Capturing long range correlations in two-dimensional quantum lattice systems using correlator product states

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We study the suitability of correlator product states for describing ground-state properties of two-dimensional spin models. Our ansatz for the many-body wave function takes the form of either plaquette or bond correlator product states and the energy is optimized by varying the correlators using Monte Carlo minimization. For the Ising model we find that plaquette correlators are best for estimating the energy while bond correlators capture the expected long-range correlations and critical behavior of the system more faithfully. For the antiferromagnetic Heisenberg model, however, plaquettes outperform bond correlators at describing both local and long-range correlations because of the substantially larger number of local parameters they contain. These observations have quantitative implications for the application of correlator product states to other more complex systems, and they give important heuristic insights: in particular the necessity of carefully tailoring the choice of correlators to the system considered, and its interactions and symmetries.

I. INTRODUCTION

A. Background

Modeling strongly correlated systems exactly for more than a few particles is not possible due to the rapid increase in the size of the Hilbert space with their number. However, there has been much success recently in using tensor network methods to numerically simulate strongly correlated systems. These approaches allow states with suitable properties, such as obeying an area law for their entanglement entropy, to be efficiently represented by a network of tensors. They also provide efficient methods of ‘contracting’ the network to allow expectation values of operators, or more basically wave function amplitudes, to be calculated. In particular, matrix product states (MPS) can be used to accurately describe large systems in one dimension and can be exactly and efficiently contracted. Crucially this means that expectation values can be calculated in a number of steps that only grows as a low degree polynomial with system size and tensor size. This has been crucial for the success of MPS based algorithms like the density-matrix renormalization group and time-evolving block decimation. Tensor networks can be extended to higher dimensions using the projected entangled pair states (PEPS) construction but unlike MPS these do not permit efficient exact contraction. Approximate contraction procedures in two dimensions, such as those based on MPS methods or tensor renormalisation group (TRG), must be employed at the cost of having a much less favorable, though polynomial, scaling. Recently a procedure for performing PEPS simulations combining TRG and Monte Carlo sampling has been proposed, and it shows promising results for treating large bond dimensions. In contrast to the PEPS construction, both the multiscale entanglement renormalisation ansatz (MERA) and tensor tree networks (TTNs) utilize a hierarchical structure where the tensors are interconnected by bonds according to a tree pattern, and can be contracted exactly and efficiently. Like PEPS, these methods scale as a high degree polynomial.

An alternative less computationally expensive approach to represent two-dimensional systems is to consider classes of states that while not being efficiently contractible exactly, are efficiently and exactly samplable, i.e. for any given configuration the amplitude can be found exactly. The tensor network formalism provides a powerful framework from which to devise new and physically tailored ansätze with these properties for which the closest approximate ground state can be found using variational Monte Carlo methods. For example, string bond states use a product of overlapping MPS ‘strings’ and acquire their exact samplability from the contractibility of the underlying MPS. An alternative ansatz is formed by building the wave function using superpositions of all possible coverings of singlets i.e. resonating valence bond (RVB) states, and it has been shown that RVB states have a PEPS description. Recently a new ansatz within this class has been proposed for simulating lattice systems: so-called correlator product states or entangled plaquette states, which are equivalent and are hereafter referred to as CPS. A similar approach was also proposed in some earlier works. Correlator product states can be thought of as a more basic form of tensor network states, where correlations between sites are encoded explicitly in “correlator” building blocks so that the amplitude for each configuration is given by a product of their scalar elements. They provide a slightly simpler ansatz than string bond states, but with similar power and properties.
B. Motivation

The CPS construction has been applied to a variety of models, and in all cases the energies found compared well with those found using other methods, e.g. PEPS, MPS, stochastic series expansion (SSE), but with a much reduced computational cost. As we will discuss in detail below, the CPS description is incredibly versatile, and the correlator type can be varied without adversely affecting the complexity of the calculation. Thus, the type of correlator can be chosen to best match the properties or symmetries of the system. It is also possible to go beyond PEPS, which are composed of short ranged bonds, and consider CPS that contain long ranged bonds and that are area-law violating.

