Comb tensor networks

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(Dated: March 4, 2019)

In this paper we propose a special type of a tree tensor network that has the geometry of a comb—a 1D backbone with finite 1D teeth projecting out from it. This tensor network is designed to provide an effective description of higher dimensional objects with special limited interactions, or, alternatively, one-dimensional systems composed of complicated zero-dimensional objects. We provide details on the best numerical procedures for the proposed network, including an algorithm for variational optimization of the wave-function as a comb tensor network, and the transformation of the comb into a matrix product state. We compare the complexity of using a comb versus alternative matrix product state representations using density matrix renormalization group (DMRG) algorithms. As an application, we study a spin-1 Heisenberg model system which has a comb geometry. In the case where the ends of the teeth are terminated by spin-1/2 spins, we find that Haldane edge states of the teeth along the backbone form a critical spin-1/2 chain, whose properties can be tuned by the coupling constant along the backbone. By adding next-nearest-neighbor interactions along the backbone, the comb can be brought into a gapped phase with a long-range dimerization along the backbone. The critical and dimerized phases are separated by a Kosterlitz-Thouless phase transition, the presence of which we confirm numerically. Finally, we show that when the teeth contain an odd number of spins and are not terminated by spin-1/2’s, a special type of comb edge states emerge.

I. INTRODUCTION

Modern condensed matter physics to a large extent relies on numerical simulations. The Density Matrix Renormalization Group (DMRG)$^{1,2}$ algorithm has established itself as one of the most powerful numerical tools for strongly-correlated one-dimensional systems. The method is variational and in most cases its accuracy is well controlled by the number of states kept and the number of sweeps—two external parameters of the algorithm. In contrast to Wilson’s numerical renormalization group approach, the selection of the effective basis vectors is not based on energies, but on the eigenvalues of the density matrix, or equivalently, on the entanglement. The success of DMRG in one-dimension inspires its further application to two-dimensional systems by mapping the 2D lattice to a 1D path in a snake-line way. However, according to the area law$^3$ the entanglement for multiple leg ladders grows exponentially with the number of legs. This limit the width $L$ of these quasi-two dimensional lattices, and the current state of the art for two-dimensional cylinders with DMRG is about $L \approx 6 - 14$, depending primarily on the number of states per site in the model.

The reformulation of conventional DMRG in terms of matrix product states (MPS)$^{12}$ led to a number of generalizations, most notably for higher dimensions. Probably the best known example is the projected entangled pair states (PEPS)$^{13,14}$ network, where the tensors living on each physical site are coupled by nearest-neighbor links. These higher dimensional tensors split the entanglement uniformly over many bonds, so the network satisfies the area law regardless of the width of the system. On the other hand, the presence of loops in the network, and the higher number of indices on each tensor lead to much higher complexity in terms of the number of states per link, compared to DMRG. Currently, for many 2D problems, DMRG and PEPS are complimentary.

In this paper we study another tensor network that goes beyond 1D—the comb, shown in Fig. 1. A comb consists of a 1D backbone, with 1D teeth projecting out from it. A comb is a special case of a tree tensor network; it can also be considered a generalization of a Y-junction network, for which DMRG algorithms were developed for Heisenberg spin systems.$^{15}$ It is also related to fork tensor networks introduced recently for multi-orbital Anderson impurity model.$^{16}$ It may be that the physical interactions and sites in a system appear in a comb geometry, in which it is natural to use a comb tensor network to study the system—see Fig. 1(a). Another, less obvious case where a comb may be particularly useful is for a 1D system composed of complicated, highly entangled but finite units. In this case a comb geometry can be effective in isolating the intra-unit entanglement into a tooth, leaving the backbone less entangled—see Fig. 1(b). To be studied with the comb tensor network the model has to satisfy the one-dimensional area law (at least along the backbone); otherwise the algorithm has the same limitations as a snake-like DMRG for a two-dimensional system.

The motivation to develop a comb tensor network is two-fold. From a numerical point of view, the complexity of the algorithm scales with an auxiliary bond dimension $D$ as $D^3$, which is much lower than two-dimensional PEPS. On the other hand, for a fixed bond dimension the complexity of the comb tensor network is higher than in DMRG. However, the higher dimensional tensors placed
along the backbone, split two channels of entanglement: one within the clusters and one along the backbone. In DMRG the entanglement from these two channels is superimposed, which results in a large bond dimension. Thus, when the entanglement along the backbone and within the teeth are of the same order of magnitude, the comb tensor network requires much lower bond dimension than conventional DMRG.

From the point of view of possible applications, the comb tensor network is also very promising. First of all, a comb geometry itself can lead to exotic phases and phase transitions even for the simplest Heisenberg model. Among them are various junctions of spin chains\cite{12,13} and quantum wires\cite{16–19} that have attracted a lot of attention over the past decade. Modern technologies allow one to realize comb lattices in cold atom and adatom experiments. In addition, combs may be used to study coupling effects in clustered materials, such as the recently discovered 24-spin boleite\cite{24–26}.

