Cross-over from an incommensurate singlet spiral state with a vanishingly small spin-gap to a valence bond solid state in dimerized frustrated ferromagnetic spin-chains

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Motivated by the magnetic properties of the spin-chain compounds LiCuSbO4≡LiSbCuO4 and Rb2Cu2Mo3O12, we study the ground state of the Heisenberg chain with dimerized nearest-neighbor ferromagnetic (FM) (J1, J′ 1 < 0) and next-nearest-neighbor antiferromagnetic (J2 > 0) couplings. Using the density-matrix renormalization group technique and spin-wave theory we find a first-order transition between a fully-polarized FM and an incommensurate spiral state at 2α = β/(1 + β), where α is the frustration ratio J2/J1 and β the degree of dimerization J′ 1/J1. In the singlet spiral state the spin-gap is vanishingly small in the vicinity of the FM transition, corresponding to a situation of LiCuSbO4. For larger α, corresponding to Rb2Cu2Mo3O12, and smaller β there is a crossover from this frustration induced incommensurate state to an Affleck-Lieb-Kennedy-Tasaki-type valence bond solid state with substantial spin-gaps.

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Introduction.— The exotic phenomena emerged by magnetic frustration have long been fascinating subjects of research in condensed matter physics [1]. Nowadays, quasi one-dimensional (1D) frustrated systems, despite their simple structure, are at the center of attention as a playground for novel ground states that can emerge from frustration and strong quantum fluctuations due to low dimensionality. So far, various unconventional magnetic states such as quantum spin liquids [2–4], spin-Peierls states [4], and Tomonaga-Luttinger (TL) liquid phases [5] have been investigated. Currently, among the hottest topics are magnetic multipoles and in particular spin-nematic states [6–11] in which magnon bound states are formed from a subtle competition between geometrical balance of ferromagnetic (FM) and antiferromagnetic (AFM) correlations among spins.

Very recently, a magnetic field-induced “hidden” spin-nematic state was reported in the anisotropic frustrated spin-chain cuprate LiCuSbO4 [12]. By the nuclear magnetic resonance technique, a field-induced spin gap was observed above a field ≈ 13T in the measurements of the 7Li spin relaxation rate T1 −1, supported by static magnetization and electron spin resonance data. This material has a unique crystal structure: In the CuO2 chain, four nonequivalent O2− ions within a CuO4-plaquette give rise to two kinds of nonequivalent left and right Cu-Cu bonds along the chain direction. This gives rise to alternating nearest-neighbor transfer integrals (t1 ≠ t′ 1). As a result, a sizeable splitting of the two nearest-neighbor FM exchange integrals was estimated: J1 ≈ −160K and J′ 1 ≈ −90K, whereas the next-nearest-neighbor AFM coupling is J2 ≈ 37.6K [see Figure 1(a)]. Another example of a FM dimerized chain compound is Rb2Cu2Mo3O12 which has CuO2 ribbon chains. Here its ribbon chains are twisted, so that the Cu-Cu distances and the Cu-O-Cu angles are slightly alternating. Accordingly, a small dimerization of the nearest-neighbor exchange integrals is expected. Assuming no dimerization, the values of the FM nearest- and AFM next-nearest-neighbor exchanges have been estimated as -138K and 51K, respectively, by the fitting of susceptibility and magnetization [13]. Besides, a non-magnetic ground state with energy gap E0 ≈ 1.6K has been experimentally detected [4]. So far, the 1D dimerized AFM Heisenberg has been extensively studied in connection to the celebrated spin-Peierls compound CuGeO3 [15]. In contrast, the dimerized FM case has been hardly ever discussed. Recently, only the weakly dimerized case has been investigated [16], and theoretical studies are definitely required.

Motivated by the above observations, we study a dimerized FM Heisenberg chain with next-nearest-neighbor AFM couplings by using the spin-wave theory (SWT) and the density-matrix renormalization group (DMRG) method. The ground-state phase diagram is obtained as a function of dimerization and frustration strengths, based on the numerical results of total spin, spin gap, spin-spin correlation function, and Tomonaga-Luttinger (TL) liquid exponent. We establish the presence of a frustration induced incommensurate singlet state with a spin-gap that is vanishingly small close to the vicinity to a first order FM transition, corresponding to the situation of LiCuSbO4. Despite the vanishingly small gap, correlation lengths are comparable to those in the large-gap region in the phase diagram. For larger α and smaller β there is a crossover from this frustration induced incommensurate state to an Affleck-Lieb-Kennedy-Tasaki-type valence bond solid (realized at β = 0) with substantial spin-gaps. We also confirm the presence of finite spin gap in the uniform J1 = J2 limit.