In previous work, the estimated ground state energy as a function of the plaquette size has been determined. What has so far been lacking is a systematic study of the relative behavior of different correlator types for describing the important physical properties of different systems. In particular bond correlators, where correlators are arranged to connect pairs of sites across the lattice, have not been studied in any detail. Bond correlators allow the possibility to directly encode long-range correlations between distant sites using a very small number of parameters. This is not possible with plaquette correlators, due to the exponential increase in the number of correlator elements with plaquette size, or with matrix product states, where distant correlations are mediated by intermediate nearest neighbor bonds. However, plaquette correlators do provide a more straightforward ansatz that imposes a less rigid short ranged structure. It is thus interesting to investigate what effect the choice of correlator product ansatz has on the ability to describe systems that possess long-range correlations, as well as the difficulties they may present during minimization.

The CPS construction seems promising for describing a wide variety of spin models, and modeling these systems could provide insights for cold-atom simulations of quantum magnetism. In previous works it has also been noted that certain important states such as the Laughlin wave function which cannot be efficiently described with other types of tensor network states, can have an exact CPS description when bond correlators between all sites are included. This illustrates that bond correlators can efficiently describe complex topological systems, and may be able to describe other fractional quantum Hall states. To aid future studies which will apply the CPS ansatz to more complicated systems, in this work we test their effectiveness using systems whose behavior is well-known. We examine in detail the performance of correlator types at describing the long range order and antiferromagnetism by applying them to the quantum transverse Ising model (TIM) on a square lattice. We find that bond correlators are better able to predict the critical point and represent expected long range correlations than plaquette correlators for the two-dimensional TIM, where bond correlators allow a larger range of sites to be covered for a given number of parameters and computational effort. In section IV we extend the treatment to different lattice geometries, finding similar behavior for the performance of the two correlator types. In section V we investigate the ability of plaquette and bond correlators to describe a more complex type of long range order and antiferromagnetism by applying them to the antiferromagnetic Heisenberg model (AFHM). We find that plaquette correlators are more successful at describing the AFHM, since they provide a much larger number of local parameters to better capture the complicated local structure encoding the antiferromagnetic order in this system. Finally, we conclude and summarize the results in section VI.

II. Method

The exact and efficient samplability of CPS makes them ideal for variational minimization (for details of the exact-efficient samplability of CPS see appendix A). Previously CPS have been minimized using a generalized eigenvalue method and a deterministic method. Here we use a Monte Carlo based stochastic minimization method to determine the ground state, which has previously been successfully applied to one-dimensional systems represented by matrix product states. The correlator elements that best approximate the ground state are found by estimating the derivative of the energy with respect to each correlator element, and then updating each correlator element according to the direction of the derivative with a random step size. This allows minimization using only the first derivative of the energy (so requiring fewer computation steps), and ameliorates the error associated with updating the correlator elements.
numbers. Unlike with PEPS, the amplitude the wave function amplitude is a simple product of number of sites. One advantage of using CPS is that represents four sites in a plaquette, or a string of a given any form of correlator, for example each correlator could correlators, for a $2 \times 2$ system.

We illustrate this in Fig. 1, for the case where the sub-

cartation is given by \( \{ s \} \) where we term \( s = (s_1, s_2, \cdots) \) the configuration of the state. The many-body wave function is \( |\psi\rangle = \sum_s W(s)|s\rangle \), where \( W(s) \) is the amplitude or weight of a given configuration \( s \). In this work we will for simplicity consider only spin-$\frac{1}{2}$ systems, but the methods used can be applied straightforwardly to lattice systems possessing a larger on-site dimension.

In the correlator product state description, the weight \( W(s) \) is given by the product of correlator elements \( C_s \) over the lattice

\[
W(s) = \prod_{\langle i \rangle} C_{s(i)},
\]

where each correlator element is a $c$-number that describes the amplitude of a configuration \( s(i) \) of a subgroup of sites \( \{ i \} \). The wave function in the CPS representation is given by

\[
|\psi\rangle = \sum_{\{ s \}} \prod_{\{ i \}} C_{s(i)} |s_1 \cdots s_L\rangle.
\]

We illustrate this in Fig. 1 for the case where the subgroup of sites is a nearest neighbor pair i.e. ‘bond’ correlators, for a $2 \times 2$ system.

