \sum_{n=0}^{\infty} \frac{1}{n!} [H,S]^n, \] (20)
where \( [H,S]_n \), the \( n \)-fold nested commutator, is defined iteratively as
\[ [H,S]_n = [[H,S]_{n-1},S]; \quad [H,S]_0 = H, \] (21)
actually terminates exactly in the present case at the term with \( n = 2 \). Similarly, all GS expectation values, such as the magnetic order parameter \( M \) of Eq. (12), may be evaluated without any truncations of the similarity transform \( e^{-S} s_k^+ e^S \).

Hence, the sole approximation made is in the choices of which subsets of multispin-flip configurations \( I \) to retain in the expansions of correlation coefficients for both the GS wave functions, as in Eqs. (4), (5), and the ES wave function considered, as in Eq. (16). In the present case we
restrict attention to the lowest-lying spin-triplet excitation, so that \( \Delta_n \) now becomes equal to the spin-triplet gap, which we denote by \( \Delta \). For both the GS and ES calculations we employ the well-known lattice animal-based subsystem (LSUBn) truncation hierarchy, which has been much studied and utilized for a wide variety of spin-lattice problems in the past (and see, e.g., Refs. [15,41,54-56,68,69,95] and references cited therein).

At the LSUBn level of approximation one retains all multispin-flip configurations \( f \) in the CCM correlation operators \( \mathcal{S}, \tilde{\mathcal{S}} \), and \( \mathbf{X}^n \) defined in Eqs. (4), (5), and (16), respectively, which are restrained to all distinct locales on the lattice that contain no more than \( n \) contiguous sites. A cluster of sites is said to be contiguous (or to form a lattice animal or polyomino) when each site of the cluster is NN to at least one other, taking into account the geometrical definition of NN sites employed. For the GS calculation we restrict the multispin-flip configurations at each LSUBn level to have \( s_j^z = 0 \), where \( s_j^z = \sum_{k=1}^{N} s_j^z_k \) defined in the local spin axes (i.e., before the local rotations discussed above have been performed).

Similarly, for the ES calculation of the triplet spin gap \( \Delta \), we restrict the comparable configurations to have \( s_j^z = 1 \). In both cases, we utilize the space- and point-group symmetries of the Hamiltonian and the CCM model state \( |\Phi\rangle \) being employed to reduce the number of fundamentally distinct configurations to a minimum, \( N_f(n) \). Since the number \( N_f(n) \) increases rapidly as a function of the truncation index \( n \), it soon becomes necessary to use supercomputing resources and massive parallelization, together with custom-made computer-algebra packages, to enumerate the independent cluster configurations retained and to derive the corresponding CCM equations at a given LSUBn level, and then finally to solve them [68,95].

For the present bilayer model, we are thereby able to perform both the GS and ES calculations reported in Sec. IV at LSUBn levels of approximation with \( n \leq 10 \). For the GS calculation, we have \( N_f(10) = 70\,118(175\,223) \) using the bilayer Néel (striped) states described previously as CCM model states. For the corresponding ES calculation of \( \Delta \), we have \( N_f(10) = 121\,103(320\,476) \) using the bilayer Néel (striped) states as CCM model states. Both numbers are larger for the striped state than the Néel state since the former has fewer symmetries than the latter. We note that for this model the geometric definition of contiguous sites for the LSUBn configurations simply corresponds to NN pairs connected by \( J_1 \) or \( J_1^\perp \) bonds.

Finally, to obtain estimates for our results in the formally exact limit, \( n \rightarrow \infty \), we need to extrapolate the raw LSUBn data. Such extrapolations thereby comprise the sole approximation that we make. While exact extrapolation rules are not known, much practical experience from applications of the LSUBn hierarchy to many different spin-lattice models has shown the widespread accuracy of the consistent use of rather simple schemes for the relevant physical parameters. In this respect, the magnetic order parameter \( M \) is of special interest, since two different schemes have been employed for differing situations.

