# Phase diagram and spin correlations of the Kitaev-Heisenberg model: Importance of quantum effects

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We explore the phase diagram of the Kitaev-Heisenberg model with nearest neighbor interactions on the honeycomb lattice using the exact diagonalization of finite systems combined with the cluster mean field approximation, and supplemented by the insights from analytic approaches: the linear spin-wave and second-order perturbation theories. This study confirms that by varying the balance between the Heisenberg and Kitaev term, frustrated exchange interactions stabilize in this model either one of four phases with magnetic long range order: Néel phase, ferromagnetic phase, and two other phases with coexisting antiferromagnetic and ferromagnetic bonds, zigzag and stripy phase, or one of two distinct spin-liquid phases. Out of these latter disordered phases, the one with ferromagnetic Kitaev interactions has a substantially broader range of stability as the neighboring competing ordered phases, ferromagnetic and stripy, have very weak quantum fluctuations. Focusing on the quantum spin-liquid phases, we study spatial spin correlations and dynamic spin structure factor of the model by the exact diagonalization technique, and discuss the evolution of gapped low-energy spin response across the quantum phase transitions between the disordered spin liquid and phases with long range magnetic order.

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### I. INTRODUCTION

Frustration in magnetic systems occurs by competing exchange interactions and leads frequently to disordered spinliquid states [1–3]. Recent progress in understanding transition metal oxides with orbital degrees of freedom demonstrated many unusual properties of systems with active  $t_{2g}$  degrees of freedom—they are characterized by anisotropic hopping [4–8] which generates Ising-like orbital interactions [9–17], similar to the orbital superexchange in  $e_g$  systems [18,19]. Particularly challenging are 4*d* and 5*d* transition metal oxides, where the interplay between strong electron correlations and spin-orbit interaction leads to several novel phases [20,21]. In iridates the spin-orbit interaction is so strong that spins and orbital operators combine to new S = 1/2 pseudospins at each site [22], and interactions between these pseudospins decide about the magnetic order in the ground state.

The  $A_2$ IrO<sub>3</sub> (A = Na, Li) family of honeycomb iridates has attracted a lot of attention as these compounds have  $t_{2g}$ orbital degree of freedom and lie close to the exactly solvable S = 1/2 Kitaev model [23]. This model has a number of remarkable features, including the absence of any symmetry breaking in its quantum Kitaev spin-liquid (KSL) ground state, with gapless Majorana fermions [23] and extremely short-ranged spin correlations confined to nearest neighbors [24]. We emphasize that below we call a KSL also disordered spin-liquid states which arise near the Kitaev points in presence of perturbing Heisenberg interactions  $\propto J$ .

By analyzing possible couplings between the Kramers doublets it was proposed that the microscopic model adequate to describe the honeycomb iridates includes Kitaev interactions accompanied by Heisenberg exchange in the form of the Kitaev-Heisenberg (KH) model [25]. Soon after the experimental evidence was presented that several features of the observed zigzag order are indeed captured by the KH model [26–34]. Its parameters for  $A_2$ IrO<sub>3</sub> compounds are still under debate at present [35,36]. One finds also a rather unique crossover from the quasiparticle states to a non-Fermi-liquid behavior by varying the frustrated interactions [37]. Unfortunately, however, it was recently realized that this model is not sufficient to explain the observed direction of magnetic moments in Na<sub>2</sub>IrO<sub>3</sub>, and its extension is indeed necessary to describe the magnetic order in real materials [38,39]. For example, bond-anisotropic interactions associated with the trigonal distortions have to play a role to explain the differences between Na<sub>2</sub>IrO<sub>3</sub> and Li<sub>2</sub>IrO<sub>3</sub> [40], the two compounds with quite different behavior reminiscent of the unsolved problem of NaNiO<sub>2</sub> and LiNiO<sub>2</sub> in spin-orbital physics [19]. On the other hand, the KH model might be applicable in another honeycomb magnet  $\alpha$ -RuCl<sub>3</sub>, see, e.g., a recent study of its spin excitation spectrum [41].

Understanding the consequences of frustrated Heisenberg interactions on the honeycomb lattice is very challenging and has stimulated several studies [42–45]. The KH model itself is highly nontrivial and poses an even more interesting problem in the theory [25,34,46,47]: The Kitaev term alone has intrinsic frustration due to directional Ising-like interactions between the spin components selected by the bond direction [23]. In addition, these interactions are disturbed by nearest neighbor Heisenberg exchange which triggers long-range order (LRO) sufficiently far from the Kitaev points [25,34,46,47]. In general, ferromagnetic (FM) and antiferromagnetic (AF) interactions coexist and the phase diagram of the KH model is quite rich as shown in several previous studies [25,34,46–49]. Finally, the KH model has also a very interesting phase diagram on the triangular lattice [50–53]. These studies motivate better

understanding of quantum effects in the KH model on the honeycomb lattice in the full range of its competing interactions.

The first purpose of this paper is to revisit the phase diagram of the KH model and to investigate it further by comparing the exact diagonalization (ED) result [34] with the self-consistent cluster mean field theory (CMFT), supplemented by the insights from the linear spin-wave theory (LSWT) and the second-order perturbation theory (SOPT). The main advantage of CMFT is that it goes beyond a single site mean field classical theory and gives not only the symmetry-broken states with LRO, but partly includes quantum fluctuations as well, namely the ones within the considered clusters [43,54,55]. In this way the treatment is more balanced and may allow for disordered states in cases when frustration of interactions dominates.

We present below a complete CMFT treatment of the phase diagram which includes also the Kitaev term in MF part of the Hamiltonian and covers the entire parameter space (in contrast to the earlier prototype version of CMFT calculation on a single hexagon for the KH model [56]). Note that the CMFT complements the ED which is unable to get symmetry breaking for a finite system, but nevertheless can be employed to investigate the phase transitions in the present KH model by evaluating the second derivative of the ground state energy to identify phase transitions by its characteristic maxima [25,34]. The ED result can be also used to recognize the type of magnetic order by transforming to reciprocal space and computing spin-structure factor. The second purpose is to investigate further the difference between quantum KSL regions around both Kitaev points mentioned in Ref. [34] and LRO/KSL boundaries.

