Second-order Peierls transition in the spin-orbital Kumar-Heisenberg model

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We add a Heisenberg interaction term $\infty\lambda$ in the one-dimensional $SU(2)\otimes XY$ spin-orbital model introduced by Kumar. At $\lambda=0$ the spin and orbital degrees of freedom can be separated by a unitary transformation leading to an exact solution of the model. We show that a finite $\lambda>0$ leads to spontaneous dimerization of the system which in the thermodynamic limit becomes a smooth phase transition at $\lambda\to0$, whereas it remains discontinuous within the first-order perturbation approach. We present the behavior of the entanglement entropy, energy gap, and dimerization order parameter in the limit of $\lambda\to0$ confirming the critical behavior. Finally, we show the evidence of another phase transition in the Heisenberg limit, $\lambda\to\infty$, and give a qualitative analytical explanation of the observed dimerized states both in the limit of small and large λ .

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I. INTRODUCTION

A Peierls transition being a distortion of the periodic lattice of a one-dimensional (1D) crystal has been known for more than 80 years [1]. Peierls' theorem states that a 1D equally spaced chain with one electron per ion is unstable to the dimerization, i.e., a more favorable configuration can be achieved if every ion moves closer to its one neighbor and further away from the other. This is a purely kinetic effect related to the nesting of the 1D Brillouin zone. Such dimerization involves charge and lattice degrees of freedom but a similar process can be observed in the spin-orbital models [2,3] where the interaction within the orbital sector dimerize the spin order. Such an effect was observed by a neutron scattering in the perovskite vanadate YVO₃ [4] as theoretically described by Horsch and coworkers [5]. Similarly, for an iridate triangular-lattice superconductor $Ir_{1-x}Pt_xTe_2$ the photoemission and model studies have revealed that an orbitally induced Peierls effect governs the charge and orbital instability in this compound [6]. A strict, 1D spin-orbital analog of the Peierls' dimerized state was found in NaTiSi₂O₆ [7], as shown by Raman scattering measurements, where the dimerized spin state at low temperature is driven by the orbital fluctuations at high temperature. A quite different situation emerges when the orbital fluctuation do not drive the spins' dimerization but the orbital order makes the electrons' hopping quasi-1D so that the classical Peierls transition can occur [8]. Such an orbital-selective Peierls dimerization was found in the t_{2g} spinel MgTi₂O₄ by an optical measurement. On the other hand it was shown that the notion of the Peierls instability also applies to the strongly interacting electrons coupled to the phonons within the Holstein model [9] which allows one to explain the orbital order in the undoped manganite LaMnO₃ [10].

Spin-orbital models are crucial from the point of view of frustrated magnetism [11–13] as the orbital interactions, being typically of lower symmetry than the SU(2)-symmetric spin interactions, can lead to frustration in both sectors even on a square lattice. This often happens in the transition-metal

oxides. When the 3d orbitals are partly filled, electrons localize due to large on-site Coulomb interaction and superexchange between magnetic ions includes both spin and orbital degrees of freedom that are strongly interrelated [14]. The orbital degeneracy leads then in many cases to a dramatic increase of quantum fluctuations [15], which may trigger exotic order [16,17], or may stabilize a spin liquid [18,19] when different states compete near a quantum critical point. In general these fluctuations make the ground states of the spin-orbital models entangled, as it happens in the archetypal Kugel-Khomskii model in one-, two-, three-dimensional and bilayer cases, respectively [16,17,20,21]. Similarly, in the $S = 1/2 \text{ SU}(2) \otimes \text{SU}(2)$ chain [22,23], both ground state [24] and excited states [25] are entangled and the $S = 1 \text{ SU}(2) \otimes$ SU(2) chain which plays a prominent role in the vanadium perovskites [13,26]. In exceptional cases such 1D models can be solved exactly, for example, at the SU(4) point [27] or for a valence-bond state [28] of alternating spin and orbital singlets similar to the Majumdar-Ghosh state in a 1D J_1 - J_2 spin chain [29], but even in these situations the spins and orbitals cannot be separated from each other.

In real materials the symmetry between spin and orbital interactions is absent. Orbital interactions generically have lower symmetry than spin ones [30], being usually Isingor XY-like [31]. The XY case is quantum and in general the orbitals cannot be separated from the spins [32]. Therefore the 1D SU(2) \otimes XY model introduced by Kumar [33] is very exceptional because (i) it is exactly solvable and (ii) by a change of basis, the S = 1/2 spins decouple from the orbitals in an *open* chain. The orbital interactions remain formally unchanged but the spin ones are gauged away. The spins then appear free and the ground state has large degeneracy $(2^L \text{ for chain length } L)$ [33]. It is worth mentioning that spin-orbital separation has been also recently demonstrated for Sr₂CuO₃ [34] although this separation occurs in the physical Hilbert space of spin and orbitals, which is not the case for the Kumar model.

