Stripes in the two-dimensional $t$-$J$ model with infinite projected entangled-pair states

Philippe Corboz,1,2 Steven R. White,3 Guifrè Vidal,4 and Matthias Troyer1

1Theoretische Physik, ETH Zurich, 8093 Zurich, Switzerland
2Institut de théorie des phénomènes physiques, École Polytechnique Fédérale de Lausanne, CH-1015 Lausanne, Switzerland
3Department of Physics and Astronomy, University of California, Irvine, CA 92697-4575 USA
4School of Mathematics and Physics, The University of Queensland, QLD 4072, Australia

(Dated: January 20, 2013)

We simulate the $t$-$J$ model in two dimensions by means of infinite projected entangled-pair states (iPEPS) generalized to arbitrary unit cells, finding results similar to those previously obtained by the density-matrix renormalization group (DMRG) for wide ladders. In particular, we show that states exhibiting stripes, i.e., a unidirectional modulation of hole-density and antiferromagnetic order with a $\pi$-phase shift between adjacent stripes, have a lower variational energy than uniform phases predicted by variational and fixed-node Monte Carlo simulations. For a fixed unit-cell size the energy per hole is minimized for a hole density $\rho_l \sim 0.5$ per unit length of a stripe. The superconducting order parameter is maximal around $\rho_l \sim 0.75 - 0.8$.

PACS numbers: 02.70.-c, 71.10.Fd, 71.10.Hf

I. INTRODUCTION

The simulation of strongly correlated electron systems in two dimensions remains one of the biggest challenges in condensed matter physics. The infamous negative sign problem3 prevents accurate quantum Monte Carlo simulations of large fermionic systems at low temperature. An alternative route without a sign problem is provided by variational Monte Carlo (VMC). Typically, several ansatz wave functions, biased towards different orders, are optimized and the one with lowest energy is considered as the best approximation to the ground state. A powerful way to improve upon a variational wave function is to use it as a guiding wave function for the fixed-node Monte Carlo (FNMC) method2 which yields the best variational wave function with the same nodal structure as the guiding wave function.

In recent years, a new class of variational wave functions for two-dimensional systems have been proposed: the so-called tensor network states, including e.g. the multi-scale entanglement renormalization ansatz (MERA)4 and projected entangled-pair states (PEPS)5 also called tensor product states (TPS). These tensor networks have recently been generalized to fermionic systems4 and can be seen as generalizations of the density-matrix renormalization group (DMRG)5 method to two dimensions. In contrast to other variational wave functions, tensor network states are largely unbiased11 with an accuracy which can be systematically controlled by the so-called bond dimension $D$ (typically called $m$ in DMRG). DMRG yields very accurate results for quasi one-dimensional ladder systems with cylindrical boundary conditions up to a width around $8 - 12$, but becomes numerically inefficient for larger widths. The computational cost of MERA and PEPS is polynomial in system size and in $D$, however, with such a high power in $D$ that one may question if these methods are competitive to solve hard problems in condensed matter physics.

In this paper we show that tensor networks indeed can compete with the best known variational methods, in particular with FNMC based on Gutzwiller projected ansatz wave functions. Specifically, we simulate the doped $t$-$J$ model for $J/t = 0.4$ in the thermodynamic limit with infinite PEPS (iPEPS5) and with DMRG for systems up to a width $10$, and find significantly lower variational energies than obtained with FNMC. It was previously shown that iPEPS yields lower energies than VMC11,12 but the values were still higher than the ones from FNMC. Here we find that extending the ansatz to larger unit cells than the $2 \times 2$ cells previously used leads to a considerable improvement of the variational energy, which is an indication that the ground state may break translational invariance on a larger scale than $2 \times 2$. By inspection of local order parameters we find that the ground state of the two-dimensional $t$-$J$ model exhibits stripes, i.e., a unidirectional modulation of the hole density and the antiferromagnetic order, as previously found with DMRG in cylinders up to width $8$. This is in contrast to the findings from VMC11,12 and FNMC5 which have suggested a uniform phase.