The paper is organized as follows: In Sec. II we introduce the KH model and define its parameters. In Sec. III we present three methods of choice: (i) the exact diagonalization in Sec. III A, (ii) the self-consistent CMFT in Sec. III B, and (iii) linear spin wave theory in Sec. III D. An efficient method of solving the self-consistence problem obtained within the CMFT is introduced in Sec. III C. The numerical results are presented and discussed in Sec. IV: (i) the phase transitions and the phase diagram are introduced in Sec. IVA, and (ii) the phase boundaries, the values of the ground state energies, and the magnetic moments obtained by different methods are presented and discussed in Secs. IV B and IVC, and (iii) we discuss the impact of the Kitaev interaction on different spin ordered states in Sec. IVC. Spin correlations obtained for various phases are presented in Sec. V. The dynamical spin susceptibility and spin structure factor are introduced and analyzed for different phases in Sec. VI. Finally, in Sec. VII we present the main conclusions and short summary. The paper is supplemented with the Appendix where we explain the advantages of the linearization procedure implemented on the CMFT on the example of a single hexagon.

# **II. KITAEV-HEISENBERG MODEL**

We start from the KH Hamiltonian with nearest neighbor interactions on the honeycomb lattice in a form,

$$\mathcal{H} \equiv K \sum_{\langle ij \rangle \parallel \gamma} S_i^{\gamma} S_j^{\gamma} + J \sum_{\langle ij \rangle} S_i \cdot S_j, \qquad (2.1)$$

where  $\gamma = x, y, z$  labels the bond direction. The Kitaev term  $\propto K$  favors local bond correlations of the spin component

interacting on the particular bond. The superexchange J is of Heisenberg form and alone would generate a LRO state, antiferromagnetic or ferromagnetic, depending on whether J > 0 or J < 0. We fix the overall energy scale,

$$J^2 + K^2 = 1, (2.2)$$

and choose angular parametrization:

$$K = \sin \varphi, \quad J = \cos \varphi,$$
 (2.3)

varying  $\varphi$  within the interval  $\varphi \in [0, 2\pi]$ . This parametrization exhausts all the possibilities for nearest neighbor interactions in the KH model.

While zigzag AF order was observed in Na<sub>2</sub>IrO<sub>3</sub> [28–32], its microscopic explanation has been under debate for a long time. The *ab initio* studies [35,57] give motivation to investigate a broad regime of parameters *K* and *J*, see Eqs. (2.3). Further motivation comes from the honeycomb magnet  $\alpha$ -RuCl<sub>3</sub> [41]. Note that we do not intend to identify the parameter sets representative for each individual experimental system, but shall concentrate instead on the phase diagram of the model Eq. (2.1) with nearest neighbor interactions only.

### **III. CALCULATION METHODS**

# A. Exact diagonalization

We perform Lanczos diagonalization for an N = 24-site cluster with periodic boundary conditions (PBC). This cluster respects all the symmetries of the model, including hidden ones. Among the accessible clusters it is expected to have the minimal finite-size effects.

#### B. Cluster mean field theory

A method which combines ED with an explicit breaking of Hamiltonian's symmetries is the so-called self-consistent CMFT. It has been applied to several models with frustrated interactions, including the Kugel-Khomskii model [54,55]. The method was also extensively used by Albuquerque *et al.* [43] as one of the means to establish the full phase diagram of the Heisenberg- $J_2$ - $J_3$  model on the honeycomb lattice.

Within CMFT the internal bonds of the cluster [connecting the circles in Fig. 1(a)] are treated exactly. The corresponding part  $H_{IN}$  of the Hamiltonian is the nearest neighbor KH Hamiltonian, Eq. (2.1). The external bonds that connect the boundary sites (•) with the corresponding boundary sites of periodic copies of the cluster ( $\Box$ ) are described by the MF part of the Hamiltonian,

$$H_{\rm MF} \equiv K \sum_{[ij]\parallel z} \langle S_i^z \rangle S_j^z + J \sum_{[ij]} \langle S_i^z \rangle S_j^z, \qquad (3.1)$$

where [ij] marks the external bonds. Since the ordered moments in the KH model align always along one of the cubic axes *x*, *y*, *z* (see, e.g., Ref. [25]), we have put

$$\langle \vec{S}_i \rangle \cdot \vec{S}_j \equiv \langle S_i^z \rangle S_j^z \tag{3.2}$$

in  $H_{\rm MF}$  to simplify the calculations.

The averages  $\langle S_i^z \rangle$  generate effective magnetic fields acting on the boundary sites of the cluster. The total Hamiltonian

$$\mathcal{H} \equiv H_{\rm IN} + H_{\rm MF} \tag{3.3}$$



FIG. 1. (a) 24-site cluster and the introduction of the mean fields. Gray (black) circles indicate internal (boundary) sites. In CMFT the internal bonds of the cluster are treated exactly while the external bonds crossing the cluster boundary (dashed) are treated on the MF level. The sites marked by  $\Box$  generate effective magnetic fields on the boundary sites  $\bullet$ . Labels x, y, and z stand for three inequivalent bond directions determining the active products  $S_i^{\gamma} S_i^{\gamma}$  in the Kitaev part of the Hamiltonian (2.1), e.g., bonds of x direction contribute with the  $S_i^x S_i^x$  product to the Hamiltonian, etc. The pseudospin axes used here are parallel to the cubic axes indicated in the top view of a single octahedron. (b) Unit cells: for honeycomb lattice (coinciding with a single hexagon of that lattice), for triangular lattice (inner dotted hexagon), and zigzag magnetic unit cell (dashed rectangle). Black and white circles stand for up/down spin and indicate one of three equivalent zigzag patterns. (c) Corresponding Brillouin zones and special q points for the lattice constant a = 1. The q vectors compatible with the 24-site cluster in (a) are also shown.

is diagonalized in a self-consistent manner, taking a slightly different approach than the one presented in Ref. [43]: Instead of starting with a random wave function our algorithm begins with expectation values  $\langle S_i^z \rangle_{in}$  on each boundary site *i* of the cluster. These can represent a certain pattern (zigzag, stripy, Néel, FM) or be set randomly to have a "neutral" starting point. After diagonalizing the Hamiltonian (3.3) (again by the ED Lanczos method) the ground state of the system is obtained and we recalculate the expectation values  $\langle S_i^z \rangle$  to be used in the second iteration. The procedure is repeated until self-consistency is reached.