Quite surprisingly, the properties of the Kumar model are determined by topology. When the chain is closed and the

model becomes periodic the exact solution still exists, as shown in Ref. [35], but the spins can no longer be gauged away completely. Contrarily, the gauge is accumulated on the closing orbital bond so that the whole spin sector enters the orbital problem only through the boundary condition, in analogy to Aharonov-Bohm magnetic flux through the ring. This partially lifts the macroscopic degeneracy of the ground state of the open chain and splits the manifold of the 2^L states into a multiplet of low-lying states. These are the states with topological excitations of a nonlocal nature having quadratic dispersion in the total quasimomentum carried by the spins [35]. Such a multiplet of states connected by the nonlocal excitations is a characteristic feature of another model originating from the orbital exchange, namely the two-dimensional orbital compass model (OCM) [36], which was considered, inter alia, in the context of topologically protected qubits [37]. From this point of view it was important to study how the small admixture of the SU(2) Heisenberg interactions, which is unavoidable in the physical implementations employing arrays of Josephson junctions, will modify the multiplet structure in OCM [38].

In this paper we introduce an extension of the Kumar model by including a Heisenberg term, which removes the degeneracy in the spin sector. It is worth noting that another possibility for the extension of the Kumar model to perturb the orbital degrees of freedom, e.g., by adding an orbital Heisenberg or Ising term, which could lead to a higher symmetry [27]. As we will see further on, this model is no longer exactly solvable but the Kumar gauge transformation still can be applied. As a result the pure Kumar term gets completely or partially decoupled depending on the topology of the system but the Heisenberg term is subject to the spin permutation that depends on the configuration of orbitals in their canonical basis. In other words, under the transformation the NN spin interaction gets smeared out and this smearing is governed by the delocalization of the up/down orbitals caused by the quantumness of the XY orbital interactions. This situation is interesting because in this way the spin interactions can get frustrated—the smeared interaction remains AF but now the further neighbors are involved. On the other hand, as the pure Kumar model leaves the spins practically completely uncorrelated, we expect that any small admixture of spin interactions λ will order the spins in some way. This is a favorable situation as it will enable us to use perturbation expansion in the manifold of degenerate ground states of the pure Kumar model. To examine the region, where the coupling between the spins cannot be treated as a small perturbation, we use the density matrix renormalization group method (DMRG) [39–43], which gives an accurate description of the ground state in one dimension. This also provides an independent way to verify the perturbational results.

The paper is organized as follows. In Sec. II we introduce the Kumar-Heisenberg (KH) model, in Sec. III we show the exact solution of the periodic KH model based on Ref. [35], in Sec. IV we apply this solution to the ferromagnetic, $\lambda < 0$, case and get the exact ground state, and in Sec. V we explain the spin dimerization within a first-order expansion in a small $\lambda > 0$ which is a valid description in the limit of short chains. In Sec. VI we explain why this limit cannot describe an infinite system where a second-order phase transition occurs, as shown in Sec. VII. In Sec. VIII we describe a dimerized state and

give a mean-field mechanism of the orbital dimerization for large λ . In this section we also show the evidence of a second phase transition in the limit of $\lambda \to \infty$. The conclusions are presented in Sec. IX and in Appendix we give further details on the effective spin couplings introduced in Sec. V.

II. KUMAR-HEISENBERG MODEL

The Hamiltonian of the KH model reads

$$\mathcal{H} = \mathcal{H}_0 + \mathcal{V},\tag{1}$$

where

$$\mathcal{H}_0 = J \sum_{l=1}^{L} X_{l,l+1} (\tau_l^+ \tau_{l+1}^- + \tau_l^- \tau_{l+1}^+)$$
 (2)

is the Kumar model and

$$\mathcal{V} = \lambda \sum_{l=1}^{L} \vec{\sigma}_{l} \vec{\sigma}_{l+1} \tag{3}$$

is the Heisenberg term. Here τ 's are orbital Pauli matrices and $X_{i,j} = (1 + \vec{\sigma}_i \vec{\sigma}_j)/2$ is a *spin transposition* operator, defined by another set of Pauli matrices σ . In the case of periodic boundary conditions (PBC) $\vec{\sigma}_{L+1} = \vec{\sigma}_1$ and $\tau_{L+1} = \tau_1$. For open boundary conditions (OBC) $\vec{\sigma}_{L+1} = 0$ and $\tau_{L+1} = 0$.

The Hamiltonian can be transformed into the base where \mathcal{H}_0 appears as a purely orbital contribution. For this we use the unitary operator,

$$\mathcal{U} = \prod_{l=1}^{L-1} \left[\frac{1 - \tau_{l+1}^z}{2} + \frac{1 + \tau_{l+1}^z}{2} \chi_{l+1,l} \right],\tag{4}$$

where $\chi_{l+1,l}$ is a *spin permutation* operator composed of the spin transpositions $X_{i,j}$:

$$\chi_{l+1,l} = X_{l+1,l} X_{l,l-1} \dots X_{3,2} X_{2,1}. \tag{5}$$

Now according to Ref. [35] the transformed Hamiltonian takes a form,

$$\mathcal{H}' = \mathcal{U}^{\dagger} \mathcal{H} \, \mathcal{U} = \mathcal{H}'_0 + \mathcal{V}', \tag{6}$$

where

$$\mathcal{H}'_{0} = J \sum_{l=1}^{L-1} (\tau_{l}^{+} \tau_{l+1}^{-} + \text{H.c.}) + J (R_{1}^{(1)} R_{1}^{(2)} \tau_{1}^{+} \tau_{L}^{-} + \text{H.c.}), (7)$$

and

$$\mathcal{V}' = \lambda \sum_{l=1}^{L} \vec{\sigma}_l \vec{\sigma}_{l+1}.$$
 (8)