II. METHOD

Fermionic iPEPS has been introduced and explained in detail in Ref. [1] and we here repeat only the basic ideas. A PEPS, illustrated in Fig. [1a], can be seen as an extension of a matrix-product state (MPS) in Fig. [1b]), the tensor network DMRG is based on, to two dimensions. Each blue-filled circle represents a tensor with a rank given by the number of legs attached to it, where the open leg corresponds to a physical index carrying the local Hilbert space of a lattice site, and the connecting lines are bond indices with a certain bond dimension $m$ or $D$, which characterizes the number of variational parameters in the ansatz. Tracing over all bond indices yields the coefficients of the state in the tensor product basis of the local Hilbert spaces of all sites. MPSs and PEPSs en-
able an efficient representation of states obeying the area law of the entanglement entropy in one and two dimensions, respectively. An MPS can also be used to represent states in two dimensions, e.g., by using a snake structure as in Fig. 1(d), however, the required bond dimension $m$ of the tensors grows exponentially with the system’s width, whereas in a PEPS the required $D$ is independent of system size (in the limit of large systems). In the present work we consider a PEPS of infinite size (iPEPS) made of a periodically repeated rectangular unit cell containing $L_x \times L_y = N$ different tensors, $A^{[x,y]}$, labeled by the coordinates relative to the unit cell. To obtain an approximate representation of the ground state we perform an imaginary time evolution of an initial, randomly chosen, iPEPS. To efficiently compute observables an approximate contraction scheme, discussed below, is used to evaluate the trace over all bond indices, where the accuracy can be controlled by another parameter called boundary dimension $\chi$.

For the experts, we briefly outline how to treat large unit cells, where we adopt the notation from Refs. 11, 12. First, for the imaginary time evolution we use the simple update on all bonds in the unit cell. Second, to contract the iPEPS we use the corner-transfer matrix (CTM) method, generalized to larger unit cells. The CTM method yields the so-called environment, consisting of corner tensors $C_1, C_2, C_3, C_4$ and edge tensors $T_1, T_2, T_3, T_4$, which account for the infinite system surrounding the tensors in the “bulk” of the system (Fig. 1). We again assign coordinates $[x, y]$ to each of these tensors to label the relative position in the unit cell, i.e. 4N corner tensors and 4N edge tensors in total are separately stored. Initially, the corner and edge tensors at position $[x, y]$ are constructed similarly as a reduced tensor $a^{[x,y]}$, by multiplying the tensor $A^{[x,y]}$ to its conjugate and fusing the bond indices where we trace over the legs directed toward an open boundary, as illustrated in Fig. 1. The environment tensors are iteratively built by four directional coarse-graining moves (left, right, top, bottom), similarly as proposed in Ref. 21.

An entire CTM step consists of $L_x$ left moves, $L_y$ right moves, $L_y$ top moves and finally $L_y$ bottom moves. This sequence is repeated until convergence is reached. The renormalization procedure, adopted from Refs. 12, 20, is based on a $4 \times 4$ cell of tensors to compute an appropriate isometry. These isometries are used to absorb a column (or row) of tensors into the corresponding boundary tensors, which effectively corresponds to the growth of the system by one column (or row) of sites. The left move is explained in Fig. 1 and one proceeds similarly for the other moves.

III. SIMULATION RESULTS

With the generalized iPEPS introduced in the last section we simulate the $t$-$J$ model, given by the Hamiltonian

\[
H = -t \sum_{\langle ij \rangle} \langle c_{i\sigma}^\dagger c_{j\sigma}^\vphantom{\dagger} \rangle \delta_{\sigma,\varepsilon} + J \sum_{\langle ij \rangle} \langle S_i S_j \rangle - \frac{1}{4} \hat{n}_i \hat{n}_j
\]

with $\sigma = \{\uparrow, \downarrow\}$ the spin index, $\hat{n}_i = \sum_\sigma \hat{c}_{i\sigma}^\dagger \hat{c}_{i\sigma}$ the electron density and $\hat{S}_i$ the spin 1/2 operator on site $i$, and $\hat{c}_{i\sigma} = \hat{c}_{i\uparrow} (1 - \hat{c}_{i\downarrow} \hat{c}_{i\uparrow})$, where we fix $J/t = 0.4$.

Our main result for the infinite two-dimensional $t$-$J$ model is that, in contrast to the uniform phase found by VMC and FNMC we find a striped phase in the doping regime $1/2 < \delta < 1/4$, in agreement with DMRG calculations on wide ladders. Figure 2 shows that the

![Diagram of a PEPS with bond dimension D on a 4 x 4 lattice.](image)