### C. Linearized cluster mean field theory

A single iteration of the self-consistent MF calculation may be viewed as a nonlinear mapping of the set of initial averages



FIG. 2. (a) The values of  $\lambda$  obtained by the linearization of CMFT for an embedded cluster of N = 24 sites with fixed magnetic order patterns: FM, AF, stripy, and zigzag. Leading  $\lambda > 1$  indicates the order that sets in. The disordered KSL phases near  $\varphi = \pi/2$  and  $3\pi/2$  are indicated by red. (b) Second derivative of the ground state energy,  $-d^2 E_0(\varphi)/d\varphi^2$ , obtained by ED. Adopted from Ref. [34].

 $\{\langle S_i^z \rangle_{in}\}\$  to the resulting averages  $\{\langle S_i^z \rangle_{in}\}\$ . The self-consistent solution is then a stable stationary point of such a mapping. To find the leading instability, we may consider the case of small initial averages in the CMFT calculation and identify the pattern characterized by the fastest growth during the iterations. To this end we linearize the above mapping.

In the lowest order the mapping corresponds to the multiplication of the vector of the averages  $\{\langle S_i^z \rangle_{in}\}$  by the matrix,

$$F_{ij} = \frac{\partial \langle S_i^z \rangle_{\text{fin}}}{\partial \langle S_j^z \rangle_{\text{in}}},\tag{3.4}$$

where *i* and *j* run through the cluster boundary sites. During iterations, the patterns corresponding to the individual eigenvectors of the matrix *F* grow as  $\lambda^n$  after *n* iterations for a particular eigenvalue  $\lambda$ . The ordering pattern obtained by CMFT is then given by the eigenvector with largest  $\lambda_{max} > 1$ . In the quantum KSL regimes, all the eigenvalues are less than 1 and no magnetic order emerges. An example of linearized CMFT applied to a single hexagon with PBC can be found in the Appendix.

A modified version of this method, used to obtain Fig. 2(a), assumes a particular ordered pattern (Néel, zigzag, FM, or stripy phase) and uses a single spin average  $\langle S^z \rangle_{in}$  distributed along the boundary sites outside the cluster, with the signs consistent with this pattern. The resulting values,  $\langle S_i^z \rangle_{fin}$ , are then averaged correspondingly. In this case the matrix *F* is reduced to a single value  $\lambda$  plotted in Fig. 2(a). We observe that the largest eigenvalue either drops below 1 when the disordered

### D. Linear spin-wave theory

at a quantum phase transition to a different ordered phase.

The LSWT is a basic tool to determine spin excitations and quantum corrections in systems with LRO [58]. For systems with coexisting AF and FM bonds quantum corrections are smaller than for the Néel phase on the same lattice but are still substantial for S = 1/2 spins [59]. For the KH model the LSWT [25,29,34] has to be implemented separately for each of the four ordered ground states: Néel (N), zigzag (ZZ), FM, or stripy (ST). Then for a particular ground state the Hamiltonian is rewritten in terms of the Holstein-Primakoff bosons [29,60] and only quadratic terms in bosonic operators are kept. The spectrum of such a quadratic Hamiltonian is finally obtained using the successive Fourier and Bogoliubov transformations.

While the spin wave dispersion relations are usually of prime interest [25,29,34,60], there are also two other quantities which can easily be calculated using LSWT and which will be important in the discussion that follows: (i) the value of the total ordered moment  $\langle M \rangle$  per site, and (ii) the total energy per site  $\langle E \rangle$ . These observables are calculated in a standard way [58,59] and expressed in terms of the eigenvalues, i.e., spin-wave energies  $\omega_{\mathbf{k}\alpha}$ , and the eigenvector components  $\{v_{\mathbf{k}\alpha\lambda}\}$  of the bosonic Hamiltonian *before* the Bogoliubov transformation:

$$\langle M \rangle = S - \frac{1}{LV} \sum_{\alpha, \lambda = 1, \dots, L} \int_{\mathbf{k} \in BZ} |v_{\mathbf{k}\alpha, \lambda}|^2 d^2 \mathbf{k}, \qquad (3.5)$$

and

$$\langle E \rangle = E_{\rm cl} \left[ S^2 \to S(S+1) \right] + \frac{S}{2LV} \sum_{\alpha=1,\dots,L} \int_{\mathbf{k} \in BZ} \omega_{\mathbf{k}\alpha} \, d^2 \mathbf{k}, \qquad (3.6)$$

where the choice of the sign of the eigenvalues and the normalization of their eigenvectors is described in Ref. [58]. Here  $E_{cl}$  is the classical ground state energy per site, e.g.,

$$E_{\rm cl} = -JzS^2/2,$$
 (3.7)

with z = 3 for the Néel phase at K = 0 and S = 1/2 is the value of spin quantum number. *L* in Eqs. (3.5)–(3.6) is the number of the eigenvalues of the problem (spin-wave modes) and  $\alpha$  enumerates these modes. For all cases except for the zigzag order [25], the integrals go over the two-sublattice (L = 2) rectangular Brillouin zone (BZ) [61] with its volume  $V = 8\pi^2/3\sqrt{3}$  and  $-\pi/\sqrt{3} \le k_x \le \pi/\sqrt{3}$ ,  $-2\pi/3 \le k_y \le 2\pi/3$  (as already mentioned we assume the lattice constant a = 1). For the zigzag state L = 4 and the rectangular BZ can be chosen as:  $-\pi/\sqrt{3} \le k_x \le \pi/\sqrt{3}$  and  $-\pi/3 \le k_y \le \pi/3$  and its volume is  $V = 4\pi^2/3\sqrt{3}$ .

### **IV. QUANTUM PHASE TRANSITIONS**

# A. Phase diagram

Here we supplement the ED-based phase diagram for the KH model Eq. (2.1) established in Ref. [34] with the one obtained within CMFT. Figure 3 displays the phase boundaries obtained with ED [34], within CMFT, as well as classical



FIG. 3. T = 0 phase diagram for KH model. The outer ring is composed from ED data for the 24-site cluster, reproducing the result from Ref. [34] in the new parametrization, the middle ring shows CMFT results also for 24-site cluster and the inner black circle represents the classical result. The convention used for the angular parameter  $\varphi$  which determines coupling constants [see Eqs. (2.3)] is shown in the center of the inner circle. The colors represent particular phases, shown also as mini drawings next to suitable regions of the phase diagram. Starting from  $\varphi = 0$  green colored region corresponds to Néel order, red—KSL, yellow—zigzag order, dark blue—FM, red—KSL, light blue—stripy phase, and again green—Néel phase.