Model and method.— Our spin Hamiltonian is given by

\[ H = \sum_{i=\text{even}} J_1 S_i \cdot S_{i+1} + \sum_{i=\text{odd}} J'_1 S_i \cdot S_{i+1} + J_2 \sum_i S_i \cdot S_{i+2} \]

(1)

where \( S_i \) is a spin-1/2 operator at site \( i \). The nearest-neighbor (\( J_1, J'_1 < 0 \)) and next-nearest-neighbor (\( J_2 > 0 \)) interactions are FM and AFM, respectively [see Figure 1(a)], and we use the notations of next-nearest-neighbor coupling ratio \( \alpha = J_2/J_1 \) and nearest neighbor coupling ratio \( \beta = J'_1/J_1 \).
hereafter.

When the system is undimerized \((\beta = 1)\), we are dealing with the so-called \(J_1 - J_2\) model. Increasing \(\alpha\), a phase with incommensurate spin-spin correlations follows a FM phase. The transition occurs at \(\alpha = 1/4\), both in the quantum as well as in the classical model \([17,18]\). The incommensurate (“spiral”) correlations are short ranged in the quantum model \([19,20]\). A vanishingly small gap was predicted by the field-theory analysis \([21]\) but no numerical evidence exists so far. In the limit of \(\beta = 0\), the system \([11]\) is equivalent to spin ladder with AFM legs and FM rung couplings. Since this system can be effectively reduced to an \(S = 1\) AFM Heisenberg chain with regarding two \(S = 1/2\) spins on each rung as a \(S = 1\) spin \([23,24]\), the ground state is gapped as predicted by Haldane conjecture \([22]\). Therefore, the ground state can be well described by a valence-bond-solid (VBS) picture, proposed in the Affleck-Lieb-Kennedy-Tasaki (AKLT) model \([25]\). The schematic picture is shown in Figure 1(b).

The DMRG method \([26]\) is employed to investigate the ground-state properties of the system \([11]\). We calculate the total spin with periodic boundary conditions, and spin gap, spin-spin correlation functions, Tomonaga-Luttinger (TL) spin exponent with open boundary conditions. We keep up to \(m = 6000\) density-matrix eigenstates in the renormalization procedure and extrapolate the calculated quantities to the limit \(m \rightarrow \infty\) if necessary. Furthermore, several chains with lengths up to \(L = 800\) are studied to handle the finite-size effects. In this way, we can obtain quite accurate ground states within the error of \(\Delta E/L = 10^{-9} - 10^{-10}|J_1|\).

**Ferromagnetic critical point.**—In the limit of \(\beta = 0\) and \(\alpha = 0\), the FM critical point no longer exists because the system is solely composed of isolated spin-triplet dimers. However, if \(\beta\) is finite, the FM order is expected for small \(\alpha\). Let us then consider the \(\beta\)-dependence of the critical point. Since the quantum fluctuations vanish at the FM critical point, the classical SWT may work perfectly for estimating the FM critical point. By the SWT the excitation energy for a FM ground state is given as \(2\omega_q = -\sqrt{1 + \beta^2 + 2\beta \cos(2q) + 2\alpha \cos(2q)}\). The system is in the FM ground state if \(\omega_q > 0\) for all \(q\); otherwise, it is in the spiral singlet state. Thus, the FM critical point is derived as

\[
\alpha_c,1 = \frac{\beta}{2(1 + \beta)}.
\]  

As shown in Figure 2, the FM region is simply shrunk with decreasing \(\beta\), and disappears in the limit of \(\beta = 0\) as a consequence of isolated FM dimers. It can be numerically confirmed by calculating the ground-state expectation value of the total-spin quantum number \(S\) of the whole system, \(S^2\), defined as \(S^2 = S(S + 1) = \sum_{ij} \langle S_i \cdot S_j \rangle\). In Figure 3 the normalized total spin at \(\beta = 0.5\) is plotted as a function of \(\alpha\). We can find a direct jump from \(S = 0\) to \(S = L/2\) at \(\alpha \sim 0.17\), indicating the absence of an intermediate (partially polarized) FM state. This critical value is in good agreement with that obtained by the SWT (\(\alpha_c,1 = 1/6\)). Similarly, for all \(\beta\)-values, we confirm direct transition between FM \((S = L/2)\) and singlet spiral \((S = 0)\) states as well as perfect agreement between SWT and DMRG critical points as compared in Figure 2.