The comb tensor network may also be used for quantum chemistry. There are molecules which have backbone and chain geometries resembling combs, such as triglycerides or polysulfones\cite{27,28} for which one might make Hubbard-like models. At the more ab initio level, consider quantum chemistry DMRG, where the sites in the algorithm are orbitals, including core orbitals as well as valence. The entanglement within a core gets added to the interatomic entanglement. It would be natural to have the core degrees of freedom of an atom appear as sites of a tooth, with the atoms connected along the backbone, separating the two types of entanglement. In a similar spirit, associating the teeth of the comb with finite-size Sachdev-Ye-Kitaev (SYK)\cite{10,13} clusters, intercluster entanglement can be dealt with separately.

The rest of the paper is organized as follows. In Sec. II we will set up the basic properties of the comb tensor network, including the transformation of the wave function between the comb and the traditional matrix product-state forms and the mixed canonical form of the comb network. In Sec. III we provide further details on the numerical approach and discuss the variational optimization algorithm of the wave function as a comb tensor network. In Sec. IV we apply these algorithms to study the low-energy properties of the Heisenberg spin-1 comb lattice. We summarize our results in Sec. V.

II. QUANTUM STATE AS A COMB NETWORK

A. Mixed canonical form

Putting an MPS into a mixed canonical form (MCF) simplifies drastically both optimization and the following re-usage of the wave function\cite{2}. The MCF is characterized by the combination of left-orthogonal tensors on the left and right-orthogonal tensors on the right, with a site or link between them called the orthogonality center (OC). If the OC is a link, an extra two-index diagonal tensor is associated with the OC. No orthogonality is associated with OC itself. In variational optimization, the MCF simplifies a generalized diagonalization to a simple one, and fixes the norm of the wave function. When extracting local observables, the MCF allows one to skip the contraction of the complete tensor network, utilizing only those tensors that are located at and between the local operators of interest\cite{2}.

In analogy with a one-dimensional MPS we introduce an MCF for the comb tensor network. To reduce the complexity of the algorithm we associate any physical degrees of freedom on the backbone to the first sites of the teeth, so that the backbone tensors, as well as the tooth tensors, only have three indices. Let us consider the case where the OC is on a tooth, and specifically on a tooth, so that the backbone tensors, as well as the tooth tensors, only have three indices. Let us consider the case where the OC is on a tooth, and specifically on a link, as shown in Fig. 2. The notion of the left and right normalized tensors of an MPS is generalized for the comb. We attach arrows to each link, with each arrow pointing along the path connecting to the OC. For each tensor, one link (the forward link) points away from the tensor. To give the orthogonality condition, we contract a tensor with its conjugate, contracting over all indices except the forward link. The resulting two index tensor is equated to the identity tensor, as shown in Fig. 2. Note that a Hermitian conjugated tensor has its arrows reversed.

In Fig. 2(b) we show graphical representations of the three types of orthogonality conditions. As in the case of MPS, the MCF simplifies the diagonalization in the variational optimization at each step from a generalized to a simple eigenvalue problem, reducing the computational cost and stabilizing the algorithm. The MCF also simplifies measurement of local observables; for an operator on a tooth, the complexity is equivalent to the standard DMRG complexity $D^3$. Computing correlation functions along the backbone has higher complexity $D^4$.\[a\]
however the contraction of the network on each tooth can be skipped and the complexity does not depend explicitly on the length of the teeth.

**B. From an MPS to a comb**

In principle, an arbitrary tensor network representation of a wavefunction on a finite number of sites can be transformed into an MPS, and vice-versa. The resulting MPS may, however, have large bond dimension $D$, an example being the conversion of a PEPS wavefunction for a 2D cluster into an MPS. Even in cases where the bond dimension of both representations is modest, it may not be so easy to transform back and forth, particularly if the tensor network has loops. An advantage of the comb network is that there is a very simple, robust algorithm for transforming back and forth between a comb and an MPS, using simple contractions and SVDs. In Fig. 3 we illustrate this transformation. Note that the orthogonality center (OC) has to be moved to any sites being contracted, so that any pair of tensors being contracted includes the OC. This assures that the singular values obtained after the SVD decomposition are equivalent to the Schmidt values and thus allow optimal truncation.

**III. VARIATIONAL OPTIMIZATION**

The direct mapping between the comb tensor network and an MPS described in the previous section allows one to use standard DMRG on the MPS to get a comb ground state, but in general it is more natural and probably more efficient to optimize within the comb geometry. Since the comb has no loops, the optimization is similar to that of DMRG, but there are specific techniques for the teeth versus the backbone sites. In addition, it is useful to represent the Hamiltonian in a comb tensor network form. Not wishing to introduce an additional term for this comb operator, we will call this a projected entangled pair operator (PEPO), borrowing from projected entangled pair state (PEPS) terminology. There are five main ingredients in the proposed algorithm: the representation of the Hamiltonian as a PEPO, the creation of a good initial wave-function, the recipe for a full sweep to update all tensors in the network, the recipe for a back-bone sweep to update only auxiliary tensors along the backbone, and efficient measurements of observables.
A. Hamiltonian