(Luttinger-Tisza) phase boundaries. The latter are included for completeness and to highlight the fact that the quantum fluctuations stabilize the KSL phases beyond single points, see below. To examine them in more detail it is instructive to analyze the data in Fig. 2(a) for the boundaries obtained from linearized CMFT and Fig. 2(b) for the peaks in the second derivative of energy,  $-d^2 E_0(\varphi)/d\varphi^2$ , giving phase boundaries in ED [34].

It is clearly visible that all the methods that include quantum fluctuations give quantum versions of the four classically established magnetic phases: Néel, zigzag, FM, and stripy. As the most important effect we note that when quantum fluctuations are included within a classical phase, the energy is generally lowered and that the emerging phase is expected to expand beyond the classical boundaries, but only in cases when a phase which competes with it has weaker quantum fluctuations. This implies that phases of AF nature will expand at the expense of the FM ones as the latter phases have lower energy gains by quantum fluctuations (which even vanish exactly for the FM order at K = 0 and J < 0).

We summarize the phase boundaries obtained within different methods in Table I. One finds substantial corrections to the quantum phase transitions which follow from quantum fluctuations. These corrections are quite substantial in both KSLs at the Kitaev points (K = +1,  $\varphi = \frac{1}{2}\pi$  and K = -1,

TABLE I. Phase boundaries for the KH model, parameterized by the angle  $\varphi$  (in units of  $\pi$ ), see Eqs. (2.3). Columns: classical Luttinger-Tisza approximation, second-order perturbation theory (SOPT), exact diagonalization (ED), and self-consistent cluster mean field theory (CMFT).

| Boundary    | Classical | SOPT  | ED    | CMFT  |
|-------------|-----------|-------|-------|-------|
| Néel/KSL    | 0.5       | 0.492 | 0.494 | 0.496 |
| KSL/zigzag  | 0.5       | 0.507 | 0.506 | 0.505 |
| zigzag/FM   | 0.75      | 0.813 | 0.814 | 0.825 |
| FM/KSL      | 1.5       | 1.463 | 1.448 | 1.478 |
| KSL/stripy  | 1.5       | 1.530 | 1.539 | 1.519 |
| stripy/Néel | 1.75      | 1.705 | 1.704 | 1.699 |

 $\varphi = \frac{3}{2}\pi$ , first column of Table I). Indeed, in the classical approach massively degenerate ground states exist just at isolated points, but they are replaced by disordered spin-liquid states that extend to finite intervals of  $\varphi$  when quantum fluctuations are included, see the second, third, and fourth column in Table I. The expansion of Néel and zigzag phases beyond classical boundaries is given by particularly large corrections and is well visible.

The most prominent feature in the phase diagram described above is however the difference in size between two KSL regions, already addressed before using ED [34] and also visible now in the CMFT data. Therefore, the CMFT result supports the claim from Ref. [34] that the stability of KSL perturbed by relatively small Heisenberg interaction depends on the nature of the phases surrounding the spin liquid and the amount of quantum fluctuations that they carry. In the following we discuss the above issues more thoroughly, examining: (i) ground state energy curves emerging from ED, CMFT, SOPT within the linked cluster expansion and LSWT, (ii) the ordered moment given by various methods, (iii) the spin-spin correlation functions, and (iv) the spin structure factor as well as the dynamical spin susceptibility in the vicinity of the Kitaev points.

#### **B.** Quantum corrections: Energetics

We start the discussion of quantum corrections to the energy of the ordered phases by noting that, even though it properly captures finite order parameters, the CMFT looses quantum energy on the external bonds and would therefore not provide a reliable estimate of the ground-state energy. However, if one calculates instead the energy based on the correlations on the bonds of the central hexagon, the estimate is significantly improved. Here we choose the energy obtained using the ED calculations [see Fig. 4(a)] as a reference value because of all the bonds treated in an exact manner. This observation is also supported by the fact that the ED phase boundaries were roughly confirmed by tensor networks (iPEPS) [49] and density matrix renormalization group (DMRG) results [48]: The iPEPS phase boundaries agree with ED for AF KSL/LRO transitions and the boundaries between different LRO phases differ only slightly from those found in ED (iPEPS: zigzag/FM— $0.808\pi$ , stripy/Néel— $1.708\pi$ ). For FM KSL/LRO transition however the iPEPS result deviates more, i.e., KSL/stripy—1.528 $\pi$ . On the other hand, DMRG bound-



FIG. 4. (a) Comparison between ground state energies per site obtained using various methods: classical Luttinger-Tisza approximation (dashed black), SOPT (solid red), LSWT (dashed red), ED for 24-site cluster (solid blue, see Ref. [34] for this result in a different parametrization), and CMFT (energy given by the central hexagon, solid green). (b) Ordered moment obtained from CMFT (solid green line for the central hexagon, dashed green line for the value for intermediate and boundary sites) and LSWT (dashed red line).

aries agree perfectly with ED and due to four-sublattice dual transformation [10,25] one can reproduce the FM/zigzag as well as FM/KSL boundaries. Only the extent of the AF spin-liquid phase cannot be extracted from this result, but that is already confirmed by iPEPS.