Thus, for unfrustrated spin systems or for ones with only small amounts of frustration, an appropriate extrapolation scheme is found to be [15,41,55,73–76,79–82,84,86,87]

\[
M(n) = m_0 + m_1 n^{-1} + m_2 n^{-2},
\]

from fits to which we obtain the extrapolated LSUB\( \infty \) value \( m_0 \) for \( M \). By contrast, a more appropriate scheme for systems that exhibit a GS order-disorder QPT, or for phases whose order parameter \( M \) is zero or small, is [15,41,54-56,77,78,84–87,92–94]

\[
M(n) = \mu_0 + \mu_1 n^{-1/2} + \mu_2 n^{-3/2},
\]

which yields the respective LSUB\( \infty \) extrapolant \( \mu_0 \) for \( M \).

For the triplet spin gap \( \Delta \) an extrapolation scheme with leading power of \( n^{-1} \), like that in Eq. (22) for \( M \), has been found to give a very good fit to the LSUBn approximants \( \Delta(n) \) for both of the above cases of unfrustrated (or slightly frustrated) and highly frustrated systems [41,74,88–90,93,94],

\[
\Delta(n) = d_0 + d_1 n^{-1} + d_2 n^{-2},
\]

from fits to which we obtain the extrapolated LSUB\( \infty \) estimate \( d_0 \) for \( \Delta \).

Clearly, for each of the fits of Eqs. (22)-(24), it is best to use four or more fitting points (i.e., different \( n \) values of the LSUBn sequence) to obtain robust and the most reliable results. Furthermore, the LSUB2 result is expected \textit{a priori} to be too close to mean-field theory and too far from the exact \( n \rightarrow \infty \) limit to be used in each fit, if it can be avoided. For this reason, our preferred sets of LSUBn approximants for the fits are those with \( n = \{4,6,8,10\} \) in the present case where it is computationally infeasible to calculate results for \( n > 10 \).

We also note, however, that a \((4m - 2)/4m\) stagnation effect, where \( m \in \mathbb{Z}^+ \) is a positive integer, has been observed [86,87,93] in LSUBn sequences of CCM results for various physical parameters on frustrated honeycomb-lattice monolayers. Such stagnation implies that the two subsequences with \( n = 4m - 2 \) and with \( n = 4m \) need to be extrapolated separately from one another. In some cases corresponding adjacent pairs of curves from each subsequence (e.g., those with \( n = 2 \) and \( n = 4 \), or with \( n = 6 \) and \( n = 8 \)) even cross one another as some coupling parameter is varied. Such stagnation has been possibly attributed [93] to the fact that the honeycomb lattice comprises two interlocking triangular Bravais lattices, on each of which a more well-known \((2m - 1)/2m\) (i.e., odd/even) stagnation effect is commonly seen, exactly analogous to the same effect that is well understood in perturbation theory. Indeed, this is precisely the reason why LSUBn results with odd values of the truncation index \( n \) are not included in our CCM extrapolations here. In view of these prior observations on frustrated honeycomb-lattice monolayers, we shall also compare our results in Sec. IV between extrapolations based on the preferred sequences with \( n = \{4,6,8,10\} \) and those based on \( n = \{2,6,10\} \). The latter sequence is suboptimal in the two aspects that it both includes the LSUB2 result and is based on only three fitting points to extract three parameters. Nevertheless, it avoids mixing results from the two staggered subsequences.

**IV. RESULTS**

We first show in Fig. 2 our CCM results for the GS magnetic order parameter \( M \) of Eq. (12) in the Néel phase, as a function of the scaled interlayer exchange coupling constant, \( \delta \equiv J_1^\perp/J_1 \), for three particular representative values of the intralayer frustration parameter, \( \alpha \equiv J_2/J_1 \). In each case we...
show the “raw” LSUBn data with $n = 2, 4, 6, 8, 10$, based on the Néel state of Fig. 1(c) as the CCM model state, together with various LSUB∞ extrapolations. We note in particular that the two extrapolations based on the scheme of Eq. (23), but using the two respective LSUBn data sets with $n = 2, 6, 10$ and $n = [4, 6, 8, 10]$ give results in very close agreement with one another for all three values of $\alpha$ and for most values of $\delta$. Generally, the only exception, where there is some slight sensitivity to the extrapolation input data, is the joint region of the highest values of intralayer frustration (near to where Néel order disappears) and the lowest values of interlayer AFM coupling, as seen in Fig. 2(c), for example.