Here $R_1^{(1)}$ is a cyclic permutation of spins at sites l = 1, ..., N by 1 site and $R_1^{(2)}$ is the same permutation of spins at sites l = N + 1, ..., L:

$$R_{1}^{(1)}: \{\vec{\sigma}_{1},...,\vec{\sigma}_{N}\} \to \{\vec{\sigma}_{2},...,\vec{\sigma}_{N},\vec{\sigma}_{1}\}, R_{1}^{(2)}: \{\vec{\sigma}_{N+1},...,\vec{\sigma}_{L}\} \to \{\vec{\sigma}_{N+2},...,\vec{\sigma}_{L},\vec{\sigma}_{N+1}\}.$$
(9)

Here *N* is a good quantum number of "up" $(\tau_l^z = +1)$ orbitals:

$$N = \sum_{l=1}^{L} \frac{1 + \tau_l^z}{2}.$$
 (10)

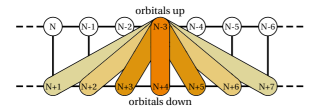


FIG. 1. (Color online) The schematic view of the bonds connecting the two spin subsystems (upper and bottom legs of the ladder) in the first-order Hamiltonian in Eq. (17). The spin bonds connecting site l=4 of the orbital-up subsystem with sites of the orbital-down subsystem are marked. Their color intensity increases with the strength of the coupling $J_{i,j}$.

The primed spins $\vec{\sigma}_l$ are entangled spin-orbital operators:

$$\vec{\sigma}_l' = |\vec{\tau}^z\rangle \vec{\sigma}_{f(l,\vec{\tau}^z)} \langle \vec{\tau}^z|, \tag{11}$$

with $|\vec{\tau}^z\rangle = |\tau_1^z, \tau_2^z, \dots, \tau_L^z\rangle$ and a permutation of chain sites,

$$f(l, \vec{\tau}^z) = l + \frac{1}{2} \left(1 + \tau_l^z \right) (1 - l) + \sum_{p=l+1}^L \frac{1}{2} \left(1 + \tau_p^z \right). \tag{12}$$

 $\vec{\sigma}_l'$ are the original spins $\vec{\sigma}_l$ after a permutation of the chain sites dictated by the orbital configuration in the τ_l^z eigenbasis.

III. SOLUTION OF THE KUMAR MODEL

The Hamiltonian \mathcal{H}'_0 is exactly solvable [35]. On all bonds but the closing one it involves only orbital degrees of freedom and even on the closing bond the spin part affects only the boundary conditions via the cyclic permutations $R_1^{(1,2)}$. These can be diagonalized and turned to the phase factors $e^{i\mathcal{K}_{1,2}}$ with quasimomenta (spin currents) $\mathcal{K}_1 = \frac{2\pi n_1}{N}$ and $\mathcal{K}_2 = \frac{2\pi n_2}{L-N}$. Here $n_1 = 0, \ldots, N-1$ and $n_2 = 0, \ldots, L-N-1$. Then the diagonalization of \mathcal{H}'_0 can be completed by the Jordan-Wigner (JW) transformation of the form $\tau_l^z = 1 - 2n_l$ and $\tau_l^+ = c_l \prod_{j < l} (1 - 2n_j)$, where $n_l = c_l^\dagger c_l$ and c_l annihilates a JW fermion. The last step is a Fourier transformation; we have

$$\mathcal{H}'_{0} = J \sum_{l=1}^{L-1} c_{l+1}^{\dagger} c_{l} + J e^{-2\pi i \Phi} c_{1}^{\dagger} c_{L} + \text{H.c.}$$

$$= 2J \sum_{k} c_{k}^{\dagger} c_{k} \cos k, \qquad (13)$$

where the phase $2\pi\Phi=\mathcal{K}_1+\mathcal{K}_2-\pi(L-N-1)$ is twisting the boundary condition, $c_{L+1}=e^{2\pi i\Phi}c_1$, just like a magnetic flux Φ through the periodic ring. The standard cosine dispersion of the fermions involves quasimomenta k quantized as $k=\frac{2\pi}{L}(m+\Phi)$ with m=0,...,L-1.

As shown in Ref. [35], for PBC the ground state (GS) of the model is the Fermi sea of the JW fermions at half-filling,

$$N = L/2, \tag{14}$$

with a zero spin current,

$$\mathcal{K}_1 + \mathcal{K}_2 = 0. \tag{15}$$

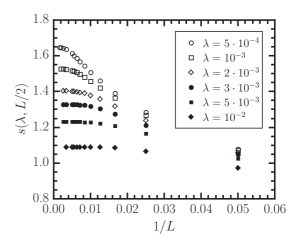


FIG. 2. Finite-size scaling results for the block entropy of an open chain as a function of 1/L (up to length L=600) for several values of λ . For each λ the entropy saturates for small enough 1/L where the thermodynamic limit is achieved.