Let us consider as an example an $N \times L$ comb lattice shown in Fig. 1(a) with Heisenberg nearest-neighbor interaction with coupling constant $J_i$ along the tooth and $J_{bb}$ along the backbone:

$$H = J_{bb} \sum_{i=1}^{N-1} S_{i,1} \cdot S_{i+1,1} + J_i \sum_{i=1}^{N-1} \sum_{j=1}^{L-1} S_{i,j} \cdot S_{i,j+1},$$  \hspace{1cm} (1)$$

where $i$ indicates which tooth and $j$ indicates the site within the tooth, starting from the backbone. The construction of the MPO for $2 \leq j \leq L$ is standard and is well described, for example, in Ref. 5. For completeness we provide the explicit form of the MPO. For $2 \leq j \leq L - 1$ it is given by

$$H(i,j) = \begin{pmatrix}
I & & & \\
S^- & & & \\
S^+ & & & \\
S^z & & & \\
hS^z & J_iS^+ & J_iS^- & J_iS^z & I
\end{pmatrix},$$  \hspace{1cm} (2)$$

where $I$ is an identity matrix, $h$ is an external magnetic field, and zero-elements are marked by dots for clarity. Both $I$ and $S$ are operators; if the operators are written in terms of indices then the total rank of the tensor is four. At the tip of the tooth $j = L$ the tensor is given by the first column of this matrix and has rank three. At the backbone the rank of the on-site tensors that represent the Hamiltonian is higher, because on top of two physical bonds, they contain three auxiliary bonds that connect them to all their neighbors. Since there are no loops in the comb, the construction of these PEPO operators is as simple as the conventional construction of the MPO. In the tensor $H_{\alpha,\beta,\gamma}$ where $\alpha$ is the label $\alpha$ for the left, $\beta$ for the upper and $\gamma$ for the right auxiliary legs. The tensor is very sparse, but we can represent the slice $\beta = 1$ as a matrix, and then the only nonzero elements left have $\alpha = 5$ and $\gamma = 1$, so these values can be written as a vector in the index $\beta$:

$$H_{\beta=1} = \begin{pmatrix}
I & & & \\
S^- & & & \\
S^+ & & & \\
S^z & & & \\
hS^z & J_{bb}S^+ & J_{bb}S^- & J_{bb}S^z & I
\end{pmatrix},$$  \hspace{1cm} (3)$$

We find it instructive to show the pictorial representation of non-zero elements of the backbone PEPO tensor (see Fig. 4), where the gray boxes state for identity operators, the green box includes three elements of the first column of the matrix in Eq. 3 [$S^-, S^+, S^z$], the blue box contain three elements of the last row of this matrix [$J_{bb}S^+, J_{bb}S^-, J_{bb}S^z$], the orange box encodes three elements for the interaction within the tooth [$J_bS^+, J_bS^-, J_bS^z$], and the yellow box corresponds to the field-term, which is set to zero throughout a paper. Of course, this form of the Hamiltonian is model-dependent, and if the interaction is more complicated and extends beyond nearest-neighbors the structure of the PEPO is not as simple.

B. Sweep

Let us first explain the variational optimization of the comb tensor network starting from some initial state. Later, we shall come back to the creation of a good initial wave-function.

Fig. 3(a) shows the full tensor network that evaluates the total energy consisting of a ket wave-function $|\psi\rangle$ (lower surface of blue and green tensors), the Hamiltonian in local representation (middle surface with yellow (MPO) and orange (PEPO) tensors) and a bra vector $\langle \psi |$, represented by the upper surface of tensors. The contraction over all physical and auxiliary links gives the total energy $E = \langle \psi | H | \psi \rangle$. We assume that the wave-function is always written in a canonical form defined above, so the normalization can be omitted here. Note that the number of tensors that represent the wave-function is larger than the number of tensors that represent the Hamiltonian, due to the use of auxiliary backbone tensors without physical indices. So, PEPO backbone Hamiltonian tensors are contracted with the first MPS tooth tensors.

In a full sweep we go through the whole network shown in Fig. 3(a) and update the on-site tensors iteratively. During the first half-sweep we move from left to right, and then reverse during the second half-sweep. Within each tooth we move from the backbone to the edge and back. At each iteration we update two on-site tensors. A generalization to single- or multiple-site update is straightforward.

In most of the cases the update can be reduced to the diagonalization of the effective Hamiltonian that consists
of the left and right environments and a pair of MPO between them. So on a tooth this is standard 1D DMRG. For a pair of auxiliary backbone tensors, one can view a fully contracted tensor network on a tooth as an effective MPO as shown in Fig. 5(b). The effective Hamiltonian for two auxiliary tensors also looks like the form in 1D DMRG; see Fig. 5(d). However, there is a special case that cannot be reduced to the standard network with two environments and two MPOs. When an auxiliary tensor is updated together with the first MPS tensor on a tooth (a connect update) the effective Hamiltonian contains three environment each of which is connected to a PEPO in the middle, as shown in Fig. 5(c).

Depending on the model, we find it useful to alternate the full sweep with a simplified one that updates only the backbone tensors. In particular this make sense when the backbone chain is critical while the teeth are gapped. This short sweep consists only of backbone updates.

We increase the number of states every half sweep. We control three parameters that restrict the number of states: $\chi$ is the bond dimension within the tooth; $\zeta$ restricts the bond in connect updates; and $\lambda$ controls the dimension of a bond along the backbone. When all bonds are equivalent $\chi \approx \lambda \approx \zeta \approx D$ the complexity of the backbone update is $D^5$, the complexity of the connect update is $D^4$, and the complexity of the tooth update is only $D^3$, as in the standard DMRG. Of course, the complexity is very different when the number of states on the tooth and on the backbone are of a different order of magnitude. For example, if the state on the tooth is close to the Affleck-Kennedy-Lieb-Tasaki state, the bond dimensions $\chi$ and $\zeta$ are small in comparison to $\lambda$ and the complexity is only $\lambda^3$.