It is interesting to note that in each case the effect of turning on the interlayer coupling is first to enhance the stability of the Néel order, up to some particular value of $\delta$, which depends on the intralayer frustration parameter $\alpha$. Increasing $\delta$ beyond this value then leads to a decrease in the Néel order parameter $M$, out to some critical value $\delta_c(\alpha)$ at which $M \to 0$. Furthermore, we note that Néel order persists in the honeycomb-lattice monolayer ($\delta = 0$) for all values of the frustration parameter in the range $\alpha < \alpha_c(0)$. In this range, we find an upper critical value $\delta^u_c(\alpha)$ of the scaled interlayer exchange coupling constant, such that Néel order persists over the range $0 < \delta < \delta^u_c(\alpha)$. Very interestingly, however, as may be seen from Fig. 2(c), for example, for higher values of $\alpha$ in the range $\alpha_c(0) < \alpha < \alpha_c(\alpha)$, we find a re-entrant type of behavior in which Néel order now exists only over the range $\delta^u_c(\alpha) < \delta < \delta^l_c(\alpha)$, with $\delta^l_c(\alpha) > 0$. The corresponding upper and lower critical values coincide when $\alpha = \alpha_c^\star$, at which point we thus have $\delta^u_c(\alpha_c^\star) = \delta^l_c(\alpha_c^\star)$. Finally, Néel order is absent for $\alpha > \alpha_c^\star$, for all values of $\delta$.

The extrapolation scheme of Eq. (23), used for the LSUB∞(1) and LSUB∞(2) results shown in each of Figs. 2(a), 2(b), and 2(c), is certainly valid for all these cases when the frustration is appreciable, especially for systems close to a QCP as here, at which the magnetic order parameter vanishes. However, for unfrustrated systems or ones that are only slightly unfrustrated, the scheme of Eq. (22) provides a better fit to the LSUBn input data. Accordingly, we also show in Fig. 2(a) alone the LSUB∞(2a) extrapolation based on Eq. (22) and the input data set $n = [4, 6, 8, 10]$. This extrapolation is then valid for this case of zero intralayer frustration ($\alpha = 0$) only for a small range of values of the scaled interlayer coupling, $\delta \lesssim 0.2$, say.

In particular, the LSUB∞(2a) curve gives a value $M = 0.2741(1)$ for the unfrustrated honeycomb-lattice monolayer (i.e., with $\alpha = 0 = \delta$), where the quoted error is simply the least-squares error associated with the fit. The corresponding result using Eq. (22) and the input data set $n = [2, 6, 10]$ is $M = 0.2761$. There is clearly a small sensitivity associated with the LSUBn input data set used of about 1% or so. In this case ($\alpha = 0 = \delta$) alone, where “minus-sign problems” are absent, we may compare our CCM results with recent values obtained from two different quantum Monte Carlo (QMC) simulations, which yielded the corresponding respective values $M = 0.2681(8)$ [96] and $M = 0.2688(3)$ [97]. The agreement with our own CCM estimates is very good, and we expect a corresponding accuracy (i.e., of the order of 2%) for all the results that we present. Indeed, for the monolayer case, where we are also able to perform LSUBn calculations with $n = 12$, our corresponding extrapolated value, based on the data set $n = [8, 10, 12]$, again with only three fitting points, for example, is $M = 0.2715$, in even better agreement (1%) with the QMC result.

In Fig. 3, we show corresponding results for the GS magnetic order parameter $M$ of Eq. (12) in the striped phase to those shown in Fig. 2 for the Néel phase. Again, we show results for $M(\delta)$ for three particular representative values of the intralayer frustration parameter $\alpha$. In each case, we show the same “raw” LSUBn data with $n = 2, 4, 6, 8, 10$, but now based on the striped state of Fig. (d) as the CCM model state. We also display in Fig. 3 the same LSUB∞(1) and LSUB∞(2) extrapolations as those shown in Fig. 2, both of
FIG. 3. CCM results for the GS magnetic order parameter $M$ vs the scaled interlayer exchange coupling constant, $\delta \equiv J^i_1/J_1$, for the spin-$\frac{1}{2} J_1$-$J_2$-$J_3$-$J^i_1$ model on the bilayer honeycomb lattice (with $J_3 = J_2$ and $J_1 > 0$), for three selected values of the intralayer frustration parameter, $\alpha \equiv J_2/J_1$: (a) $\alpha = 1.0$, (b) $\alpha = 0.8$, and (c) $\alpha = 0.56$. Results based on the striped state as CCM model state are shown in $\text{LSUB} n$ approximations with $n = [2, 6, 8, 10]$, together with two corresponding $\text{LSUB} \infty(i)$ extrapolated results using Eq. (23) and the respective data sets $n = \{2, 6, 8, 10\}$ for $i = 1$ and $n = \{4, 6, 8, 10\}$ for $i = 2$.