FIG. 1: (Color online) a) Diagrammatic representation of a PEPS with bond dimension $D$ on a 4 x 4 lattice. b) An MPS with bond dimension $m$ to represent a 4 x 4 system. c-e) Diagrams for the left move of the corner-transfer matrix method: (c) A plaquette of four reduced tensors a embedded in the environment. The coordinates $[x, y]$ are to be understood relative to the unit cell, i.e., one has to take the coordinate $x$ (y) modulo $N_x$ (modulo $N_y$). Cutting through the lines as marked in the figure yields the tensor $Q_{[x,y+1]}^{[x,y+1]}$ in panel (d): a singular value decomposition of tensor $Q_{[x,y+1]}^{[x,y+1]}$ is performed, where only the $\chi$ largest singular values are kept. The resulting isometry $U_{[x,y+1]}^{[x,y+1]}$ and its conjugate are used as an approximate resolution of the identity, $I \approx U_{[x,y+1]}^{[x,y+1]} U_{[x,y+1]}^{[x,y+1]}$. For a fixed $x = x_0$ one computes all isometries $U_{[x_0,y]}^{[x_0,y]}$ for all $y \in [1, N_y]$. These isometries are then used to obtain the renormalized corner tensors $C_1^{[x_0+1,y]}$ and $C_4^{[x_0+1,y]}$, and edge tensors $T_4^{[x+1,y]}$ for all $y \in [1, N_y]$ and fixed $x = x_0$, shown in e). This whole procedure is repeated $N_x$ times for $x_0 \in [1, N_x]$ to complete an entire left move. (f) Initialization of a boundary tensor from a PEPS tensor and its conjugate, where crossings have been replaced by swap tensors (cf. Ref. 11).
variational energies obtained with iPEPS and DMRG are considerably lower than the ones from VMC and FNMC. We consider here the energy per hole (in units of $t$), $E_{\text{hole}} = (E_s - E_0) / \delta$, with $E_s$ the energy per site and $E_0 = -0.467775$ the value at zero doping taken from Ref. 22. The VMC and FNMC energies are seen to increase slightly with system size, and thus, we expect that in the thermodynamic limit the striped states are energetically favored.

In Fig. 3 we present a convergence study of the iPEPS energies as a function of the dimensions $D$ and $\chi$, and compare it to DMRG results for ladders for a doping $\delta \sim 1/8$. While iPEPS is a variational ansatz the resulting energies may be non-variational, because of the approximate contraction of the iPEPS, which introduces an error depending on $\chi$. The convergence study in the right panel in Fig. 3 indicates that this error is smaller than the symbol size, i.e. the upper end of the symbol provides an upper bound of the ground state energy. The middle panel shows the values, extrapolated in $\chi$, as a function of $1/D$. The values have not yet converged, thus the ansatz can still be further improved by increasing $D$. Attempting a quadratic fit yields $E_{\text{hole}} = -1.59(3)$ as $D \to \infty$, which is similar to the extrapolated DMRG results, $E_{\text{hole}} = -1.61(2)$.

Examples of stripes obtained with iPEPS are presented in Fig. 4. These stripes appear without biasing the initial iPEPS to stripe order (we typically start from several random initial states). We tested different unit cell sizes with $L_y \in \{2, 4\}$ and $L_x \in \{2, 4, 5, 6, 7, 8, 10, 12\}$, where each unit cell imposes a different periodicity on the wave function. Some of the $L_y = 4$ samples exhibit a slight modulation also in $y$ direction, however, since the energy is similar to the $L_y = 2$ samples we focus on the latter.

Figure 2 shows that, for a fixed unit cell size, the energy minimum is found near $\rho_l \sim 0.5$ holes per unit length per stripe, e.g. at $\delta = 1/8$ for the $8 \times 2$ unit cell, which is in agreement with DMRG results (black stars and Ref. 16). However, the minimum for the $8 \times 2$ cell at $\delta \sim 1/8$ is higher than the energy of the $10 \times 2$ or the $12 \times 2$
sample at the same doping, which indicates that the repulsion between the stripes of width 4 in the 8 × 2 cell is too strong so that stripes with larger widths and larger \( \rho_l \) are energetically favored, in contrast to the predictions by DMRG \( \text{[10]} \). For densities \( 0.75 < \rho_l < 1 \) we observe the tendency that mixed stripes (two stripes with different densities) yield a lower variational energy than two stripes of equal density, however, we do not observe signs of phase separation between stripes with \( \rho_l = 1 \) and \( \rho_l = 0.5 \) as found with DMRG \( \text{[10]} \). A future study with larger unit cells and larger \( D \) will shed further light in this issue.

An open question is what happens at smaller doping than \( 1/12 \). The minimum at \( \rho_l \) for different unit cells is seen to decrease with increasing unit cell length \( L_x \). It is conceivable that this trend continues, so that we will see decreasing doping we obtain stripes which are increasingly more widely spaced\( \text{[10]} \), with a distance between the stripes varying as \( d \approx 1/(2\Delta) \text{[10]} \). However, from the present data we cannot rule out other phases, such as a uniform phase at small doping \( \delta < 1/12 \), or phase separation between an undoped and a doped region.