Thus the GS wave function of \mathcal{H}'_0 factorizes into a product of an orbital and spin state:

$$|0\rangle = c_{k_1}^{\dagger} \dots c_{k_N}^{\dagger} |\text{vac}\rangle_c \otimes |\mathcal{K}_1\rangle^{(1)} \otimes |-\mathcal{K}_1\rangle^{(2)}, \qquad (16)$$

with $|\mathrm{vac}\rangle_c$ being a vacuum of the JW fermions. The orbital state is a Fermi sea with occupied quasimomenta $k_j=\frac{\pi}{N}(-j+\frac{1}{2})-\frac{\pi}{2}$ for $j=1,\ldots,N$ such that $\cos k_j<0$. The spin states $|\mathcal{K}\rangle^{(1,2)}$ are the \mathcal{K} -momentum eigenstates of the operators $R_1^{(1,2)}$ with an eigenvalue $e^{i\mathcal{K}}$. The ground state is degenerate with respect to the N values of \mathcal{K}_1 .

For OBC the spin state is completely arbitrary and the orbital state is a half-filled Fermi sea with OBC. Since in this paper we are interested in the thermodynamic limit, from now on we focus on OBC.

The Kumar model is the special case of the Heisenberg coupling $\lambda = 0$ in Eq. (3). Its 2^L -fold degenerate ground state is very sensitive to the perturbation. In the following we scan different regimes of λ .

IV. FERROMAGNETIC COUPLING: $\lambda < 0$

This case is easy to solve in the Kumar basis, where the ferromagnetic spin state is the ground state of both the Heisenberg term (8) and the Kumar model (7). The orbital ground state is a half-filled Fermi sea.

Back in the physical representation, permuted spins are still ferromagnetic, the exchange operators $X_{i,j} = 1$, and the orbitals are in the ground state of the antiferromagnetic XY chain; see Eq. (2). The ground state is a product of the ferromagnetic spin state (FM) and the orbital state—there is no spin-orbital entanglement.

V. PERTURBATIVELY WEAK COUPLING: $0 < \lambda \ll 1/L$

On the antiferromagnetic side the solution is no longer so simple. In order to get a rough idea how the highly degenerate Kumar ground state may respond to a weak Heisenberg perturbation, it is tempting to try a perturbation theory in weak λ . In the Kumar representation for OBC, the spin state is

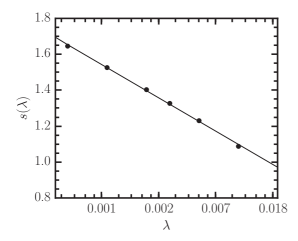


FIG. 3. The entanglement entropy of a half-chain as a function of λ in the thermodynamic limit. Note that the x scale is logarithmic. The points are the DMRG results, while the solid line is the best fit using Eq. (20).

arbitrary and the orbital state is a half-filled Fermi sea with a gap $\simeq 1/L$. When the perturbation is weaker than the orbital gap, $0 < \lambda \ll 1/L$, one can treat the spins with the degenerate perturbation theory.

To first order in λ , the orbital state does not change, protected by the orbital gap, and the spin degeneracy is removed by a Hamiltonian,

$$\mathcal{H}'_{1} = \langle \mathcal{V}' \rangle = \lambda \sum_{i,j=1}^{L} \frac{1}{2} \mathbf{J}_{i,j} \vec{\sigma}_{i} \vec{\sigma}_{j}, \tag{17}$$

where the average $\langle ... \rangle$ is taken in the orbital Fermi sea. In the physical representation only nearest-neighbor (NN) spins are coupled.

After the Kumar transformation, each physical NN coupling contributes to (is smeared over) many $J_{i,j}$ (see Appendix for more details). The transformation is a permutation that maps the N consecutive spins on empty sites (orbitals up) to spins i = N,...,1 and the N consecutive spins on occupied sites (orbitals down) to spins i = N + 1,...,2N. The exact matrix $J_{i,j}$ is described in more detail in Appendix. It turns out to be dominated by the couplings between the two N-spin subsystems, as shown in Fig. 1. The strongest bonds are along the rungs of the spin ladder. When we keep only them, then the ground state will become a product of spin singlets along the rungs. We compared the energy of this trial state to the energy from \mathcal{H}'_1 with the full matrix J. For L=16 the energies differ by less than 0.00004 in the case of $\lambda=0.01$ and by slightly more than 0.0003 for $\lambda=0.1$.

A full trial state is the spin trial state times the orbital Fermi sea. It can be transformed back to the physical representation. For L=16 the result shows strong dimerization of spins, i.e., $\langle \vec{\sigma}_l \vec{\sigma}_{l+1} \rangle < 0$ on odd NN bonds and =0 on even ones. The odd bonds take large values: $\{-2.590, -2.183, -2.063, -2.018, -2.018, -2.063, -2.183, -2.590\}$. This perturbative result uncovers a tendency of the system towards spin dimerization. The transition to the dimerized state is discontinuous (first order) by the very nature of the degenerate perturbation theory.

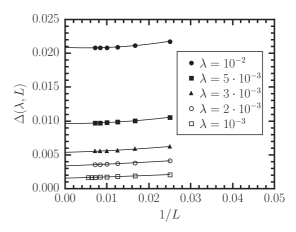


FIG. 4. The finite-size scaling of the spin gap for several values of λ . The solid lines denote the result of our fit using Eq. (21).