which are based on the appropriate scheme of Eq. (23), but with the two respective $\text{LSUB} n$ input data sets $n = [2, 6, 10]$ and $n = [4, 6, 8, 10]$.

Again, we note that the two extrapolations are in remarkable agreement with each other in every case, despite the $\frac{(4m - 2)}{4m}$ staggering that has been discussed in Sec. III and which is now clearly visible in each of Figs. 3(a), 3(b), and 3(c), where it manifests itself most vividly as actual crossings of corresponding adjacent pairs of curves (viz., $\text{LSUB} 2$ and $\text{LSUB} 4$, corresponding to $m = 1$, and $\text{LSUB} 6$ and $\text{LSUB} 8$, corresponding to $m = 2$). Presumably, the very close agreement between the $\text{LSUB} \infty(1)$ and $\text{LSUB} \infty(2)$ extrapolations, the former of which takes the staggering into account and the latter of which does not, is related to the fact that the crossings occur only for unphysical values of $\delta$, far beyond the associated QCP at which the (extrapolated) striped order parameter has vanished.

Figure 3 exhibits a re-entrant behavior very similar to that seen in Fig. 2, and discussed above, for the Néel phase. Thus, striped order is present in the honeycomb-lattice monolayer ($\delta = 0$) for all values of the intralayer frustration parameter in the range $\alpha > \alpha_c(0)$. In this range we observe from Fig. 3 that as the AFM bilayer coupling is turned on, striped order persists over the range $0 < \delta < \delta^*_c(\alpha)$ of the scaled interlayer exchange coupling. However, now for somewhat lower values of $\alpha$ in the range $\alpha < \alpha^*_2 < \alpha < \alpha_c(0)$, we again observe a re-entrant behavior in which striped order reappears over the range $\delta^*_c(\alpha) < \delta < \delta^*_2(\alpha)$, with $\delta^*_c(\alpha) > 0$. Similar to the Néel case, these respective upper and lower critical values for the striped phase coalesce when $\alpha = \alpha^*_2$, such that $\delta^*_c(\alpha^*_2) = \delta^*_2(\alpha^*_2)$. Striped order is then absent for all values of $\delta$ for $\alpha < \alpha^*_2$.

The re-entrant behavior for both the Néel and striped phases is also demonstrated more graphically in Fig. 4. Here we show sequences of extrapolated results for the

FIG. 4. CCM results for the GS magnetic order parameter $M$ vs the scaled interlayer exchange coupling constant, $\delta \equiv J^i_1/J_1$, for the spin-$\frac{1}{2} J_1$-$J_2$-$J_3$-$J^i_1$ model on the bilayer honeycomb lattice (with $J_3 = J_2$ and $J_1 > 0$), for a variety of values of the intralayer frustration parameter, $\alpha \equiv J_2/J_1$, using (a) the Néel state and (b) the striped state as the CCM model state. In each case we show extrapolated results, obtained from using Eq. (23) with the corresponding $\text{LSUB} n$ data sets $n = [2, 6, 10]$.
magnetic order parameter of both phases as functions of $\delta$, for a variety of values of $\alpha$ in both cases. All of the extrapolations shown are based on the scheme of Eq. (23), used with the corresponding LSUB$n$ input data sets with $n = [2,6,10]$. With this particular extrapolation, the two QCPs for the monolayers are $\alpha_c(0) \approx 0.379$ and $\alpha_c(0) \approx 0.595$. These may be compared, for example, with the corresponding results [53], $\alpha_c(0) \approx 0.41$ and $\alpha_c(0) \approx 0.6$, from using a rotationally invariant version of Schwinger boson mean-field theory, which has proven itself to be a fairly accurate technique for taking quantum fluctuations into account. By contrast, lowest-order (or linear) spin-wave theory gives the less accurate results [53], $\alpha_c(0) \approx 0.29$ and $\alpha_c(0) \approx 0.55$. Our own corresponding CCM results for $\alpha_1^e$ and $\alpha_2^e$ are $\alpha_1^e \approx 0.487$ and $\alpha_2^e \approx 0.556$, again based on the extrapolation scheme of Eq. (23) used with the LSUB$n$ input data set with $n = [2,6,10]$.

