Entanglement Perturbation Theory for Infinite Quasi-1D Quantum Systems

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We develop Entanglement Perturbation Theory (EPT) for infinite Quasi-1D quantum systems. The spin 1/2 Heisenberg chain with ferromagnetic nearest neighbor (NN) and antiferromagnetic next nearest neighbor (NNN) interactions studied as a prototypical system. The obtained accurate phase diagram is compared with a recent prediction [Phys.Rev.B, 81,094430(2010)] that dimer and Néel orders appear alternately as the XXZ anisotropy $\Delta$ approaches the isotropic limit $\Delta = 1$. The first and second transitions (across dimer, Néel, and dimer phases) are detected with improved accuracy at $\Delta \approx 0.722$ and 0.930. The third transition (from dimer to Néel phases), previously predicted to be at $\Delta \approx 0.98$, is not detected at this $\Delta$ in our method, raising the possibility that the second Néel phase is absent.

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In a recent article, Furukawa et al.\textsuperscript{[1]} reported exotic Néel and dimer orders in a spin 1/2 Heisenberg chain described by the Hamiltonian,

$$
H = \sum_{\delta=1}^{2} \sum_{i=1}^{L} J_{\delta} \left( S_{i}^{x} S_{i+\delta}^{x} + S_{i}^{y} S_{i+\delta}^{y} + \Delta S_{i}^{z} S_{i+\delta}^{z} \right)
$$

characterized by a ferromagnetic NN exchange coupling $J_{1} < 0$ and an antiferromagnetic NNN exchange coupling $J_{2} > 0$, with an easy-plane anisotropy $\Delta < 1$. Let us describe the effect of the NNN exchange by $\mu \equiv J_{2}/J_{1}$. Fig.\textsuperscript{[1]} shows an equivalent coupled 2 spin-chains (zigzag chain) suited for EPT. The theoretical tools employed are the bosonization-Sine-Gordon (SG) theory\textsuperscript{[2]}, the RG (renormalization group)-level spectroscopy analysis augmented by exact diagonalization\textsuperscript{[3, 4]} and numerical RG analysis based on iTEBD (infinite time evolving block decimation)\textsuperscript{[5]}, a variance of DMRG (density matrix renormalization group) method\textsuperscript{[6]} for infinite 1D systems. The key finding is that the Néel order and dimer order alternate numerous times when approaching the ferromagnetic transition point $(J_{2}/|J_{1}|, \Delta) = (1/4, 1)$ along the Lifshitz line which separates the character of the spin-spin correlation in the xy-plane from commensurate to incommensurate. The reported Néel order is indeed counterintuitive since neither ferromagnetic NN nor antiferromagnetic NNN exchange couplings favor antiferromagnetism.

When $J$s are both positive, the ground state phase diagram of the model is well understood\textsuperscript{[2, 4, 5]}. Particularly, for small $\mu$ and with easy-plane anisotropy $\Delta < 1$, it is a gapless Tomonaga-Luttinger liquid (TLL). The gapless TLL phase has instabilities toward ordered states\textsuperscript{[2]}. Néel order will be induced by easy-axis anisotropy $\Delta > 1$, whereas dimer order appears when $\mu$ is larger than some critical $\Delta$-related value. For instance, transition takes place when $\mu \approx 0.24$ at $\Delta = 1$. The quantum phase transition to these ordered phases can be understood within a bosonization-SG theory\textsuperscript{[2]}.

Now in the case $J_{1} < 0$ of our interest, the bosonization-SG theory of the TLL phase at $J_{2} = 0$ is known exact, and the “infinite” alternation of energies of the Néel and dimer states was observed in the parameter space $0 < \Delta < 1$, and based on a perturbative treatment, from the Gaussian Hamiltonian, of the cosine term in the SG theory, the authors of\textsuperscript{[1, 9]} claimed that the same should occur deep inside the frustrated regime $|\mu| \sim 1/4$, and thus the realization of the true alternating appearance of the Néel and dimer phases. This prediction was first checked by the level spectroscopy\textsuperscript{[3, 4]}, showing 5 times of phase alternation between the two phases. This level spectroscopy analysis was, however, still based on the effective SG theory whose validity in the parameter regime $|\mu| \sim 1/4$ is correct only in a perturbative sense such as neglecting a higher frequency term $\cos(\sqrt{8}/3)$. Thus, authors of\textsuperscript{[1]} also carried out an independent iTEBD calculation of the original model (1), confirming only one dimer to Néel transition at $\Delta \sim 0.72$.
and one Néel to dimer transition at $\Delta \sim 0.93$. It was reported that iTEBD experienced a poor convergence for $0.95 \leq \Delta < 1$ and $0.25 < |\mu| < 0.35$.