However, the degenerate perturbative treatment is not justified when λ becomes comparable or stronger than the orbital gap $\simeq 1/L$, hence it must fail, at least quantitatively, in the thermodynamic limit. This is why in the following we use DMRG for OBC to obtain numerically exact results in this limit.

VI. CROSSOVER TO THE THERMODYNAMIC LIMIT: $\lambda \simeq 1/L$

In our DMRG calculation we used the dynamic block-state selection algorithm [44,45]. The *a priori* value of the quantum information loss was set to $\chi=10^{-7}$, which required to keep block states up to 4000. Typical truncation errors were in order of 10^{-8} . We considered finite chains with OBC up to length L=800.

The thermodynamic limit is reached when the system size is longer than its correlation length ξ . The crossover to this regime can be seen, e.g., in the entanglement (von Neumann) entropy, $s(\lambda, L/2)$, between the two halves of the chain [46–48]. This quantity can be obtained from the appropriate reduced density matrix, $\rho_{L/2}(\lambda)$:

$$s(\lambda, L/2) = -\text{Tr}\rho_{L/2}(\lambda)\ln\rho_{L/2}(\lambda), \tag{18}$$

where $\rho_{L/2}(\lambda)$ denotes the reduced density matrix of the subsystem containing sites 1,...,L/2. In Fig. 2, we show the entropy for different system sizes and small $\lambda \ll 1$.

For small system sizes, or large 1/L, the entropies for different λ collapse. This is the regime of validity of the degenerate perturbation theory, where the actual strength of a weak $\lambda \ll 1/L$ is irrelevant.

For each λ the entropy is growing with the system size L until it saturates below a crossover value of 1/L. This saturation is characteristic for the thermodynamic limit. Interestingly, the weaker λ is the longer L is needed for the saturation, hence the correlation length ξ must increase or even diverge when $\lambda \to 0$. In the following section, we consider mainly "saturated" results in the thermodynamic limit for all physical observables.

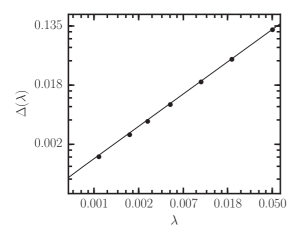


FIG. 5. The spin gap extrapolated to the thermodynamic limit as a function of λ . Note that both scales are logarithmic. The points are the DMRG results, while the solid line is the best fit using Eq. (22).

VII. THERMODYNAMIC LIMIT: SECOND-ORDER TRANSITION FOR $\lambda \rightarrow 0^+$

First, we consider the behavior of the block entropy of a half-chain $s(\lambda)$ in the thermodynamic limit:

$$s(\lambda) = \lim_{L \to \infty} s(\lambda, L/2). \tag{19}$$

This is shown in Fig. 3, where the x scale is logarithmic.

The entropy diverges as we approach the critical point. Near the critical point at $\lambda \to 0^+$ the entropy is expected to behave as [48]

$$s(\lambda) = \frac{c}{6} \ln(\xi) = -\frac{c\nu}{6} \ln(\lambda/\lambda_0), \tag{20}$$

where c is the central charge and $\xi = (\lambda/\lambda_0)^{-\nu}$ the correlation length with the exponent ν . The best fit is $\lambda_0 = 3.9 \pm 0.5$ and $c\nu = 1.11 \pm 0.02$.

The diverging block entropy indicates a diverging correlation length ξ in the system and a vanishing gap $\Delta = \Delta_0 \lambda^{z\nu}$, where z is the dynamical exponent. Therefore we investigated the first few low-lying excitations in the model. We found that the first excitation for $\lambda \ll 1$ is a spin flip, that is, the smallest

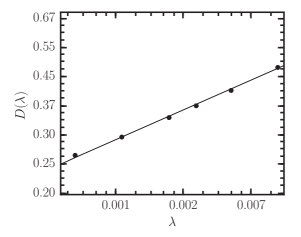


FIG. 6. The extrapolated dimerization order parameter D as a function of λ . The points are the DMRG results, while the solid line denotes the fit using Eq. (24). Note that both scales are logarithmic.

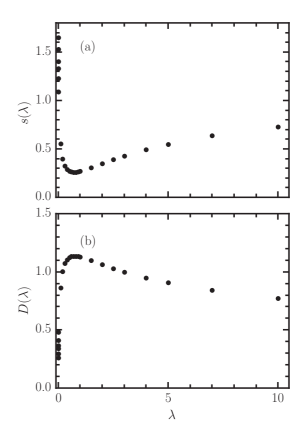


FIG. 7. The extrapolated block entropy $s(\lambda)$, and dimerization order parameter $D(\lambda)$, as a function of λ are shown in (a) and (b), respectively.

gap is given by the spin gap. To obtain the bulk values of the gap, $\Delta(\lambda)$, we performed extrapolation to the thermodynamic limit with the fitting ansatz,

$$\Delta(\lambda, L) = \Delta(\lambda) + A/L + B/L^2, \tag{21}$$

where $\Delta(\lambda)$, A, and B are free parameters. This ansatz is motivated by the fact that the finite-size corrections are expected to be algebraic for OBC. The finite-size scaling procedure is demonstrated in Fig. 4.

The extrapolated gap as a function of λ is shown in Fig. 5 with a log-log scale.