We turn now to our respective results for the triplet spin gap $\Delta$. We first show in Fig. 5 results based on the Néel state as our CCM model state as a function of the scaled interlayer exchange coupling constant, $\delta \equiv J^\perp_2/J_1$, for the same three representative values of the intralayer frustration parameter, $\alpha \equiv J^\perp_2/J_1$. We note that the two sets of approximants, both based on Eq. (24) but using the two LSUB$n$ input data sets with $n = [2,6,10]$ and $n = [4,6,8,10]$, are in excellent agreement with one another. Both give results for $\Delta$ that are zero, within very small numerical errors, when Néel order is present, as expected, i.e., for $\delta < \delta_c^e(\alpha)$. Furthermore, the values obtained for $\delta_c^e(\alpha)$ are in good agreement with the independent values obtained from Fig. 2 for where the GS magnetic order parameter $M$ vanishes. It is clear too that in each case the nonmagnetic phase that opens up after the melting of Néel order is gapped, consistent with it having a VBC character.

Our corresponding CCM results for the triplet spin gap $\Delta$ based on the striped state as the model state are shown in Fig. 6. Once again we display results as a function of $\delta$, now for the same three representative values of $\alpha$ as were shown in Fig. 3 for the magnetic order parameter $M$. In this case the extrapolated LSUB$n$ results, based on the input LSUB$n$ data set $n = [2,6,10]$, give results that $\Delta$ is zero, within extremely small numerical errors, for all three values of $\alpha$ shown, over essentially the same ranges of values of $\delta$, viz., $0 < \delta < \delta_c^e(\alpha)$ for $\alpha > \alpha_c(0)$ and $\delta_c^e < \delta < \delta_c^e(\alpha)$ for $\alpha < \alpha < \alpha_c(0)$, for which the striped magnetic order parameter $M$ is positive in Fig. 3. By contrast, for the striped state, the LSUB$n$ extrapolation, based on the input LSUB$n$ data set with $n = [4,6,8,10]$, is definitely not as accurate or reliable as the LSUB$n$ extrapolation in this respect. This is surely due now to the $(4m-2)/4m$ staggering effect that is clearly visible in the LSUB$n$ sequences of results shown in Figs. 6(a)–6(c).

By way of further elucidation of the extrapolation of sequences of approximants that display a staggering as in Fig. 6, it is instructive to consider an analogous situation in $n$th-order perturbation theory (PT), for which exact extrapolation laws can be derived for various physical quantities. However, as is very well known, the even ($n = 2m$, where $m \in \mathbb{Z}^+$) and odd ($n = 2m - 1$) sequences of PT approximants involve an additional staggering effect, exactly as is also seen in corresponding CCM LSUB$n$ approximants. In both cases the even and odd sequences obey an extrapolation scheme of the same sort (i.e., with the same leading exponent), but one should not mix even and odd terms together in a single approximation scheme, unless the staggering is incorporated somehow, since the coefficients are not identical for both sequences. The explicit inclusion of the staggering is always very difficult to achieve in a robust manner, and hence in practice one always extrapolates only the even-order terms or the odd-order terms, as mentioned in Sec. III. For honeycomb-lattice models of the sort considered here, it has been observed previously, as discussed above, that there is an additional staggering in the even-order sequence of LSUB$n$ between those with $n = 4m$ and those with $n = 4m - 2$. This staggering can clearly be seen by visual inspection of the LSUB$n$ curves.
shown in Fig. 6. Once again, both subsequences still separately obey Eq. (24). If we do, however, despite the staggering, mix terms from the latter two subsequences, as in our LSUB∞(2) extrapolation scheme, we get a poorer fit, leading, for example, to the observed negative values for the spin gap in Fig. 6. Such an obviously incorrect result is simply due to the staggering effect not having been incorporated.