The purpose of this letter is to study the controversial parameter regime in the above by EPT (entanglement perturbation theory), a recent many body method [10–13]. Our finding in this paper is summarized as follows: (1) EPT gives a converging, and hence exact phase diagram, including exact order parameters of Néel and dimer phases for the entire $0 \leq \Delta < 1$ along the Lifshitz line, see Fig.4 below. The dimer phase and Néel phase alternate only once, and for $\Delta \geq 0.93$, only the dimer phase exists. This means two points. (2) First, the perturbative SG analysis when applied to the deeply frustrative regime $|\mu| \sim 1/4$, is missing some physics, and (3) second, iTEBD, even with a large entanglement $\chi \sim 300$, is not as accurate as EPT with $\chi \sim 80$. This conclusion is not a surprise. Indeed, TEBD operates only on a few local spins, and therefore the deterioration of accuracy due to loss of information arising from successive Hilbert space truncation, typical of real-space RG, is expected worse than DMRG.

The EPT algorithms EPT-g1 and EPT-g2 both solve for ground state properties of systems with translational inhomogeneity will be reported elsewhere). EPT-g1 is especially suitable for infinite systems and is our starting point. It was previously applied to the $J_2 = 0$ case [14, 15], and its superb accuracy is demonstrated in the first confirmation of a prediction made by the conformal field theory [16, 17] on a long-range spin-spin correlation which can only be seen for thousands of lattice sites, and therefore the deterioration of accuracy is not a surprise. Indeed, TEBD operates only on a few local spins, and therefore the deterioration of accuracy due to loss of information arising from successive Hilbert space truncation, typical of real-space RG, is expected worse than DMRG.

The central idea of EPT-g1 is as follows. First is the system wave function in the matrix product state (MPS) [10, 19, 23] representation

$$|\psi\rangle = \sum_{\{\sigma\}} \text{Tr}[\xi_1^\sigma \cdot \xi_2^\sigma \cdots \xi_N^\sigma] |\sigma_1\rangle \otimes |\sigma_2\rangle \otimes \cdots \otimes |\sigma_N\rangle$$

(2)

where $\xi_i^\sigma$ is the local wave function matrix at site $i$, and $N$ is the number of sites. $\sigma_i(=\uparrow, \downarrow)$ refers to local states. The dimension $p$ of the square matrices $\xi_i^\sigma$, the entanglement, controls the accuracy of the wave function $|\psi\rangle$. In principle $|\psi\rangle$ with optimized $\xi_i^\sigma$ becomes exact with $p \to \infty$. The matrices $\xi_i^\sigma$ are optimized to maximize the variational energy:

$$\frac{\partial}{\partial \xi_i} \langle \psi (\xi_1, \xi_2, \cdots, \xi_N) | e^{-\beta H} | \psi (\xi_1, \xi_2, \cdots, \xi_N) \rangle = 0$$

(3)

where $\beta \to 0 [11]$. We then arrive at a generalized eigenvalue equation

$$X_i \langle \xi_1, \cdots, \xi_{i-1}, \xi_{i+1}, \cdots, \xi_N \rangle \xi_i = \epsilon Y_i \langle \xi_1, \cdots, \xi_{i-1}, \xi_{i+1}, \cdots, \xi_N \rangle \xi_i \quad (i = 1, 2, \cdots, N),$$

(4)

where $X_i$ and $Y_i$ are symmetric matrices depending on $\{\xi_i\}$, and $\epsilon$ is the eigenvalue. The eigenfunction $\{\xi_i\}$ with the largest $\epsilon$ describes the ground state wave function $|\psi\rangle$.