The data points can be fitted well with a linear, that is, the gap exhibits a power-law dependence on λ :

$$\Delta(\lambda) = \Delta_0 \lambda^{z\nu},\tag{22}$$

where $\Delta_0 = 3.12 \pm 0.02$ and $z\nu = 1.088 \pm 0.002$. This means that the gap vanishes continuously in contrast to the naive prediction of the perturbation calculation, and a second-order phase transition occurs at $\lambda = 0^+$.

The same applies for the dimerization D, which is defined as

$$D(\lambda) = \lim_{L \to \infty} |s(\lambda, L/2) - s(\lambda, L/2 - 1)|. \tag{23}$$

This is shown in Fig. 6 with a log-log scale. The dimerization—similarly to the gap—behaves as

$$D(\lambda) = D_0 \lambda^{\beta}, \tag{24}$$

where $D_0 = 1.22 \pm 0.03$ and $\beta = 0.204 \pm 0.004$.

VIII. DIMERIZATION FOR FINITE λ BOTH LARGE AND SMALL

The previous section focused on the critical behavior in the limit $\lambda \to 0^+$. This is a second-order phase transition, where the symmetry broken by dimerization is restored. In this section we paint a general picture of the dimerized phase at $\lambda > 0$. We distinguish two regimes, of large and small λ , where the nature of the dimerization is qualitatively different. They are separated by a smooth crossover.

It is interesting to analyze the behavior of the block entropy and the dimerization order parameter for larger λ before going into the details of the dimerized phase. They are shown in Fig. 7. It is seen that around $\lambda \simeq 0.7$ the block entropy and the dimerization order parameter exhibits a local minimum and maximum, respectively. This can be understood as follows: When the dimerization is the strongest the ground state is the less entangled, which manifests in the minimum value of the block entropy. We address what happens for $\lambda \to \infty$ later in this section.

A. Numerical findings

Numerical observations are as follows. For any λ we observe both spin and orbital dimerization in the sense that the NN correlations alter between large (small) values for odd (even) bonds as shown in Fig. 8. The difference between the spin and orbital sectors is that the spin correlations remain always AF while the orbital ones can alternate between positive

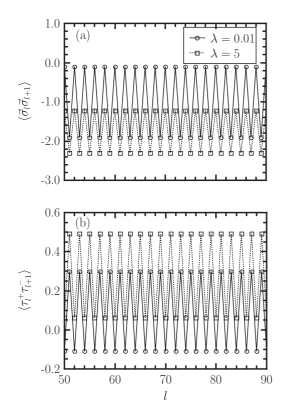


FIG. 8. DMRG results for the NN spin (a) and orbital (b) correlations for the open chain of L=140 sites as a function of the site number l for $\lambda=0.01$ and $\lambda=5$.

TABLE I. The correlation lengths of the spin and orbital correlation functions for several values of λ .

λ	ξorbital	ξ _{spin}
0.01	5.52	4.65
5	4.72	18.18

and negative values for small λ and remain FM only for large λ . The behavior of the long-range spin and orbital correlations is different in the small and large λ regime. While the correlation $\langle \vec{\sigma}_l \vec{\sigma}_{l+R} \rangle$ always decays exponentially with R it remains constantly negative for small λ while for large λ it alternates between negative and positive values. Thus we say that for small λ we have strong dimerization of the spins. Because of the negative sign of the correlations this is a spin-singlet dimerization. The behavior of the orbital correlations seems to be complementary, the long-range correlations $\langle \tau_l^+ \tau_{l+R}^- \rangle$ are constantly positive for large λ , while for small λ they alternate with R. In both regimes they decay exponentially with R. Thus, analogically to the spin sector, we say that for large λ the orbitals exhibit strong dimerization in the orbital-triplet state. Summarizing, we have found strong spin-singlet dimerization for small λ and strong orbital-triplet dimerization for large λ .

Finally, we note that the transition between the two regimes, of strong dimerization of either spins or orbitals, is a crossover without any features of a phase transition. However, one can easily observe a change in the behavior of the entanglement entropy of the half-system $s(\lambda)$ in Fig. 7(a); while it is rapidly decreasing for small λ , it becomes slowly increasing for larger values of λ with a shallow minimum at $\lambda \simeq 0.7$, where the dimerization is the strongest.

At the same time, in addition to the qualitative change of the spin correlation triggered by increasing λ , there is a big difference in their correlation lengths. They are shown in Table I. From the semilogarithmic plot of Fig. 9 we can learn that for small λ both correlations decay with almost the same exponent, while for large λ the spin correlation function has much larger correlation length than that of the orbital

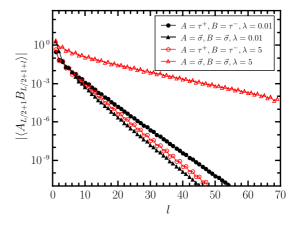


FIG. 9. (Color online) DMRG results for the spin and orbital correlations for the open chain of L=140 sites as a function of the site number l for $\lambda=0.01$ and $\lambda=5$. The vertical axis has a logarithmic scale.