Figure 4(a) shows a quite remarkable agreement between the energy values and critical values of  $\varphi$  obtained by the simplest SOPT [25] and our reference ED results. This suggests that this analytical method can be utilized to get better insight to the quantum contributions to the ground state energy. For a phase X with LRO, the energy per site  $\mathcal{E}_X$ , written as a sum of the classical energy  $E_{cl}$  and the quantum fluctuation contribution  $\Delta E_X$ , is obtained as:

$$\mathcal{E}_{\rm N} = -\frac{1}{8}(K+3J) - \frac{1}{16}(K+3J),$$
 (4.1)

$$\mathcal{E}_{ZZ} = -\frac{1}{8}(K-J) - \frac{1}{16}(K-J),$$
 (4.2)

$$\mathcal{E}_{\rm FM} = +\frac{1}{8}(K+3J) + \frac{1}{16}\frac{K^2}{K+2J},$$
 (4.3)

In addition, to get the LRO/KSL phase boundary points in Table I, we estimate the energy of the KSL phase as

$$\mathcal{E}_{\text{KSL}} \simeq \frac{3}{2} (K+J) \langle S^{\gamma} S^{\gamma} \rangle_{\text{Kitaev}},$$
 (4.5)

using the analytical result for the Kitaev points [24],  $\langle S^{\gamma} S^{\gamma} \rangle_{\text{Kitaev}} \approx \pm 0.131.$ 

The two spin-liquid phases in the phase diagram of the KH model differ strongly in their extent, despite the formal equivalence of the FM (K = -1) and AF (K = 1) Kitaev points provided by an exact mapping of the Kitaev Hamiltonian [23]. As mentioned earlier, this is due to the fact that the two KSLs compete with LRO phases of a distinct nature. Here we give a simple interpretation based on the strength of the quantum corrections of the LRO phases estimated using Eqs. (4.1)–(4.4). Later, in Secs. V and VI we illustrate the different nature of the transitions between FM and AF KSL and the surrounding it LRO phases in terms of spin correlations and spin dynamics.

Let us now compare the quantum fluctuation contribution and the classical one. For the LRO phases surrounding the AF spin liquid—Néel and zigzag—we always have  $\Delta E/E_{cl} = \frac{1}{2}$ as deduced from Eqs. (4.1) and (4.2), i.e., only  $\frac{2}{3}\mathcal{E}_N$  and  $\frac{2}{3}\mathcal{E}_{ZZ}$ are found in the classical approach. This guarantees that the quantum phase transition between these two types of order occurs at the same value of  $\varphi = \pi/2$  in SOPT and in the classical approach that do not capture the spin-liquid phase in between these ordered states, see Fig. 4(a). In contrast, the phases neighboring to the FM spin liquid-FM and stripywould reach the value of  $\Delta E/E_{\rm cl} = \frac{1}{2}$  only at the FM Kitaev point with J = 0 and away from this point the contribution of quantum fluctuations decreases rapidly allowing for a large extent of the FM spin-liquid phase. Note that both these latter phases contain a point which is exactly fluctuation free-for the FM phase when frustration is absent (K = 0), and for the stripy phase it is related to the FM one by the interaction transformation [39] at K = -2J.

Moving to the CMFT energy analysis one should also keep in mind that within the CMFT method the external bonds between  $\langle S_i^z \rangle$  and  $S_i^z$  do not include quantum fluctuations fully. This implies a worse estimate of the energy (of the whole cluster) for regions of the phase space that allow quantum fluctuations. As a consequence the region of stability of FM spin-liquid phase is smaller than that obtained in the ED. Significantly better energy estimate is given by the central hexagon, for which all the bonds experience exact interactions. As a result, this CMFT energy curve [green line in Fig. 4(a)] lies almost as close to ED energy as the SOPT one. Finally, the estimates obtained from LSWT, which represents a harmonic approximation to the quantum fluctuations, are not as good as those from central hexagon via CMFT and SOPT, see dashed red lines in Fig. 4(a). As expected, the energy obtained from the LSWT agrees well with ED curve for phases with less quantum fluctuations, FM and stripy phase, and starts to diverge when these phases are unstable beyond quantum phase transitions within Néel and zigzag phases.

### C. Quantum corrections: Ordered moment

As usual, getting the correct value of the ordered moment turns out to be a more difficult task than estimating the ground state energy. This is primarily due to the fact that the ED does not capture the symmetry-broken states and the ordered moment can only be indirectly extracted from the  $m^2$ ; moreover, the SOPT may not be reliable here. Hence, we are mostly left with the results obtained with CMFT and LSWT. We discuss the corresponding data [shown in Fig. 4(b)] together with the several values given already in the literature.

Let us begin with the Heisenberg AF point  $\varphi = 0$ : Here it is expected that the ordered moment should be strongly reduced by quantum fluctuations. LSWT estimates the ordered moment value at 0.248 [61]. Similar values were extracted from  $m^2$  in quantum Monte Carlo (0.268 [62–64]) and ED (0.270 [43]) calculations. In the last case however the authors admit that the set of clusters for finite size scaling was chosen so as to make the best agreement with quantum Monte Carlo. Another method-series expansion (high order perturbation theory) [47] sets ordered moment value at a somewhat higher value of 0.307. While all the above results seem roughly consistent, CMFT value obtained from the boundary sites seems to stand out (0.374 for  $\varphi = 0$ ). Nevertheless, the central-hexagon value (0.330 for  $\varphi = 0$ ) lies much closer to the results from the methods mentioned above. Moreover, one should note that the ordered moment estimated from  $m^2$  for 24-site cluster ED equals 0.45 [43] which is above the CMFT value. This suggests that at this point the finite size scaling is important.

Before moving to the frustrated regime we briefly mention that the trivial ordered moment value at  $\varphi = \pi$  is here correctly reproduced by both CMFT and LSWT. Besides, for the regions around the fluctuation-free FM (and stripy) point the ordered moments predicted by CMFT and LSWT also match. Following the ground state energy analysis, LSWT gives the correct result because quantum fluctuations contribution is small compared to the classical state. The further one moves towards the Kitaev points, however, the more incorrect the LSWT approximation should be because of the strong reduction of the ordered moment due to increasing frustration.

In contrast, the lack of quantum fluctuations on the external bonds generates systematic errors within CMFT except for FM and stripy phases. The ordered moment obtained from the boundary sites experiences the errors discussed above. However, the ordered moment values for intermediate sites and the central hexagon become largely reduced in the whole Néel and zigzag regions due to the fact that for the internal part of the cluster the fluctuations are fully included. Still, the best estimate comes from the central hexagon where quantum fluctuations on the bonds are included and CMFT gives more realistic results than LSWT in frustrated parts of the phase diagram. Here it is also important to stress that the series expansion captures correctly the fluctuation-free point at  $\varphi = \pi$  (FM) and  $\varphi = -\arctan 2$  (stripy) and predicts a broader region of the FM KSL phase [47]. The order parameter is also qualitatively correctly estimated and is reduced more to  $m \simeq 0.3$  for both Néel and zigzag phases [47]. However, while the ordered moment values obtained by CMFT are consistent with the four-sublattice dual transformation, the ordered moment data from the high-order perturbation theory [47] are not as the values of ordered moment differ at the points connected by the mapping. Unfortunately the largest difference appears near the FM LRO/KSL boundaries. This observation uncovers certain shortcomings of the high-order perturbation theory.