**Haldane gapped state.**—So far, the spin gapped state has been verified in the limit of \(\beta = 0\) \([23,24]\). This can be interpreted as a realization of the AKLT VBS state. However, it is a nontrivial question what happens to the spin gap for finite \(\beta\). In our DMRG calculations, the spin gap \(\Delta\) is evaluated as the energy difference between the lowest triplet state and the singlet ground state,

\[
\Delta(L) = E_0(L, S^z = 1) - E_0(L, S^z = 0), \quad \Delta = \lim_{L \rightarrow \infty} \Delta(L),
\]

where \(E_0(L)\) is the ground-state energy for a given number of system length \(L\) and \(z\)-component of total spin \(S^z\).

First, we focus on the case of \(\beta = 0\), namely, a ladder consisting of two AFM leg chains and FM rungs. In Figure 4(a) the extrapolated values of \(\Delta/|J_1|\) is plotted as a function of
where system is exactly reduced to a S∆Log-log plot of βα is consistent with the prediction by the bozonization method [24]∆α. The gap opens at α = 0.6 and saturates at a certain value scaled by Hαeff = 0 and increases monotonously with α = 0.9. The Haldane gap of the system (4) has been calculated as ∆/J2 = 0.2045 in the limit α = 0. The Haldane gap of the system [4] has been calculated as ∆/Jeff = 0.410479 [28]. Thus, we can confirm Jeff = J2/2 numerically for the mapping from Eq.(1) to Eq.(4) at the limit |J1|/J2(= 1/α) → 0 and β = 0.

Next, we look at the effect of β on the spin gap. Figure 4(b) shows a log-log plot of ∆/|J1| as a function of 1 − β for α = 0.6 and 0.9. The behaviors are nontrivial but ∆ decays roughly in power law with decreasing 1 − β. As a result, the gap is vanishingly small near the uniform J1 − J2 limit (β ≈ 1). Besides, it is interesting that ∆ for α = 0.6 is larger than that for α = 0.9 at larger β and opposite at smaller β, which may suggest that the gapped state near β = 1 is no longer the AKLT-type VBS state but the frustration-induced one (see below). This is consistent with a maximum gap around α = 0.6 at weak dimerization (β = 0.9). On the other hand, an adiabatic connection of the AKLT-type VBS state from β = 0 to 1 was predicted by the field-theoretical analysis for |J1| ≪ J2 [29]. A contour plot of the magnitude of ∆ is given in Figure 2. We can see a rapid decay of ∆ with approaching the FM phase. However, ∆ is too small to figure out whether it remains finite, e.g. ∆ ≈ 10−3, in the vicinity of the FM critical boundary. Therefore, to verify the presence or absence of the gap, we checked the asymptotic behavior of spin-spin correlation function ⟨|S_i |S_j |⟩. In Figure 5(a) the semi-log plot of ⟨|S_i |S_j |⟩ as a function of distance |i − j| is shown for some parameters near the the FM critical boundary. The distances |i − j| are taken about the mid-point of the systems to exclude the Friedel oscillations from the system edges, i.e. (i + j)/2 locates around the mid-point of the systems. All of them exhibit exponential decay of ⟨|S_i |S_j |⟩ with distance, which clearly indicates the presence of a finite spin-gap. The curves are well-fitted with the expression ⟨|S_i |S_j |⟩ ∝ cos[Q(i − j)]|i − j|−ξ exp[−ξ |i − j|] for long distances [30, 31]: the correlation lengths ξ are estimated as ξ = 11.6 (α = 0.1, β = 0.12), ξ = 8.6 (α = 0.2, β = 0.3), and ξ = 7.3 (α = 0.3, β = 0.7). In the AFM J1 − J2 model [30], a region with ξ ≈ 10 still have a spin gap of order of 10−1J1. This may imply the spin velocity of our system is more than two digits smaller than that of the AFM J1 − J2 model since ∆ = v_s/ξ where v_s is the spin velocity.

Uniform J1 − J2 model.— In the uniform case (β = 1) the existence of a tiny gap for α ≳ 3.3 was predicted by the field-theory analysis [21]. However, the investigation for smaller α is lacking. Therefore, to verify the presence or absence of a gap at smaller α, we investigated the TL liquid spin exponent Kσ. For our system having four Fermi points (±k_F1, ±k_F2), we here assume the asymptotic behavior of the spin-spin correlation function to be a power-law decay, like

\[ \langle S_i^z S_j^z \rangle \sim -\frac{K_σ}{2\pi^2 r^2} + \frac{A \cos[2(k_{F1} - k_{F2})r]}{r^2K_σ} + \cdots, \]

in analogy with the case of two coupled chains [32], because the low-energy excitation spectra are similar to those of our