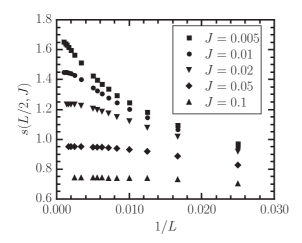


FIG. 10. Finite-size scaling results for the block entropy of an open chain as a function of 1/L (up to L=800) for several decreasing values of J and $\lambda=1$. For each J the entropy saturates for small enough 1/L where the thermodynamic limit is achieved.

correlation function. This suggests that there is a second phase transition in the system located at $\lambda \to \infty$ or J=0. This is not totally unexpected as in this limit the KH model becomes simply the Heisenberg model in the spin sector which is known to be critical. This observation is confirmed by the behavior of the block entropy of the half system, s(L/2,J), which exhibits similar scaling properties as $J\to 0$ for $\lambda=1$ (see Fig. 10), to what we have seen for $\lambda\to 0$ in Eq. (20). In the present case we fit the product $cv=1.85\pm0.01$. Since c=1 for the Heisenberg chain, we obtain an estimate for the correlation length exponent $v=1.85\pm0.01$.

B. Qualitative explanation

The dimerization of spins involving constantly negative long-range correlation function can be understood in terms of the first-order perturbative expansion of Sec. V and especially looking at Fig. 1. As we can see, in the Kumar basis the singlet bonds are mainly on the ladder's rungs connecting spins at sites N+i and N-i+1. In the physical basis the rung singlets become, in first approximation, spin singlets on every second bond. In this approximation the correlation function is strongly negative only within a given singlet and zero anywhere else. However, when we take into account fluctuating positions of the up- and down-orbitals in the orbital Fermi sea, then the singlets on every second bond become smeared over neighboring sites. This smearing implies that (i) singlets placed every second bond are not perfect, i.e., $0 > \langle \vec{\sigma}_l \vec{\sigma}_{l+1} \rangle > -3$ and (ii) there are negative spin correlations between sites land $m \in [l - \delta l, l + \delta l]$ where δl is the length scale at which a single up-orbital is delocalized within the Fermi sea. In this framework the strongly spin-dimerized ground state of the KH model can be understood as a spin liquid of strongly resonating

The triplet dimerization of the orbitals for large λ can be easily understood in the mean-field way similarly as it was done in Ref. [2]. In this limit the leading part of the Hamiltonian is a pure Heisenberg term $\mathcal V$ for the spins. Thus we can use a

mean-field decoupling of a spin-orbital term in \mathcal{H}_0 :

$$X_{l,l+1}(\tau_{l}^{+}\tau_{l+1}^{-} + \tau_{l}^{-}\tau_{l+1}^{+})$$

$$\approx \langle X_{l,l+1}\rangle(\tau_{l}^{+}\tau_{l+1}^{-} + \tau_{l}^{-}\tau_{l+1}^{+}) + X_{l,l+1}\langle \tau_{l}^{+}\tau_{l+1}^{-} + \tau_{l}^{-}\tau_{l+1}^{+}\rangle$$

$$-\langle X_{l,l+1}\rangle\langle \tau_{l}^{+}\tau_{l+1}^{-} + \tau_{l}^{-}\tau_{l+1}^{+}\rangle. \tag{25}$$

The spins are AF with $\langle \vec{\sigma}_l \vec{\sigma}_{l+1} \rangle < -1$, hence for each bond we have $\langle X_{l,l+1} \rangle < 0$ and, according to the first line of Eq. (25), the orbitals order ferromagnetically. The instability towards dimerization, meaning here formation of the orbital triplets, can be demonstrated by imagining a self-consistency loop for a nonuniform MF Hamiltonian. If, for any reason, a bond has increased the orbital correlation, then, according to the second line of Eq. (25), the spins will feel an increased tendency towards antiferromagnetism. Then again, the first line of Eq. (25) implies that the orbital bonds will be increased and so the spin bond will follow. Such a loop clearly leads to maximalization of the orbital correlations at some bonds and, since we cannot have a triplet at every bond, we must have the dimerization of the orbitals. We verified this argument by solving numerically a nonuniform MF Hamiltonian for L=16.

IX. CONCLUSION

We have found that an admixture of the Heisenberg interactions λ between the spins causes a spontaneous dimerization of the Kumar model. The dimerization is both in the spin and orbital sector but it becomes strong for the spins in the regime of small λ and for the orbitals in the opposite limit. We have shown that the dimerization of spins can be understood by means of the Kumar transformation, i.e., in a basis where the spins and orbitals in the Kumar Hamiltonian get decoupled. On the other hand we show that the dimerization of orbitals for large λ can be understood be a mean-field mechanism in the physical basis.

We have seen that the perturbative expansion in the Kumar basis, however useful for understanding the spin dimerization, fails to capture the second-order phase transition found by DMRG in the thermodynamic limit. Whereas the perturbative prediction is always a discontinuous transition, the numerical study of the system of sizes up to L = 600 allowed us to observe the divergence of the block entropy, vanishing of the gap and the dimerization order parameter in the regime where we approach the transition point at $\lambda = 0$ slower than the gap in the pure Kumar model closes, i.e., in the regime where the perturbative approach must fail. We note that the orbital gap in the pure Kumar model will open again on adding a small admixture of $J^z \tau_{l+1}^z \tau_{l+1}^z$ term to \mathcal{H}_0 . Such a term is usually present in the spin-orbital Hamiltonians for the t_{2g} systems, for instance, vanadates [5]. Then the perturbation approach will be valid in the thermodynamic limit as well and the smooth transition at $\lambda = 0$ will be lost; however, since τ_i^z operators transform trivially under the Kumar transformation, the dimerization at finite λ will remain. Finally, by observing that the spin correlation length is strongly increased as λ grows we have deduced that there is another phase transition in the limit of $\lambda \to \infty$ (or equivalently J=0). This we have confirmed by the finite-size scaling of the block entropy for decreasing J. The phase diagram is shown schematically in Fig. 11.

$$\begin{array}{c|c}
\text{Kumar} \\
\hline
\text{FM} & \text{dimerized} & \text{Heisenberg} \\
\hline
\lambda = 0 & \lambda = \infty
\end{array}$$

FIG. 11. Schematic phase diagram of the Kumar-Heisenberg model for J=1.