We note, with regard to our CCM spin gap results, that the LSUBn curves in both Figs. 5 and 6 clearly show a linear increase with δ for large values of this parameter. This is precisely as expected for an IDVBC phase, as expressed by Eq. (3). We note too that for true quantum critical behavior we expect the spin gap Δ to vanish (for a fixed value of the frustration parameter α, say) at some critical value δc, with a critical exponent ε and with κ a constant. Our LSUBn curves shown in Figs. 5 and 6, both for the “raw” curves with n finite and the extrapolated values with n → ∞, are clearly more consistent with a value ε > 1 (i.e., so that Δ vanishes with a zero slope, rather than the infinite slope expected for ε < 1) at both critical points δc(α) and δc2(α).

While the phase boundaries obtained from our CCM results for M and Δ for both quasiclassical magnetic phases are thus clearly in excellent agreement with each other, those obtained from M are surely more accurate. This is simply due to the respective shapes of the curves for M and Δ as functions of the parameters α and δ. Thus, the (extrapolated) curves for Δ, which are zero (within numerical errors) in the magnetically ordered phase, generally depart from zero (to indicate a gapped state) with zero slope. Thus, the estimates for the QCPs from the results for Δ have much larger associated errors than those obtained from the vanishing of the order parameter M, since the slope of the curve for M as a function of the corresponding coupling constant is generally nonzero at the points where M → 0.

Thus, finally, in Fig. 7 we show our best estimate for the T = 0 phase diagram of the spin-1/2 J1-J2-J3-J4 model on the AA-stacked bilayer honeycomb lattice with J1 > 0, J2 = J3 ≡ αJ1 > 0, J4 ⊥ J1 = δJ1 > 0. The leftmost (sky blue) and the rightmost (sea green) regions are the quasiclassical AFM phases with Néel and striped order respectively, while the central (gray) region is a gapped paramagnetic phase. The red cross (×) and the green plus (+) symbols are points at which the extrapolated GS magnetic order parameter M vanishes. These points are determined from extrapolating our LSUBn results for both quasiclassical AFM phases with Eq. (23), and using the data sets with n = [2, 6, 10], which overcome any errors associated with the (4m − 2)/4m staggering effects discussed above, as input. Points on the two (i.e., Néel and striped) quasiclassical AFM phases are marked by LSUBn states with δc(α) and δc2(α), respectively.

FIG. 6. CCM results for the triplet spin gap Δ (in units of J1) vs the scaled interlayer exchange coupling constant, δ ≡ J4 /J1, for the spin-1/2 J1-J2-J3-J4 model on the bilayer honeycomb lattice (with J1 = J2 and J1 > 0), for three selected values of the intralayer frustration parameter, α ≡ J3/J1: (a) α = 1.0, (b) α = 0.8, and (c) α = 0.56. Results based on the striped state as CCM model state are shown in LSUBn approximations with n = 2, 4, 6, 8, 10, together with two corresponding LSUB∞(i) extrapolated results using Eq. (24) and the respective data sets n = [2, 6, 10] for i = 1 and n = [4, 6, 8, 10] for i = 2.
stripped) phase boundaries are shown both at fixed values of $\alpha$ and fixed values of $\delta$. The former (indicated by green plus (+) symbols) are obtained from curves such as those shown in Fig. 4. They thereby represent the corresponding points $\delta_\alpha^\alpha(\alpha)$ [and also $\delta_\alpha^\alpha(\alpha)$ for fixed $\alpha$ in the range $\alpha_c(0) < \alpha < \alpha_c^\alpha$] for the Néel phase, and $\delta_\alpha^\alpha(\alpha)$ [and also $\delta_\alpha^\alpha(\alpha)$ for fixed $\alpha$ in the range $\alpha_c^\alpha < \alpha < \alpha_c(0)$] for the striped phase. The latter [indicated by red cross (x) symbols] are similarly obtained from corresponding extrapolated curves for $M$ as a function of $\alpha$ for various fixed values of $\delta$. They are hence the respective points $\alpha_c(\delta)$ for the Néel phase and $\alpha_c(\delta)$ for the striped phase.