C. Initial wave-function

We start our simulation by forming a rough initial wave-function. As in 1D DMRG there are various different approaches and depending on the problem one or another can be preferable. The simplest one is to start with a product state - tensor network with auxiliary bond dimension $D = 1$. If Abelian symmetry is used, the chosen product state should satisfy it. This approach is efficient assuming one slowly increases the bond dimension in multiple sweeps.

In variational optimization of an MPS it is well established that one of the most accurate and unbiased guesses can be obtained through infinite-size DMRG. It starts with small clusters that can be solved exactly, say four spins, and at each iteration the size of the 1D chain increases by two spins. Infinite-size DMRG produces a guess wave-function with finite bond dimension (typically $D \approx 10-50$) that naturally preserves the symmetry of the network with respect to the center of a chain and provide an excellent starting point for the following finite-size routine. We find that an excellent initial wave-function comes from using infinite-size DMRG on a chain with $2L$ sites to produce an initial guess for the MPS tensors on the teeth. The Hamiltonian used in infinite-size DMRG should be as close as possible to the original MPO on a tooth. One can translate the $2L$ MPS tensors to define the tensors of two adjacent teeth. Since this wave-function ignores the backbone interactions, one can slightly truncate the MPS bond dimension at the backbone (the middle) to make a subsequent initial set of backbone sweeps fast and inexpensive. These backbone sweeps can start either with the product-teeth states, or with an infinite-size DMRG along the backbone.

One can also use an infinite-size DMRG to initialize the backbone after constructing guesses for the tooth tensors. By treating a fully contracted tooth as an MPO with large 'physical' bond dimension, one can initialize the auxiliary guess tensors by performing a standard infinite-size DMRG on the backbone.

In the present paper we produce a guess wave-function by performing infinite-size DMRG in both directions.

D. Implementation of Abelian symmetries

When allowed by the model, the implementation of Abelian symmetries in the comb tensor network allows one to select a specific symmetry sector and thus to compute the energy gap to magnetic excitations in a trivial way. Moreover, this reduces significantly an effective size
of the Hilbert space and therefore speeds-up the convergence at each iteration. Finally, in the presence of a $U(1)$ symmetry the tensors have block-diagonal structure that simplifies their storage and access.

In order to keep track of Abelian symmetry, one can group the states on each index by quantum number (say $S^z$). Then one stores only nonzero blocks in the tensors, and the arrows in the diagrams in Fig. 2(a) indicate how one adds up the quantum numbers. For the nonzero blocks, the sum of the in-going quantum numbers must equal the outgoing quantum number on each tensor. At the OC, where all arrows point inward, one can regard the OC tensor as a wave-function in an orthonormal basis. Each link connecting to the OC tensor represents an orthonormal set of basis states living on the associated branch of the comb, and the whole basis is the direct product over all the branches. The total quantum number of the OC-tensor wavefunction is the sum of all incoming quantum numbers. If OC is associated with the tensor that contains physical index, its contribution should also be added to the total quantum number. All nonzero blocks of the OC tensor have this same quantum number.

**E. DMRG versus comb**

Let us now compare the complexity of optimizing the comb versus an equivalent MPS. First, we will consider a square spin-1/2 comb with antiferromagnetic nearest-neighbor Heisenberg interactions, where we set both coupling constants of the Hamiltonian $\hat{H}$ to be equal $J_{\text{hh}} = J_{t} = 1$.

The algorithms split the system into two parts in different ways. It is useful to compare the singular values (square root of the Schmidt values) for these various cuts. Results for $10 \times 10$ and $20 \times 20$ combs are shown in Fig. 6. Note that in both algorithms the cut $\alpha$ (or $\tilde{\alpha}$) across the backbone leads to the same bi-partition of a lattice and thus the singular values are equal (red dots). In variational optimization of a comb tensor network it is natural to cut all or part of a tooth from the rest of the system as sketched in the inset of Fig. 6(a). Here we show singular values for three example cuts: the whole tooth, which is the same as tooth-connect (cut $\delta$, blue color); a two-thirds cut ($\gamma$, green color); and a one third cut ($\beta$, magenta color) of the tooth. We see from Fig. 6(a,c) that the entanglement is largest along the backbone and, as expected, decays very fast upon approaching the end of the tooth.

By contrast, in DMRG applied to a comb lattice the entanglement is smallest on the bond that connects the upper and lower edges of two teeth. It is significantly larger within the teeth, where the entanglement from both the backbone and from within the tooth are present. We stress that the drastic difference between the decay of Schmidt values in DMRG and the comb network is caused by the different cuts, not any details of the optimization.

We measure the number of states necessary at each bond to keep the truncation error below $10^{-4}$ for singular values (and $10^{-8}$ for Schmidt values). The results are summarized in Fig. 7(a). The bond dimension along the backbone $\lambda$ grows slowly with the system size. Such slow decay qualitatively agrees with the slow logarithmic divergence of the bond dimension for critical systems. The same is true for the bond within the tooth and the bond that connects the tooth with the backbone for the comb network. On the other hand, the bond dimension of the MPS wave-function optimized with DMRG grows much faster with the system size.