#### D. Quantum corrections: Naive interpretation

Let us conclude the discussion of the quantum corrections with the following more general observation: Developing the argumentation presented by Iregui, Corboz, and Troyer [49], the dependence of the quantum corrections to the energy and to the ordered moment on the angle  $\varphi$  suggests that the Kitaev interaction is less "compatible" with the FM/stripy ground states than with the Néel/zigzag ones. This can be understood in the simple picture of the KH model on a four-site segment of the honeycomb lattice consisting of three bonds attached to a selected lattice site, as presented below.

Starting with  $\varphi = \pi$  (FM ground state, e.g. along the *z* quantization axis), increasing  $\varphi$  leads to gradual increase of the FM Kitaev term which favors ferromagnetically aligned spins along the *x*, *y*, and *z* quantization axes for the *x*, *y*, and *z* directional bonds, respectively. It can easily be seen that, e.g., for the *x* bond, the eigenstate of the FM Kitaev-only Hamiltonian on that bond  $(|\uparrow_x\uparrow_x\rangle)$  has a 25% overlap with the FM ground state,  $|\langle\uparrow_z\uparrow_z|\uparrow_x\uparrow_x\rangle|^2 = \frac{1}{4}$ . While again a similar situation happens for the *y* bond, the overlap between such states for the *z* bond is maximal, i.e., these states are identical (we assume the same phase factors 1).

Next, we perform a similar analysis for  $\varphi = 0$  and firstly assume that we deal with a *classical* Néel ground state,  $|\uparrow_z \downarrow_z\rangle$ . In this case for the "unsatisfied" bonds from the point of view of the increasing AF Kitaev interaction we also obtain that the eigenstate of the AF Kitaev-only Hamiltonian  $(|\uparrow_x \downarrow_x \rangle)$  on that bond has a 25% overlap with the classical Néel ground statee.g.,  $|\langle \uparrow_z \downarrow_z | \uparrow_x \downarrow_x \rangle|^2 = \frac{1}{4}$ . However, this situation changes once we consider that the spin quantum fluctuations dress the classical Néel ground state. This can be best understood if we assumed the unrealistic but insightful case of very strong quantum fluctuations destroying the classical Néel ground state: then for the x bond a singlet could be stabilized and the overlap between such a state and the state "favored" by the Kitaev term increases to 50%:  $|\langle 0|\uparrow_x\downarrow_x\rangle|^2 = \frac{1}{2}$ . This suggests that the Néel ground state, which contains quantum spin fluctuations, is more "compatible" with the states "favored" by the Kitaev terms than the FM ground state, resulting in more stable values of ordered moment for Néel phase. It seems that the above difference is visible in CMFT data but not in LSWT ones. We shall discuss this issue further by analyzing spin correlations below.

#### V. SPIN CORRELATIONS

Additional information about the ground state is given by spin-spin correlation functions. In Fig. 5(a) one can observe isotropic stable  $\langle S_i^{\gamma} S_j^{\gamma} \rangle$  correlations in almost the entire AF phase (with  $\langle S_i \cdot S_j \rangle \approx -0.36$  for  $\varphi = 0$ ), while for the FM phase the anisotropy quickly develops when moving away from the FM Heisenberg point  $\varphi = \pi$  (here  $\langle S_i \cdot S_j \rangle$  reaches



FIG. 5. (a) Spin correlations  $\langle S_i \cdot S_j \rangle$  obtained within ED for the bonds between nearest neighbors (black line), spin correlations of the components active in the Kitaev interaction,  $\langle S_i^{\gamma} S_j^{\gamma} \rangle$  (blue line), and complementary spin components,  $\langle S_i^{\tilde{\gamma}} S_j^{\tilde{\gamma}} \rangle$  (red line). Below further neighbor spin correlations  $|\langle S_i \cdot S_j \rangle|$  are shown (colors correspond here to different neighbors). ED: (b) near the AF spin-liquid phase, and (c) for the angle  $\varphi$  interval including the FM spin-liquid phase. CMFT—the neighborhood of the: (d) AF spin-liquid, and (e) FM spin-liquid region.

the classical value 0.25). This again demonstrates that the AF (and zigzag) phase is more robust and uniform than the FM (and stripy) phase.

Moreover, spin-spin correlations allow us to confirm the disordered regions around the Kitaev points as critical cases of quantum spin liquid [65]. At the Kitaev points (J = 0) we observe the expected undisturbed KSL pattern: non-zero values of nearest neighbor correlations between spin components active in the Kitaev interaction [blue curve in Fig. 5(a)] and vanishing correlations between complementary components (red curve). In contrast, the next nearest and further neighbor correlations disappear, see Figs. 5(b) and 5(c). While moving away from the Kitaev points the absolute values of the correlations enter the regions of slow growth—these are signatures of the critical spin-liquid phases and they look

similar in AF and FM spin liquid cases. At some point however proceeding further results in rapidly growing absolute values which mark KSL/LRO boundaries.

Furthermore, Figs. 5(b) and 5(c) prove that there is a qualitative difference between the two spin-liquid regimes. This is observed in the rapid growth of spin correlations at the onset of LRO: The steplike jump visible in Fig. 5(b) contrasts with the smoother crossover seen in Fig. 5(c). Below we investigate this distinct behavior by analyzing the dynamical spin susceptibility for various available phases. After Fourier transformation of the *z*-component correlations, we obtain the spin structure factor to be discussed in the context of the spin susceptibility also in Sec. VI.

As a supplement we present the further neighbor spin correlations obtained via CMFT [Figs. 5(d) and 5(e)]. One should remark that within KSL the averages  $\langle S_i^z \rangle$  are 0 and CMFT is thus equivalent to ED for an isolated cluster (open boundary conditions). This leads to stronger finite size effects and larger inhomogeneity of the correlations. Nevertheless, considering the central part of the cluster, the emergence of the longer-range correlations away from the Kitaev point presented in Figs. 5(d) and 5(e) is almost identical to that calculated by ED, see Figs. 5(b) and 5(c).