One can clearly see from Fig. 7 that on both phase boundaries the two sets of critical points, obtained from the completely independent results at fixed values of $\alpha$ and fixed values of $\delta$, agree extremely well with one another. This is definite evidence for both the consistency and high accuracy of the extrapolation procedure that we have adopted.

Perhaps the most striking aspect of the $T = 0$ phase diagram is the marked “avoided crossing” behavior of the two re-entrant phase boundaries. We note that the major portions of the upper parts of both boundaries (i.e., for $0 \leq \alpha \leq 0.4$ for the Néel case and $\alpha \geq 0.6$ for the striped case) are quite well approximated as straight lines. Thus, if these (approximate) straight lines were then to be extended, they would cross, and the Néel line would intersect the $\delta = 0$ axis at a value rather close to the monolayer striped QCP at $\alpha_c(0)$, while the striped line would intersect the $\delta = 0$ axis at a value close to the monolayer Néel QCP at $\alpha_c(0)$.

V. DISCUSSION AND SUMMARY

We have studied a frustrated spin-$1/2$ $J_1$-$J_2$-$J_3$-$J_4$ Heisenberg antiferromagnet on an AA-stacked honeycomb lattice in the case when $J_1 > 0$, $J_3 = J_2 \equiv \alpha J_1 > 0$, and $J_4 \equiv \delta J_1 > 0$. In particular, we have used the CCM implemented to very high order of approximation to give an accurate description of its $T = 0$ quantum phase diagram in the window $0 \leq \alpha \leq 1$, $0 \leq \delta \leq 1$ of the $\alpha\delta$ plane. This window includes two quasiclassical phases with AFM magnetic order (viz., the Néel and stripped phases), plus an intermediate paramagnetic phase that exhibits VBC order of various types.

Within the studied window there are thus two phase boundaries, $\delta = \delta_\alpha(\alpha)$ [or, equivalently, $\alpha = \alpha_c(\delta)$], along which Néel order melts, and $\delta = \delta_\alpha^\alpha(\alpha)$ [or, equivalently, $\alpha = \alpha_c^\alpha(\delta)$], along which striped order melts. We have seen that these two boundaries exhibit a distinct “avoided crossing” type of behavior, with both displaying a consequent re-entrant property. In the $\alpha\delta$ window under study we found that Néel order thus exists only for values of $\delta$ in the range $\delta_\alpha^\alpha(\alpha) < \delta < \delta_\alpha^\alpha(\alpha)$. Furthermore, we found that, whereas $\delta_\alpha^\alpha(\alpha) = 0$ in the window for values of $\alpha$ in the range $\alpha < \alpha_c(0)$, $\delta_\alpha^\alpha(\alpha) > 0$ for values of $\alpha$ in the respective range $\alpha_c(0) < \alpha < \alpha_c^\alpha$, where $\delta_\alpha^\alpha(\alpha_c^\alpha) = \delta_\alpha^\alpha(\alpha_c^\alpha)$. Similarly, we also found that in the same window striped order exists only for values of $\delta$ in the range $\delta_\alpha^\alpha(\alpha) < \delta < \delta_\alpha^\alpha(\alpha)$. Compared to the Néel phase, we also observed for the striped phase that, whereas $\delta_\alpha^\alpha(\alpha) = 0$ in the window under study for values of $\alpha$ in the range $\alpha > \alpha_c(0)$, $\delta_\alpha^\alpha(\alpha) > 0$ for values of $\alpha$ in the corresponding range $\alpha < \alpha < \alpha_c(0)$, where $\delta_\alpha^\alpha(\alpha_c^\alpha) = \delta_\alpha^\alpha(\alpha_c^\alpha)$. Our best estimates for the extremal points of the two quasiclassical phases are found to be $\alpha_c^\alpha = 0.49(1)$ on the Néel phase boundary and $\alpha_c^\alpha = 0.56(1)$ on the striped phase boundary, where the errors are estimated from a sensitivity analysis of our extrapolation procedure for the magnetic order parameter $M$.