\[
\begin{align*}
S_{i+1} \quad \text{and} \quad J_{\text{eff}} = J_2/2. \quad \text{In the inset of Figure 4(a) } \Delta \text{ is replotted in unit of } \alpha. \quad \text{We obtain } \Delta/J_2 = 0.2045 \text{ in the limit } \alpha = 0. \quad \text{The Haldane gap of the system [4] has been calculated as } \Delta/J_{\text{eff}} = 0.410479 [28]. \quad \text{Thus, we can confirm } J_{\text{eff}} = J_2/2 \text{ numerically for the mapping from Eq.(1) to Eq.(4) at the limit } |J_1|/J_2(= 1/\alpha) \rightarrow 0 \text{ and } \beta = 0.
\end{align*}
\]
The fastest convergence to the thermodynamic limit suggests a spin-gapless state with the power-law (\(K = 1\) within the TL liquid theory). In figure 5(b), \(K_\sigma\) is plotted as a function of \(\alpha\). We found that the inverse correlation length is well fitted by \(1/\xi = 0.085 \exp(-\pi \alpha)\) for large \(\alpha\). Since \(\Delta = v_0/\xi\), it may be feasible to speculate that the gap has a maximum around \(\alpha = 0.5-0.6\), decreases with increasing \(\alpha\), and smoothly connects to the tiny gap region.

Finally, let us explicitly address the relevance of the calculations above for the two spin-chain materials mentioned in the introduction. For LiCuSbO\(_4\), \(\alpha = 0.235\) and \(\beta = 0.56\) are estimated from the density-functional calculations: \(J_1 \approx -160\) K, \(J_2 \approx -90\) K, and \(J_3 \approx 37.6\) K \([12,38]\). The system is in the gapped spiral state, but very close to the FM phase where the spin-gap is vanishingly small. Thus, the spin gap may be too small to be detected experimentally. The second compound is Rb\(_2\)Cu\(_2\)Mo\(_3\)O\(_{12}\). If we use the previously estimated parameters \(J_1 = -138\) K and \(J_2 = 51\) K (\(\alpha = 0.37\)), a substantial dimerization (\(\beta = 0.65\)) of \(J_1\) and \(J_2\) is necessary to reproduce the experimentally observed gap \(E_g \approx 1.6\) K, namely, \(J_1 = -138\) K and \(J_2 = 90\) K. Furthermore, if it is more appropriate to consider the value \(-138\) K as an averaged FM coupling \((J_1 + J_2)/2\), then an even larger dimerization would be needed. In practice, the actual \(J_1\) should be somewhat smaller or \(J_2\) should be larger. A detailed analysis of the experimental data that explicitly takes into account the dimerization can clarify this point. In the context of these two compounds and also in general the influence of an external magnetic field is of considerable interest and will be addressed elsewhere.

**Conclusion.**—We considered a frustrated \(J_1-J_2\) spin chain with/without dimerization of nearest-neighbor FM coupling
and determined its phase diagram. The FM critical point was analytically determined to be \( \alpha_c = (\beta/2)/(1 + \beta) \) by applying the linear spin-wave theory, which was confirmed by the numerical calculation of the total spin. The transition between the fully polarized FM and the singlet spiral states is of the first order and no partially polarized FM state exists. The spin-gap in the vicinity of the FM boundary was confirmed to be finite by the exponential decay of the spin-spin correlation functions but it is vanishingly small. In the uniform \( J_1 - J_2 \) chain, the gapped state appears at least around \( J \approx 0.5 - 0.6 \) where the TL liquid exponent \( K_\sigma \) goes to 0 in the thermodynamic limit. Near \( \beta = 0 \) the spin-gap increases with increasing \( \alpha \); whereas, near \( \beta = 1 \) it has a maximum value around the strongest frustration region \( \alpha = 0.5 - 0.6 \). Therefore, the gap opening in the entire incommensurate singlet phase may be interpreted as a crossover from the AKLT-type valence bond solid state near \( \beta = 0 \) to the frustration-induced dimerized state near \( \beta = 1 \).

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[38] Thereby fixed FM contributions to the total \( J_1 \) and \( J'_1 \)-values have been assumed, i.e. taking into account the modification of the AFM superexchange contributions to the former, only. An LDA+U analysis available in future might provide somewhat refined numbers. Within a \( pd \)-multiband Hubbard model this corresponds to fixed direct ferromagnetic exchange interactions \( K_{pd} \) and ferromagnetic Goodenough-Kanamori exchange interactions involving the Hund’s rule coupling involving the intermediate oxygen sites generic for edge-shared \( CuO_2 \) chain compounds. The latter might be checked by advanced quantum chemistry calculations for small chain clusters. In the absence of results such calculations at present, the model considered here provides a convenient tool to investigate phenomenologically \( LiSbCuO_4 \) and other related compounds.