### VI. SPIN SUSCEPTIBILITY AND EXCITATIONS IN THE VICINITY OF THE KITAEV POINTS

Below we study the spin dynamics within the KH model by analyzing the dynamical spin susceptibility at T = 0,

$$\chi_{\alpha\alpha}(\boldsymbol{q},\omega) = i \int_0^\infty \langle \Phi_0 | \left[ S_{\boldsymbol{q}}^\alpha(t), S_{-\boldsymbol{q}}^\alpha(0) \right] | \Phi_0 \rangle \, \mathrm{e}^{i\,\omega t} \, \mathrm{d}t, \quad (5.1)$$

with the Fourier-transformed spin operator defined via

$$S_{\boldsymbol{q}}^{\alpha} = \frac{1}{\sqrt{N}} \sum_{\boldsymbol{R}} \mathrm{e}^{-i\boldsymbol{q}\cdot\boldsymbol{R}} S_{\boldsymbol{R}}^{\alpha} \,, \tag{5.2}$$

and  $|\Phi_0\rangle$  denoting the cluster ground state. For  $\omega > 0$ , the imaginary part of  $\chi(\boldsymbol{q},\omega)_{\alpha\alpha}$  reads as

$$\chi_{\alpha\alpha}^{\prime\prime}(\boldsymbol{q},\omega) = -\mathrm{Im} \langle \Phi_0 | S_{\boldsymbol{q}}^{\alpha} \frac{1}{\omega + E_{\mathrm{GS}} - \mathcal{H} + i\delta} S_{-\boldsymbol{q}}^{\alpha} | \Phi_0 \rangle,$$
(5.3)

which can be conveniently expressed as a sum over the excited states  $\{|\nu\rangle\}$ ,

$$\chi_{\alpha\alpha}^{\prime\prime}(\boldsymbol{q},\omega) = \pi \sum_{|\nu\rangle} \left| \langle \nu | S_{-\boldsymbol{q}}^{\alpha} | \Phi_0 \rangle \right|^2 \delta(\omega - E_{\nu}), \qquad (5.4)$$

where the excitation energy  $E_{\nu}$  is measured relative to the ground state energy  $E_{GS}$ . We have evaluated  $\chi_{\alpha\alpha}(\boldsymbol{q},\omega)$  by ED on a hexagonal cluster of N = 24 sites. In the ED approach, the exact ground state of the cluster  $|\Phi_0\rangle$  is found by Lanczos diagonalization, the operator  $S^{\alpha}_{-\boldsymbol{q}}$  is applied, and the average of the resolvent  $1/(z - \mathcal{H})$  is determined by Lanczos method using normalized  $S^{\alpha}_{-\boldsymbol{q}}|\Phi_0\rangle$  as a starting vector [66].

In our case of the KH model, the calculation generally requires a relatively large number of Lanczos steps (up to one thousand) to achieve convergence of the dense high-energy part of the spectrum. Having the advantage of being exact, the method is limited by the q vectors accessible for a finite cluster



FIG. 6. (a) Dynamical spin susceptibility  $\chi''(q,\omega)$  obtained by ED near the AF KSL phase at the characteristic wave vector of the AF order,  $q = \Gamma'$ . (b) The same for the zigzag wave vector q = M. (c) Brillouin zone portraits of the spin-structure factor  $\langle S_{-q}^z S_q^z \rangle$  at  $\varphi = 87.5^\circ$ , 90°, and 92.5° (interpolated from the ED data). The inner hexagon is the Brillouin zone of the honeycomb lattice; the outer one corresponds to the triangular lattice with the missing sites filled in. (d), (e) The same as in panels (a), (b) but for the interval containing the FM ( $q = \Gamma$ ) and stripy (q = X) phase. (f) Brillouin zone portraits of the spin-structure factor obtained at  $\varphi = 255^\circ$ , 270°, and 285°.

and compatible with the PBC, and by finite-size effects due to small N. These concern mainly the low-energy part of  $\chi''$  and lead, e.g., to an enlarged gap of spin excitations in LRO phases of AF nature. Nevertheless, a qualitative understanding can still be obtained.

The evolution of numerically obtained  $\chi''(q, \omega)_{\alpha\alpha}$  (5.4) with varying  $\varphi$  is presented in Figs. 6(a) and 6(b) for the region including AF spin-liquid phase, as well as in Figs. 6(d) and 6(e) for the region including the FM spin-liquid phase. The transitions are well visible at the characteristic q vectors of the individual LRO phases. The structure factor pattern, see Figs. 6(c) and 6(f), changes accordingly between the sharply peaked one in LRO phases and a wavelike form characteristic for nearest neighbor correlations in the spin-liquid phases.

After entering the spin-liquid phase, further changes of the spin response are very different for the AF and FM case. In the

AF case, there is a sharp transition—a level crossing for our cluster, so that the ground state changes abruptly. The original intense pseudo-Goldstone mode as well as many other excited states become inactive in the spin-liquid phase. The observed low-energy gap in  $\chi''(q, \omega)_{\alpha\alpha}$  varies only slightly with  $\varphi$ .

In contrast, when entering the FM spin-liquid phase the excitation that used to be the gapless magnon mode is characterized by a gradually increasing gap which culminates at the Kitaev point. Starting from the Kitaev point, the gradual reduction of the low-energy gap in  $\chi''(q,\omega)_{\alpha\alpha}$  due to the Heisenberg perturbation manifests itself by a development of finite spin correlations beyond nearest neighbors (already reported in Fig. 2 of Ref. [25]) and an increase of the static susceptibility to the magnetic field Zeeman coupled to the order parameter of the neighboring LRO phase. This susceptibility then diverges at the transition point (see also Fig. 3 of Ref. [25]).

## VII. SUMMARY AND CONCLUSIONS

In the present paper we report a study of the phase diagram of the Kitaev-Heisenberg model by a combination of exact diagonalization and cluster mean field theory (CMFT), supplemented by the insights from linear spin-wave theory and the second-order perturbation theory. Both methods allowed to stabilize previously known phases with long range order: Néel, zigzag, FM, and stripy. Moreover, the ordered moment analysis provided by cluster mean field approach demonstrates Néel-zigzag and FM-stripy connections described before [34]. Compared to the previous CMFT studies utilizing N = 6site cluster (see Ref. [56] or the Appendix), we have used a sufficiently large cluster of N = 24 sites preserving the lattice symmetries and improving the ratio between internal and boundary bonds. This led to a balanced approach which allowed us to treat both ordered and disordered (spin-liquid) states on equal footing.