We estimate the complexity of the DMRG and variational optimization of a comb tensor network based on the bond dimensions. For DMRG the most consuming term is the contraction of the network in the middle of the tooth with the single-iteration complexity $D^3$, that has to be performed about $N^2$ times per half-sweep, so the total complexity is $D^3 N^2$. For a comb tensor network, the leading term of the complexity is given by simultaneous optimization of two backbone tensors that has single-iteration complexity $\lambda^3 C^2$ and has to be performed only $N$ times, so the total complexity is given by $\lambda^3 C^2 N$. In the large $N$ limit we expect the comb representation to be generally more efficient; for smaller $N$ the comparison depends on the difference in bond dimensions for the MPS along the backbone versus in the middle of the comb.
For a spin-1/2 Heisenberg comb, the optimization of the wave-function directly in a comb geometry gives slightly lower complexity only starting from the lattice with 30 \times 30 sites as shown in Fig. 7(b). However the two curves follow each other very closely and we expect these behavior to persist even for larger clusters.

Now we compare the computational cost of a spin-1 comb with strong coupling along the teeth $J_t = 1$ and weak backbone interaction $J_{bb} = 0.1$. The main difference from the previous case, is that now the teeth are essentially gaped sub-systems, each of which corresponds to the Haldane finite-size chain. At the edge of each tooth, spin-1/2 edge states emerge. As soon as a backbone coupling is non-zero these edge states interact between themselves. While spin-1/2 edge-states emergent at the backbone interact with the coupling $J_{bb}$, the effective coupling between the edge states at the end of the teeth is exponentially suppressed with the length of the tooth. This leads to a massive degeneracy of the ground state with exponentially small splitting between many in-gap states. In order to avoid this, we replaced spin-1 degrees of freedom at the end of each tooth by a spin-1/2. Below we will provide the detailed study of the model, but here we only focus on the algorithm efficiency.

Since the teeth are expected to be in a gapped state, according to the area law the corresponding bond dimension within the teeth is expected to approach a constant in the thermodynamic limit. Indeed one can observe it in the green and magenta lines in panel (c) of Fig. 7. On the other hand, spin-1/2 edge states are expected to form a critical chain and so the bond dimension across a backbone diverges logarithmically with the number of teeth. In DMRG the entanglement induced by the critical chain is also seen inside the teeth, so in DMRG each bond dimension diverges with the length of the comb. For the chosen set of parameters, comb tensor network turns out to be more efficient than standard DMRG starting from very small systems of 8 \times 8 sites (see Fig. 7).

To summarize, there exist classes of models, for which the optimization of the wave-function directly in the comb geometry has lower computational cost than standard one-dimensional algorithms. Moreover, in some particular cases, e.g. dimerized state on the backbone, the comb tensor network is less likely to get stuck in local minima than the DMRG, in which the dimers are formed between sites that are far apart.

In the following section we use this algorithm to study the Heisenberg model on a spin-1 comb lattice.

\section{IV. SPIN-1 HEISENBERG MODEL ON A COMB LATTICE}

\subsection{A. The model}

The spin-1 Heisenberg chain has long been known to have a finite bulk gap\cite{22} and spin-1/2 edge states\cite{23,24}. This is one of the simplest examples of a topologically non-trivial state realized in spin systems. Here we couple a set of these spin-1 chains into a comb with Heisenberg nearest-neighbors interactions defined by Eq. 1.

In the absence of any backbone interaction the edge states of a tooth couple. If the number of spins per tooth is even then they form a singlet and the first excited state is a Kennedy triplet\cite{22}. When the number of sites on a tooth is odd, the ground-state is a triplet, and the first excitation is a singlet. In both cases, the energy splitting between the ground-state and the in-gap excited state(s) decays exponentially with the length of the tooth. Introducing a backbone interaction, one couples the backbone edge-states into a spin-1/2 chain. This chain is then decorated with the spin-1/2 degrees of freedom at the tips with an effective coupling constant that decays exponentially fast with tooth length. So, for large $L$ the pendant spins cause a macroscopic degeneracy of the ground state with $2^N$ states, where $N$ is the number of teeth. To avoid
this massive degeneracy, for most of our calculations we remove the emergent spin-1/2 degrees of freedom at the end of each tooth by replacing the last spin-1 site by a spin-1/2. We will come back to the model with all spins-1 and study the effect of the edge spins at the end of the paper.

B. Critical spin-1/2 chain

When the backbone coupling is small the emergent spin-1/2 edge states, coupled to each other along the backbone, are only slightly changed and form a spin-1/2 chain. According to conformal field theory (CFT) the Heisenberg spin-1/2 chain is critical and can be described by the Wess-Zumino-Witten SU(2)\(_1\) critical theory. As a confirmation to that, we extract the critical exponent \(d\) from the decay of the Friedel oscillations. The latter naturally appears in the finite-size chain, because open edges of a critical spin-1/2 chain favor dimerization, and therefore fix the boundary conditions. We define the local dimerization order parameter as an absolute value of a difference between spin-spin correlations on the neighboring bonds on a backbone:

\[
D_{bb}(i, N) = |\langle S_{i,1} \cdot S_{i+1,1} \rangle - \langle S_{i+1,1} \cdot S_{i+2,1} \rangle|,
\]

where \(i\) is the tooth index and \(N\) is the total number of teeth. Then according to boundary CFT the dimerization decays away from the boundary as:

\[
D_{bb}(i, N) \propto \frac{1}{[N \sin(\pi i/N)]^d},
\]

where \(d = 1/2\) is a critical exponent of the WZW SU(2)\(_1\).