Finally, we study the pair field \( \Delta = 1/\sqrt{2}(\hat{c}_i \hat{c}_j - \hat{c}_i^\dagger \hat{c}_j^\dagger) \) between nearest-neighbor sites \( (i, j) \) which is modulated along the \( x \)-direction and predominantly forms a \( \delta \)-wave pattern as shown in Fig. 4. The \( \delta \)-wave order between neighboring stripes has the same phase, however, we have also observed states with a similar energy where the \( \delta \)-wave order exhibits a \( \pi \)-phase shift between neighboring stripes. Thus, it seems that this \( \pi \)-phase shift has only little influence on the energy, and from the present data we cannot determine which state is preferred. The mean pairing amplitude (averaged over the unit cell) in Fig. 5 exhibits a maximum around \( \rho_l = 0.75 \) (\( \rho_l = 0.8 \) in DMRG). For \( \rho_l = 1 \) the stripes are insulating with a vanishing pairing amplitude. For \( \rho_l = 0.5 \) the pairing amplitude for \( D = 8 \) is finite, but decreases quickly with increasing \( D \), and is possibly insulating for larger \( D \), too. In between these two insulating states, excess holes (or electrons) form pairs, leading to an increase of the pairing.

\[ \text{IV. CONCLUSION} \]

In this paper we presented iPEPS simulation results for the \( t-J \) model in the thermodynamic limit, where we obtained striped states, in qualitative agreement with DMRG calculations on wide ladders. The iPEPS variational energies are considerably lower than the ones from the uniform states obtained with VMC and FNMC based on Gutzwiller projected ansatz wave functions, and compatible with DMRG results for finite systems. This demonstrates that iPEPS is a competitive variational method for strongly correlated electron systems\( \text{[24]} \).

The differences between iPEPS and DMRG only involve quantitative details, such as the precise stripe linear filling and spacing as a function of overall filling. Using larger unit cells in iPEPS than the usual \( 2 \times 2 \) is essential to study ground states which break translational invariance on a larger scale than only 2 lattice sites.

The iPEPS wave functions for the accessible values of \( D \) include unbiased quantum fluctuations over short distance and high energy scales. Over longer length and lower energy scales, they revert to a more mean-field-like description. Thus, with the currently accessible \( D' \text{[25]} \) we cannot expect them to resolve between, e.g., static versus fluctuating stripes. On the other hand, our results provide significant evidence that approaches that ignore stripes do not give reliable descriptions of the \( t-J \) model.

We acknowledge inspiring discussions with F. Becca, T. Nishino, and H.-Q. Zhou and funding from the Swiss National Science Foundation and from the NSF under DMR-0907500. The simulations were performed on the Brutus cluster at ETH Zurich.

\[ \begin{align*}
1 & \quad \text{M. Troyer and U.-J. Wiese, Phys. Rev. Lett. 94, 170201 (2005).} \\
2 & \quad \text{D. F. B. ten Haa, H. J. M. van Bemmel, J. M. J. van Leeuwen, W. van Saarloos, and D. M. Ceperley, Phys. Rev. B 51, 13039 (1995).} \\
3 & \quad \text{M. Lugus, L. Spanu, F. Becca, and S. Sorella, Phys. Rev. B 74, 165122 (2006).} \\
4 & \quad \text{G. Vidal, Phys. Rev. Lett. 99, 220405 (2007); G. Vidal, Phys. Rev. Lett. 101, 110501 (2008); L. Cincio, J. Dziar-} \\
\]
Since we consider systems of infinite size, symmetries may be spontaneously broken.

Although charge- and spin-density waves have already been found in iPEPS simulations with $D = 6$, stripes could not be represented due to a too small unit cell.

A larger bond dimension $D$ can be achieved by exploiting symmetries of the model (we only used parity conservation in the present simulations). For a fixed $D$ the results can also be improved by going beyond the simple update used in the present work (cf. Ref. [21]).

Competitive variational energies (compared with VMC) have recently also been obtained with the 2D MERA for the Heisenberg model on the kagome lattice.

A larger bond dimension $D$ can be achieved by exploiting symmetries of the model (we only used parity conservation in the present simulations). For a fixed $D$ the results can also be improved by going beyond the simple update used in the present work (cf. Ref. [21]).