The density matrix $K \equiv e^{-\beta H}$ can also be written in matrix product form, the matrix product operator (MPO) [14, 15]. A generalization to quasi-1D naturally inherits this MPO of density matrix along with MPS for the wave function. But now, cf. Fig.2, MPS is defined on a 2-spin composite (grey and black vertical two circles) and MPO is in a complex shell by shell (inner product direction) and layer by layer structure (vertical direction). We first deform a zigzag chain into two chains shown in Fig[1]. The Hamiltonian is rewritten as a summation of local bond Hamiltonians,

$$H = \sum_{\text{bond}} H_{\text{bond}}$$

(5)

There are four types of bonds. Bond $\{1, 2\}$, $\{3, 4\}$ etc form the first series. $\{2, 3\}$, $\{4, 5\}$ etc belong to the second. These two series account for NN interactions. The third is $\{1, 3\}, \{2, 4\}, \{5, 7\}, \{6, 8\}$ etc and the forth is $\{3, 5\}, \{4, 6\}, \{7, 9\}, \{8, 10\}$ etc. They account for NNN interactions. The way to group is not unique, but it does not affect the result. It is immediately seen that $[H_{\text{bond}}^i, H_{\text{bond}}^j] = 0$ if they are in the same series. This is the criteria to group the bonds. Small $\beta$ safely separates series in the density matrix as follows

$$e^{-\beta H} = \prod_{\text{bond}} e^{-\beta \xi_{\text{bond}}} + \mathcal{O}(\beta^2)$$

$$\approx \cdots e^{-\beta \sum_{1-st \text{ series}} H_{\text{bond}}^1} e^{-\beta \sum_{2-nd \text{ series}} H_{\text{bond}}^2} \cdots$$

(6)

And for each series, we have

$$e^{-\beta \sum_{\text{series}} H_{\text{bond}}} = \cdots f_\alpha \otimes g_\alpha \otimes f_\gamma \otimes g_\gamma \cdots + \mathcal{O}(\beta^2)$$

(7)

where repeating indexes follow Einstein summation convention. $f$, for instance for the XXZ model, takes four $2 \times 2$ matrices: $1, \sqrt{3}S^x, \sqrt{3}S^y, \sqrt{3}S^z$ on a site. And $g$ likewise on the other site of a bond. We use $\text{shell}$ to denote a series in $[17]$. There are four shells in total. They are coupled in the direction of in and out of the paper in Fig[2] by the inner product of the operators on the same site. Also note that MPO contains two layers, composed of two vertically aligned spins, one in the top chain and the other in the bottom chain. We finally have an explicit
and \( \Gamma \) come 4-leg tensors. The two effective legs run from 1 to \( l \) and \( l' \) are the indexes involved in the direct product. If they 

\[ \xi \]

lipse in Fig. 3. It is also noted that, reflecting the \( \Gamma \) repetition structure, the local wave functions are also of two species \( \xi \). The generalized eigenvalue problem (4) is solved iteratively for the local wave functions, and after convergence, energy etc can be calculated as before [11, 13].

The calculation on zigzag chains is more expensive compared with a single spin chain because one now has to deal with the larger number of local states and larger indexes in MPO. Let us discuss the major time consumption. There are two. The first are generalized eigenvalue equations whose size is \( mp^2 \) with \( n \) being the number of effective local states. The second is an eigenvalue decomposition of a transfer matrix in the horizontal direction in Fig.3, a basis unit involving \( \Gamma^1 \) and \( \Gamma^2 \) and in total \( 4 \) 2-spin composites connected to \( \Gamma^{1,2} \). Its size is \( mp^2 \) where \( m \) is the size of the 3 bonds in the direct product between \( \Gamma^1 \) and \( \Gamma^2 \). In contrast to \( n = 2 \) and \( m = 4 \) in a spin chain, they are 4 and 64 now. It is seen that the eigenvalue decomposition of \( mp^2 \times mp^2 \) transfer matrix becomes especially large.

Fortunately, there is a simple way to overcome this difficulty. The key point is in the original idea of EPT-g1, namely convert the Hamiltonian eigenvalue problem to that for the density matrix \( K \equiv e^{-\beta H} \), noting \( e^{-\beta H} \rightarrow 1 - \beta H \), leading to an MPO representation of the density matrix. A simple example is the product of two local density matrixes, \((1 - \beta H_{ij}) \otimes (1 - \beta H_{kl}) \sim 1 - \beta H_{ij} - \beta H_{kl} + \beta^2 H_{ij} \otimes H_{kl} \), where the last term is unimportant in the \( \beta \rightarrow 0 \) limit. \( \beta = 10^{-6} \sim 10^{-7} \) is used in this work like before [11]. Likewise, \( \Gamma^{1,2} \) should contain such higher order terms in \( \beta \) which we can consistently discard. A rule to do so is in the fact that, in \( f_{\alpha} \) and \( g_{\alpha} \) matrixes, \( \alpha = 1 \) is identity matrix while other three terms are all order \( \sqrt{\beta} \). We should keep in mind that whenever there is a term of order \( \sqrt{\beta} \), there should be a companion order \( \sqrt{\beta} \) term at the other end of the bond. This amounts to keep, among 64 index combinations in \( \alpha \beta \gamma \), those terms with only one of \( \alpha \beta \gamma \) not being 1, namely, 111, 112, 121, 211, 113, 131, 311 and 114, 141, 411, there are total 10.