Finally we note that the full dimerization in orbital space can be easily understood by the mean-field approach, similarly as it was done for YVO₃ [2] at finite temperature, but here in zero temperature, the spin dimerization is more complex and its mechanism can be revealed only by the Kumar transformation. This is a feature that was not found in the similar spin-orbital models, namely the SU(2) \otimes XY [31] and SU(2) \otimes SU(2) [31,49] ones, where in principle the dimerization could be captured by a simple variational wave functions in the physical basis and confirmed via DMRG. We argue that such a dimerization can be observed in the (quasi) one-dimensional compounds with active spin and orbital degrees of freedom.

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APPENDIX: THE EFFECTIVE SPIN COUPLINGS

The effective, first-order couplings between the spins can be expressed as

$$J_{i,j} = \sum_{p_1, \dots, p_N} P(\vec{p}) \sum_{l=1}^N \delta_{i,f(l,\vec{p})} \delta_{j,f(l+1,\vec{p})},$$
(A1)

where the first sum is an average over a probability distribution,

$$P(\vec{p}) = \prod_{1 \le i < j \le N} \left\{ 4 \sin^2 \frac{\pi(p_j - p_i)}{L} \right\}, \quad (A2)$$

for empty sites $\{p_1,...,p_N\}$ in a half-filled Fermi sea. Here we consider periodic boundary conditions to minimize boundary effects. For N=L/2 the permutation maps the N consecutive spins on empty sites (orbitals up) to spins N,...,1 and the N consecutive spins on occupied sites (orbitals down) to spins N+1,...,2N. In the physical representation only nearest-neighbor spins are coupled. After the transformation \mathcal{U} each physical NN coupling (PNNC) contributes to (is smeared over) many $J_{i,j}$.

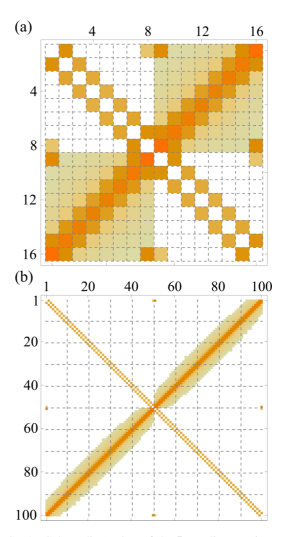


FIG. 12. (Color online) Plots of the **J** coupling matrices of the effective spin Hamiltonians $\tilde{\mathcal{H}}_1$ for system sizes: (a) L=16 and (b) L=100. Colors tending to red mean high positive values; white fields mean zero values. Dashed grid is a guide for the eye.

The matrix $J_{i,j}$ is shown in Figs. 12(a) and 12(b) as color plots for a small and large L, respectively. The matrix has two diagonal and two off-diagonal $N \times N$ blocks. The diagonal block i, j = 1, ..., N originates from PNNC's between spins on empty sites. Since only a pair of consecutive empty sites can happen to be NN's, the block couples only NN's. The weakest coupling $J_{1,N}$ originates from the relatively rare situation when the first and last empty sites also happen to be, respectively, the first and last sites of the chain, i.e., they are not only NN's (across the closing bond) but also each of them occupies a definite position. $J_{1,N} = 0$ for an open chain. The diagonal block i, j = N + 1, ..., 2N originates from PNNC's between spins on occupied sites. In our half-filled orbital Fermi sea the two diagonal blocks are the same.

The two identical off-diagonal blocks originate from PNNC's between pairs of empty and occupied sites. The top values of the coupling are between the first and the last site: $J_{1,L} \simeq 0.705$ for L=16 and $J_{1,L} \simeq 0.783$ for L=100. This strong coupling originates from the PNNC between the Nth empty and the Nth occupied sites that are very likely to be next to each other. The coupling $J_{1,N+1}$ originates from the

PNNC between the Nth empty and the first occupied site that is possible mainly across the closing bond. This coupling is much weaker for an open chain where the Nth empty and first occupied sites are rather unlikely to be NN. However, the dominant feature of each off-diagonal block is its antidiagonal belt. A coupling $J_{N-i,j}$ within this belt originates from a PNNC between the ith empty and jth occupied site with i close to

j. These sites are most likely to be NN when their difference |i-j| is not much greater than a variation of a position of the ith empty (or the jth occupied) site in the Fermi sea. This variation limits the width of the antidiagonal belt.

We found that truncating the full matrix $J_{i,j}$ to its dominant antidiagonal belt is a very good approximation. We are assuming this approximation in the main text.

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