As the main result, the present study uncovers a fundamental difference between the onset of broken symmetry phases in the vicinity of Kitaev points with antiferromagnetic or ferromagnetic interactions. While the spin liquids obtained at K = +1 and K = -1 are strictly equivalent and can be transformed one into the other in the absence of Heisenberg interactions (at J = 0), spin excitations and quantum phase transitions emerging at finite J are very different in both cases. For the antiferromagnetic Kitaev spin liquid phase  $(K \simeq 1)$ one finds that a gap opens abruptly in  $\chi''(q,\omega)$  at  $q = \Gamma'$ and q = M when the ground state changes to the critical Kitaev quantum spin liquid. This phase transition is abrupt and occurs by level crossing. In contrast, for ferromagnetic spin liquid  $K \simeq -1$  the gaps in  $\chi''(q,\omega)$  at  $q = \Gamma$  and q = Xopen gradually from the points of quantum phase transition from ordered to disordered phase. With much weaker quantum corrections for ordered phases in the regime of ferromagnetic Kitaev interactions, the spin liquid is more robust near K = -1as a phase that contains quantum fluctuations and survives in a broader regime than near K = 1 when antiferromagnetic Kitaev interactions are disturbed by increasing (antiferromagnetic or ferromagnetic) Heisenberg interactions. This behavior is reminiscent of the ferromagnetic Kitaev model in a weak magnetic field [65].

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## APPENDIX: COMPARISON BETWEEN CMFT AND LINEARIZED CMFT FOR A SINGLE HEXAGON

Here we compare linearization results for a single hexagon with full CMFT to see how well linearized CMFT performs as a shortcut method. It is important to realize that this cluster is not compatible with stripy or zigzag order because of their four-site magnetic unit cell, see Fig. 1(b), and they are suppressed within vast regions of  $\varphi$  compared to the 24-site case. The size of the system allows for quick CMFT computations and enables detailed comparison between the two approaches. Moreover, specific problems linked to the above incompatibility make the N = 6-site cluster a good test case to illustrate the linearized CMFT.

Following the procedure described in Sec. III C, 6 eigenvalues  $\lambda_i$  are produced for each value of  $\varphi$  parameter. The corresponding spin patterns are inferred by inspecting the eigenvectors. Only the patterns associated with  $\lambda_i > 1$  are able to grow during iterations and eventually stabilize as a



FIG. 7. Full linearized CMFT result for a single hexagon. Blue lines represent all emerged positive eigenvalues  $\lambda$ , while maximal  $\lambda$  larger than 1 is indicated in red.

0.0

0



π

ø

3π/2

2π

FIG. 8. (a) Spin patterns obtained for a single hexagon by CMFT. From the left: Néel, zigzag, FM, and stripy. (b) Phase diagram for a single hexagon determined by  $|\langle S_i^z \rangle|$ . Red and blue sites (see inset) are nonequivalent in the present CMFT due to the approximation given by Eq. (3.2) which generates the terms  $\propto J$  that add to the Kitaev term only on the vertical bonds  $\langle ij \rangle \parallel z$  in the MF part of the Hamiltonian (2.1).

π/2

self-consistent solution of full CMFT. Comparison of both methods presented in Figs. 7 and 8 provides the phase diagram for a single hexagon: Néel phase for  $\varphi \in [0,0.5)\pi$ , KSL for  $\varphi = \frac{\pi}{2}$ , zigzag phase for  $\varphi \in (0.5,0.555)\pi$ , disordered region I for  $\varphi \in (0.555,0.864)\pi$ , FM phase for  $\varphi \in (0.864,1.5)\pi$ , KSL for  $\varphi = \frac{3}{2}\pi$ , stripy phase for  $\varphi \in (1.5,1.62)\pi$  (linearization),  $\varphi \in (1.5,1.64)\pi$  (CMFT), disordered region II for  $\varphi \in (1.62, 1.684)\pi$  (linearization) and  $\varphi \in (1.64, 1.684)\pi$  (CMFT), and Néel phase for  $\varphi \in (1.684, 2]\pi$ . In contrast to N = 24 cluster the two spin-liquid regions are replaced by single points  $\varphi = \frac{\pi}{2}$  and  $\varphi = \frac{3}{2}\pi$ .

Striking difference between phase diagrams for 24-site and 6-site clusters is the reduction of the zigzag and stripy phases and the emergence of two regions of disorder indicated by two gray-shaded regions. Here all  $\lambda_i < 1$  and no spin pattern is strong enough to stabilize. Zigzag pattern emerges from CMFT with random initial values of  $\langle S_i^z \rangle$  without additional help. Stripy pattern however is more difficult to catch. As one can see in Fig. 7, two different  $\lambda_i$  corresponding to two stripy patterns exchange at  $\varphi = 1.568\pi$ . Unfortunately, huge parasitic oscillations make these patterns extremely difficult to stabilize within CMFT. These stem from a large negative  $\lambda_i$  that previously corresponded to FM pattern and decreased rapidly for  $\varphi > 1.5\pi$ . If one recalls that the equivalent of one iteration in the linearized version of CMFT is in fact multiplication by  $\lambda_i$ , one can easily see that large negative  $\lambda_i$  would cause oscillations with an exponentially growing amplitude when performing the iterations of the self-consistent loop. To overcome this issue we introduce a damping into a self-consistent loop by taking  $(1 - d)\langle S_i^z \rangle_{\text{fin}} + d\langle S_i^z \rangle_{\text{in}}$  as the new averages. Here d < 1 is a suitably chosen damping factor. With this modification CMFT produces one finite stripy order suggested by linearization. However since the parasitic negative  $\lambda_i$  grows enormously in magnitude as we approach the phase boundary an extreme damping has to be included making the phase boundary hard to determine by using CMFT.

In conclusion, it is evident that the ordered patterns suggested by linearization were reproduced by CMFT within regions dictated by the maximal  $\lambda_i > 1$ . Moreover, the linearized procedure indicated possible difficulties with stabilizing stripy phases that had to be cured by a strong damping introduced into the self-consistent loop.

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