In Fig. 7 we plot the dimerization profile of the Friedel oscillations on a comb with 100 teeth and both 12 (blue) and 20 (green) sites per tooth. One can immediately see that the dimerization profile is independent of the width of the comb and the data collapse is almost perfect. A fit to the CFT prediction of Eq. 5 shown in Fig. 8(a) gives the critical exponent \(d \approx 0.54\), which is higher than the CFT prediction 1/2 due to the presence of logarithmic corrections in the Heisenberg spin-1/2 chain.

In addition, we can extract the central charge from the scaling of the entanglement entropy. Following Ref. 24 we define the reduced entanglement entropy on a backbone \(\tilde{S}_{bb}(i, N)\) with the Friedel oscillations removed:

\[
\tilde{S}_{bb}(i, N) = S_{bb}(i, N) - \zeta(S_{i,1} \cdot S_{i+1,1}),
\]

where \(\zeta\) is a non-universal parameter. Then, according to CFT, the reduced entanglement entropy scales with conformal distance \(d(i) = \frac{2N}{\pi} \sin(\pi i/N)\) as \(25\):

\[
\tilde{S}_{bb}(i, N) = \frac{c}{6} \log d(i) + s_1 + \log g
\]

From the fit shown in Fig. 8(b) we find the central charge \(c \approx 0.87\). This is in reasonable agreement with the \(S = 1/2\) chain value \(c = 1\).

![Figure 8](image_url)
emergent edge states (arrows) form a critical spin-1/2 chain (dashed red line), and (b) corresponds to the limit of strong backbone interaction with the Haldane chain on the backbone and critical spin-1/2 chain on the second sites of the teeth. Interestingly, the effective size of the critical spin-1/2 chain changes: in the limit of weak and strong backbone coupling the critical chain is formed out of the edge states of all \( N \) teeth, while in the intermediate coupling the system favors the state with very long Haldane chain that includes the first tooth, the backbone and the last tooth, so the critical chain is formed by the second sites on \( N - 2 \) teeth only.

We find that the two regimes sketched at the Fig. 10(a) and (b) are connected by a smooth crossover. In both regimes the system is critical due to the presence of a critical spin-1/2 chain, regardless of its location. In the thermodynamic limit the bulk gap is closed for all values of the backbone interaction. Also, the universality class of the underlying critical theory in the two regimes is the same. All the arguments above together with the extremely smooth change of all measured observables (more details below) suggests that two regimes are indeed connected by a crossover, rather than a phase transition.

In order to approximately locate the crossover we look at the location of magnetic excitations. For that we compute the lowest energy state in the sector of total magnetization \( S^z_{\text{tot}} = 1 \) and calculate the total local magnetization along the backbone and along the chain next to the backbone. The results are summarized in Fig. 11(a). When the backbone interaction is very small, the backbone has magnetization close to \( \sum_{i=1}^{N} S^z_{i1} = 1 \). The magnetization of the second row is non-zero due to finite (non-zero) correlation length in the Haldane phase on a tooth; it takes the opposite sign to the magnetization of the backbone due to strong antiferromagnetic correlations between the first two sites on the teeth. Upon increasing the backbone interaction the localization of the magnetic excitation on the backbone is smeared out and eventually the second row carries higher magnetization than the backbone. As shown in Fig. 11(a), the two lines cross when the backbone coupling is about \( J_{bb} \approx 1.1 \). We performed the calculations on square combs with 10 × 10 and 20 × 20 sites, finding that the finite-size effects are very small.

As a alternative method, we measure nearest-neighbor correlation in the middle of the backbone and between the first two sites on the middle tooth. In Fig. 9 we already discussed a qualitative difference between the local correlation pattern in the two regimes. In Fig. 11(b) we provide a quantitative comparison. We find that the backbone correlation becomes stronger than the connect correlations (correlations between the first two sites on a tooth) when the the backbone coupling exceeds \( J_{bb} \approx 1.3 \). This value is in a decent agreement with results on spin-1 localization presented above.
D. Zig-zag backbone and Kosterlitz-Thouless transition

In this section we show that one can manipulate the emergent critical spin-1/2 chain by adding frustration such as next-nearest-neighbor interaction. In a true spin-1/2 chain, this would lead to a gapped dimerized phase when the next-nearest-neighbor coupling constant $J_2$ exceeds $J_2 = 0.241^{[28]}$. The critical phase near $J_2 = 0$ is separated from the dimerized one by a Kosterlitz-Thouless (KT) phase transition$^{[24]}$. Both the critical phase and the KT transition are characterized by the WZW SU(2)$_1$ universality class. The KT critical line can be identified by vanishing logarithmic corrections, which are present inside the critical phase due to a marginal operator. Apart from the KT transition point, there is another special point, known as the Majumdar-Ghosh point and located at $J_2 = 1/2$, where the ground-state is given by an exactly dimerized state.

Here we include an antiferromagnetic next-nearest neighbor interaction along the backbone of a comb. The most natural way to imagine this is shown in Fig. 12 with a zig-zag ladder as a backbone, decorated with chains/teeth.