To test the algorithm, we have calculated the dimerization \( d \equiv \langle \hat{S}_i \hat{S}_{i-1} - \hat{S}_i \hat{S}_{i+1} \rangle \) for \( \mu = 0.5871 \) and \( \Delta = 0 \) where \( d \) is maximum when both \( J_s \) are positive [8]. The EPT result is 0.7906137 when \( p = 35 \) vs. 0.7906135 by DMRG [8]. By comparing with their results for \( \mu = 2.5 \) and \( \Delta = 0 \), the most difficult case studied in [8] and probably the same for EPT, it is found that EPT converges faster in entanglement. Since \( p \)-equivalent \( \chi = 200 \sim 300 \) is used in iTEBD [4, 13], the entanglement needed by EPT to achieve the comparable accuracy is much smaller than iTEBD.

The main result of this letter is the phase diagram shown in Fig. 4 where \( \Delta \) approaches the isotropic point 1 from below along the Lifshitz line. The \( \xi \) order parameter is \( \langle \hat{S}_i^x \rangle \) and the dimer order parameter \( D_{xy} \) is
defined by

\[ D_{xy} = \left( S_{i-1}^x S_i^x + S_{i-1}^y S_i^y \right) - \left( S_{i-1}^x S_{i+1}^x + S_{i-1}^y S_{i+1}^y \right) \]  

The main difference between our phase diagram and that of [1] is that our calculation converged solidly for all \( \Delta \) up to 1, whereas iTEBD failed for 0.95 < \( \Delta \) < 1. It also shows no more alternation of the Néel and dimer orders for \( \Delta > 0.93 \), disagreeing the prediction of [1] that there should be an infinite number of such transitions. The failed prediction, we argue, is most likely due to the failure of perturbative analysis of the Sine-Gordon theory deep in the frustrated regime \( |\mu| \sim 1/4 \). Technically, the Lanczos diagonalization not only fails to converge due to highly degenerate nature for 0.995 \( \leq \Delta \leq 1 \) as pointed out in [1], but also, we argue, yields not quite accurate energy levels for \( \Delta > 0.95 \) so that the level spectroscopy analysis concludes incorrectly for this parameter regime. As for the nature of the Néel to dimer transition, our calculation, Fig 3 does not agree with [1] that it is first order. Rather it appears second order.

While our calculation converged solidly, there is a clear evidence of a hard competition between the Néel and dimer orders near the transition point. In Fig 5 we have plotted the Néel and dimer order parameters as a function of entanglement for \( \Delta = 0.722 \). Dimer order dies out only at relatively large entanglement. The same happens at the other phase boundary \( \Delta = 0.930 \) with even more subtle competition between the two orders since their magnitudes are both small.

To summarize, we have developed EPT for quasi-1D strongly correlated quantum systems. There are two immediate applications of interest. First is the 1D Hubbard model with long-range hopping, long-range Coulomb interaction, orbital degeneracy and Hund-coupling in the pursuit of itinerant ferromagnetism and triplet superconductivity. Second is a spin tube, namely the effective sites now contain the third spin, taking into account third nearest neighbor exchange interactions. Works are currently under way, and will be reported elsewhere.

Note added – After the completion of the paper, we have heard from the authors of [1] that the Lifshitz line along which we and they have done the calculations is actually not precise, and that their prediction of the Néel phase could still be observed close to the ferromagnetic point, \( \Delta = 1.0 \). We have explored \( \Delta = 0.98 \) and \( J_2/J_1 = 0.2552 \) (newly suggested point) to 0.2568 (our and their previous calculation point). We observed a TLL \( (J_2/J_1 = 0.2552) \) to dimer order transition of the second order type which is indistinguishable from the BKT (Berezinskii-Kosterlitz-Thouless) type [24, 25] due to an essential singularity [26], the transition point being 0.2560. We have also calculated the chiral order parameter \( S_1 x S_2 \) in this parameter region to be strictly zero. In the meantime, we have observed a finite chiral order parameter for the point \( \Delta = 0.9 \) and \( J_2/J_1 = 0.35 \) which is deeply in the chiral phase. We thus conclude that our main claim in the paper is unchanged.

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