Comparable errors are expected along most of the two phase boundaries in Fig. 7. The most sensitive region, however, which is the only exception, is that close to the $\delta = 0$ axis for the Néel boundary, as we have already noted above. For example, we have obtained the extrapolated value $\alpha_c(0) \approx 0.379$ based on Eq. (23) when used with the LSUB$n$ input data set $n = \{2,6,10\}$. By contrast, in an earlier CCM analysis of the spin-$1/2$ $J_1$-$J_2$-$J_3$ honeycomb monolayer with $J_3 = J_2 \equiv \alpha J_1$ [54], corresponding extrapolated values were obtained of $\alpha_c(0) \approx 0.466$ when based on the LSUB$n$ input data set $n = \{6,8,10,12\}$ and $\alpha_c(0) \approx 0.448$ when based on the corresponding set $n = \{6,8,10\}$. This sensitivity is surely associated with the fact that the QPT at $\alpha_c(0)$ for the monolayer, from the Néel phase to the plaquette VBC (PVBC) phase, appears to be of continuous, deconfined type [54]. A sensitivity analysis yields our best estimate for this monolayer QCP to be at $\alpha_c(0) = 0.46(2)$. Interestingly, this value is now even closer to the point where one would estimate the striped boundary curve to intersect the $\delta = 0$ axis if its crossing with the Néel boundary curve would not be avoided.

By contrast, the QPT at $\alpha_c(0)$ for the monolayer, from the PVBC phase to be striped phase, appears to be of first-order type [54], and our CCM estimates for it are accordingly much less sensitive to the extrapolation LSUB$n$ input data set. Compared with our own value here of $\alpha_c(0) \approx 0.595$ from using the LSUB$n$ set $n = \{2,6,10\}$, an earlier CCM analysis of the spin-$1/2$ $J_1$-$J_2$-$J_3$ honeycomb-lattice monolayer with $J_3 = J_2 \equiv \alpha J_1$ [54] gave the corresponding value $\alpha_c(0) \approx 0.601$ based on both sets $n = \{6,8,10,12\}$ and $n = \{6,8,10\}$. Our best overall estimate is $\alpha_c(0) = 0.600(5)$.

It is perhaps worthwhile to end by pointing out why we can assert with confidence that our extrapolation procedure is indeed robust, since this is the sole approximation in all CCM calculations. First, the method has now been used in well over 100 different papers for a wide variety of frustrated quantum magnets (and see, e.g., Refs. [15,41,54–56,68–94] and references cited therein), where the same extrapolation schemes as used here have been utilized, and in virtually all of which the results obtained have been shown to be either the best or among the best available. Second, in all of the many cases in the literature cited, where it has been possible to compare results obtained from different LSUB$n$ input sets, it has been shown that the obtained extrapolants for all physical parameters agree with one another, typically to $\sim 1\%$ or better. Third, the same is true here in the limited cases where we can test it. For example, for the limiting case $\delta = 0$ of the monolayer, where LSUB$n$ calculations can additionally be done for the case $n = 12$, the fits to all parameters stay unchanged to the same level. Lastly, in all the limited (i.e., unfrustrated) cases where comparison can be made with the essentially exact results of large-scale QMC calculations (as cited here for the case $\alpha = 0 \equiv \delta$), the extrapolated CCM values for all physical parameters typically agree with the extrapolated QMC values (i.e., after finite-size scaling to the $N \to \infty$ limit), again to $\sim 1\%$ (or better). The real point here is that the present honeycomb bilayer model is particularly challenging due to the unavoidable $(4n - 2)/4m$
staggering effects discussed. While it is therefore true that we are thereby forced to use only a few (3 or 4) points in our fits, we can nevertheless be confident of their robustness for the reasons cited.

In conclusion, we have found that the CCM, when implemented to high orders of $\text{LSUB}n$ approximation and the results suitably extrapolated to the (exact) $n \to \infty$ limit, is capable of giving very accurate descriptions of the $T = 0$ quantum phase boundaries of this frustrated $AA$-stacked honeycomb bilayer model. In the light of this it would clearly also be of interest to use the method to study comparable bilayer models with the staggered Bernal $AB$ stacking.

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[95] We use the program package cccm of D. J. J. Farnell and J. Schulenburg [http://www-e.uni-magdeburg.de/jschulen/cccm/index.html].