The traditional ways to locate the quantum phase transition, such as finite-size scaling of the order parameter or gap closing, which normally work well for gapped phases, cannot provide very accurate results for the Kosterlitz-Thouless transition, because the phase on one side of the transition is critical. Here we closely follow the method used to locate the end point on the critical lines described in Ref$^{[29]}$ and also used to locate the KT transition in the spin-3/2 chain$^{[30,31]}$. We briefly review the main idea here. From a CFT point of view, the transition between the critical and the dimerized phase is driven by tuning the coupling constant of the marginal operator in the Hamiltonian. When this coupling constant is negative (up to a convention), it can be renormalized to zero and the system can be described by WZW SU(2)$_1$ critical theory with non-vanishing logarithmic corrections that appear due to the renormalization process. By contrast, when the coupling constant of the marginal operator is positive, it cannot be renormalized to zero and lead to the gapped dimerized phase. At the KT transition this coupling constant is equal to zero and therefore the system is in the WZW SU(2)$_1$ universality class without any logarithmic corrections. So, the identification of the KT point is equivalent to the identification of the point where the critical behavior is maximally close to the WZW SU(2)$_1$ even on a finite-size system.

We consider a square comb with $20 \times 20$ sites and we set $J_t = J_{bb} = 1$. While tuning the next-nearest-neighbor coupling on the backbone we extract the apparent critical exponent and velocity. In order to extract the critical exponent we first note that the ground-state favors dimerization at the edges of the backbone. This effect is similar to one observed in the simple spin-1/2 chain and remains robust in the presence of decorating teeth. The dimerized states at the edge of a chain fix the boundary conditions and therefore induce the Friedel oscillations, which according to boundary conformal field theory takes the following form:

$$D_{bb}(i, N) \propto \frac{1}{N[\sin(\pi i/N)]^d},$$

where $D_{bb}(i, N)$ is an absolute value of the dimerization at site $i$ along the backbone with $N$ teeth. In the absence of logarithmic corrections the critical exponent predicted by boundary CFT is $d = 1/2$. When logarithmic corrections are not vanishing, the fit of the Friedel oscillations to this form gives an apparent critical exponent that deviates from the true CFT value. While approaching the thermodynamic limit the deviation slowly (logarithmically) goes to zero and in the thermodynamic limit the critical exponent is equal to $d = 1/2$ below and at the

Figure 11. (Color online) (a) Total magnetization localized at the backbone (blue) and the second row (red). (b) Local spin-spin correlations on the backbone (red) and between the first two sites on a teeth (blue). In both panels the results are for square combs of size $10 \times 10$ (squares) and $20 \times 20$ (crosses). A crossover is associated with the crossing point.

Figure 12. (Color online) An alternative representation of a comb lattice, in which both nearest- and next-nearest-neighbor interaction along the backbone naturally appears.
KT transition and zero above it. On a finite-size system, the curve is often very smooth. The apparent critical exponent extracted from a comb with \(20 \times 20\) sites is shown in Fig. 13(a). One can see that it monotonically decays and crosses the \(d = 1/2\) line around \(J_{2bb} \approx 0.37\).

The second way to locate the KT transition is based on the vanishing logarithmic corrections in the excitation spectra. For critical WZW SU(2)\(_1\) spin-1/2 chain with an even number of sites and fixed boundary conditions (see above) the ground-state is a singlet and the excitation energy scales with the length of the chain \(N\) as

\[
\Delta = \frac{\pi n v_{\text{eff}}}{N},
\]

where \(n = 1, 4, 9\) for the first triplet, quintuplet, and septuplet correspondingly. Due to logarithmic corrections the scaling can deviate on a finite-sized systems. One can therefore extract an apparent velocity as \(v_{\text{app}} = \frac{\Delta N}{\pi n}\), with the corresponding integer \(n\). When the logarithmic corrections vanish and the structure of the excitation spectrum (conformal tower) is preserved on a finite-size system, the apparent velocities extracted for various levels take the same value. Away from the KT transition inside the critical phase the velocities are different, due to the presence of the logarithmic corrections, while in the dimerized phase the spectrum is gapped and its structure changes drastically. Velocities extracted for three excited states with \(S_{\text{tot}}^z = 1, 2, 3\) as a function of the next-nearest-neighbor coupling are shown in Fig. 13(b). All velocities come close to each other around \(J_{2bb} \approx 0.37\), which agrees with the results from the analysis of the apparent critical exponent in panel (a).

Fig. [14(a) provides an example of the fit of the Friedel oscillations on a finite-size comb with \(20 \times 20\) and \(28 \times 28\) sites. For both sizes the extracted critical exponent agrees with the CFT prediction \(d = 1/2\) within 4%. We also look at the finite-size scaling of the excitation spectrum (only lowest energy states in the sectors with \(S_{\text{tot}}^z = 0, 1, 2, 3\)). The results are summarized in Fig. 14(b). The reference velocity is estimated by the singlet triplet gap of the largest available system size \((N = 28, L = N)\) as \(v_{\text{ref}} = (E_T - E_S)/N/\pi\). Then we use this non-universal constant to plot three lines lines of the CFT prediction \(\Delta = (\pi v_{\text{ref}} n)/N\) for \(n = 1, 4, 9\) and see very good agreement with our numerical data. Discrepancy between the CFT prediction and the data for \(S_{\text{tot}}^z = 3\) for \(N^{-1} > 0.05\) can be because of low-lying states with a finite gap (dashed line) which is lower than the first bulk septuplet excitation for system sizes smaller than \(N \approx 20\).

To summarize, by introducing an additional next-nearest-neighbor frustration along the backbone we were able to drive the comb into the dimerized state on the backbone passing through the Kosterlitz-Thouless critical point. For \(J_1 = J_{bb} = 1\) we locate the KT transition around \(J_{2bb} \approx 0.37\), which is significantly higher than in the simple spin-1/2 chain with \(J_1 - J_2\) interaction where the transition takes place at \(J_2/J_1 \approx 0.2411\). This suggest that partial localization of the edge states on the second sites of the teeth induce an effective ferromagnetic coupling between the next-nearest-neighbor teeth. This is confirmed by our simple calculation for \(J_{bb} = 0.1\) for which the KT transition is located approximately at...
$J_{3ab} \approx 0.25$, much closer to the KT transition of the original spin-1/2 chain.

E. Higher-order edge states

We return to our original comb model with all spins-1, including at the tooth ends. In a comb with an odd number of sites per teeth, the edge states of each tooth form a triplet as the ground-state. Therefore each tooth can be viewed as a composite spin-1 object. By tuning the interaction along the backbone one also tunes the correlation between the edge spin-1/2 at the end of the teeth, although the effective interaction between them vanishes exponentially fast with the length of the teeth. It is very natural to think here in terms of the VBS singlets: each tooth with a pair of spin-1/2 edge states corresponds to a composite spin-1 object. The ground-state is then given by the Haldane state, where each spin-1/2 is connected by a VBS to one of its neighbor. In a simple spin-1 chain two spins-1/2 inside a spin-1 object are perfectly symmetrized, so the Haldane state is not affected by the specific arrangements of a VBS singlets, as soon as each nearest-neighbor bond contains one and only one VBS singlet. The situation is a bit different on a comb, since the interaction between the edge spins-1/2 on a backbone is much stronger than the interaction between the edge spins at the end of each tooth. As we know, open boundary condition in the spin-1/2 chain favors dimerization. Most of the interactions between the composite $S = 1$'s of an effective Haldane chain is due to the backbone. So the total energy of the comb is minimized when the first VBS singlets of the composite Haldane chain is located at the first bond of the backbone. The next VBS singlets is then placed between the edge spins-1/2 on the backbone and the teeth 3 and 4 etc. This state is sketched in Fig. 15.

Within the described VBS picture it is clear that the end spins of the first and last teeth remain unpaired and form the edge states of an entire comb. On a finite-size system, these two spins-1/2 can couple to each other and form triplet and singlet in-gap states with an extremely small energy splitting. To confirm this picture we extract the local magnetization of a comb with 20 teeth and 7 spins per tooth in the sector of total magnetization $S_z^{tot} = 1$. The edge states of the Haldane state on each tooth form a triplet, so the entire comb is equivalent to the spin-1 chain, which is also in the Haldane phase with one valence bond singlet between the pair of neighboring tooth. The edge states emergent in this Haldane chain are localized at the end of the first and the last teeth - the two furthest points of a comb. On two neighboring teeth the spins equally distant from the backbone are anti-parallel. The maximal polarization is 0.15.

V. CONCLUSION

In the present paper we describe an alternative type of tree tensor network applicable to one-dimensional models.
with complicated on-site clusters. This numerical set-up partially fills the gap between one-dimensional DMRG and two-dimensional PEPS (PEPS) tensor network algorithms. While DMRG has established itself as the most efficient numerical tool for one-dimensional system of strongly interacting particles, in more complicated cases, such as a chain of clusters each with substantial entanglement, the intra and inter-cluster entanglement are both sent through one auxiliary bond, so the total bond dimension required may be large. Higher dimensional tensor networks, such as, PEPS, can distribute entanglement through many links to satisfy the area law naturally, but they involve complicated algorithms with high complexity in the bond dimension. The comb tensor network is a useful compromise, with much of the computational efficiency of DMRG but with the ability to separate intra and inter cluster entanglement. Of course, for true two- or higher-dimensional lattices, the bond dimension within the tooth would grow exponentially, and the comb is not likely to be useful.

The comb algorithms presented here have great flexibility, including allowing local changes to the Hamiltonian or disorder, alteration or variability in the local degrees of freedom (e.g., spin value), beyond nearest-neighbor interactions, the length of the teeth, etc. Along any tooth one has the full flexibility of an MPS along with the favorable complexity of an MPS, allowing, for example, each tooth to represent a finite higher dimensional or randomly connected cluster.

Of course, the most natural system for a comb tensor network is a system with a comb geometry, or one whose strong local interactions have the topology of a comb. In the present paper we discussed spin-1 Heisenberg model on a comb lattice. The teeth in this system have a bulk gap and are in a Haldane phase, which is topological and has emergent edge states. The spin-1/2 edge states of the teeth are coupled by the backbone interaction and form a critical chain. This provides a simple recipe for how half-integer-spin criticality can be realized with integer spins only. In a spin-$S = 1/2$ comb, the teeth themselves are critical. The tuning of the backbone interaction also tunes effectively the dimensionality of the entire system. We will present results for this intriguing system elsewhere\cite{33}.

VI. ACKNOWLEDGMENTS

This work has been supported by the Swiss National Science Foundation and the U.S. National Science Foundation through DMR-1812558, and the Simons Foundation through the Many-Electron Collaboration.
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