This compact description of a state can be extended to any form of correlator, for example each correlator could represent four sites in a plaquette, or a string of a given number of sites. One advantage of using CPS is that the wave function amplitude is a simple product of $c$-numbers. Unlike with PEPS, the amplitude \( W(s) \) can be calculated efficiently and exactly for any configuration \( s \). This allows the wave function to be efficiently sampled, so that Monte Carlo methods can be used to determine expectation values and minimize the wave function. The flexibility and exact-efficient sampleability of CPS is described further in appendix A.

**B. Correlator types**

In the following, we study the ground states using different types of correlator, and compare their properties. We use periodic boundary conditions with the translational invariance of the system allowing one correlator of each type to describe the entire system.

The types of correlators used for square lattices are illustrated in Fig. 2. Nearest-neighbor (n. n.) bond correlators, with one correlator for the vertical bonds and one correlator for the horizontal bonds as shown in Fig. 2(a), are the simplest correlator types used. This gives a description of the ground state using only eight parameters. We also use bond correlators with longer range bonds. In general we term a bond correlator of size \( r_{\text{max}} \) as including all bond vectors up to the maximum range \( \Delta r_{\text{max}} = (r_{\text{max}}, r_{\text{max}}) \) i.e. it includes all the diagonal bonds as well as bonds along the lattice axes. Figures 2(b) and (c) show the set-up with correlators of size \( r_{\text{max}} = 1 \) and \( r_{\text{max}} = 2 \) respectively, and in the following

FIG. 1. (Color online) Illustration of the CPS ansatz for a state \( |\psi\rangle \) describing a simple $2 \times 2$ system with open boundary conditions. The local states \( s_i \) are those of a spin-$\frac{1}{2}$ system. Each correlator \( C_s \) represents the amplitudes for different configurations of spins on two sites \( i, j \). The four-site system has a total of sixteen basis states, with the weight for an example basis state \( W(\uparrow \downarrow \uparrow \uparrow) = C_{\uparrow \downarrow \uparrow \uparrow} \).

FIG. 2. (Color online) Illustration of the different correlator types used. The blue circles represent lattice sites, and only correlators that link these $3 \times 3$ sites are represented. (a) Nearest neighbor (n. n.) bond correlators. (b) Bond correlators with \( r_{\text{max}} = 1 \). (c) Bond correlators with \( r_{\text{max}} = 2 \). (d) $2 \times 2$ plaquette correlators.
calculations we use correlators up to \( r_{\text{max}} = 8 \). For correlators of size \( r_{\text{max}} \), the number of correlator elements is \( 8r_{\text{max}}(r_{\text{max}} + 1) \). To calculate the energy, the number of computational steps scales as \( O(N^2 r_{\text{max}}^2) \), and the only part of the calculation that depends on the correlator size and type is the calculation the correlator fraction given in equation (B2).

We also use plaquette correlators, with the smallest \( 2 \times 2 \) plaquette illustrated in Fig. 2(d), up to a size of \( 4 \times 4 \) plaquettes. The correlators are set up so that they are displaced from one another by one site and overlap one another. In general, the greater the overlap between the correlators the more accurate the description of the ground state. The number of elements in an \( n \times n \) correlator scales as \( 2^{n^2} \), so the memory requirements for a given plaquette correlator size scale much faster than for bond correlators. However the number of computational steps required to calculate the energy is \( O(N^2 n^4) \), which is a mild polynomial scaling and grows much more slowly than the number of correlator elements. The reason for this is that calculating an off-diagonal expectation value only involves picking out the correct correlator element for the subgroup of sites spanned by a correlator. As a result it depends on the number of sites spanned by the correlator and the number of correlators that fall on a given site, and not on the number of elements. Note that calculation of the energy derivative formally requires a number of steps that does scale with the number of correlator elements, however in practice this part of the calculation is fast compared to estimating the energy itself and it is not rate limiting. Thus plaquette correlators allow the use of many more parameters for describing the system with only a modest increase in computational effort, although in our calculations their size is ultimately limited to \( 4 \times 4 \) by memory requirements.

Comparing the computation time for correlator types, a calculation using \( 3 \times 3 \) plaquette correlators has approximately the same number of computation steps as for bond correlators up to \( r_{\text{max}} = 4 \), and a calculation using \( 4 \times 4 \) plaquette correlators has approximately the same number of computation steps as for bond correlators up to \( r_{\text{max}} = 8 \). However, when comparing the performance of different correlator types, it is worth considering the three main differences between them: (i) The number of sites that can be reached from a given site using the correlator: \( r_{\text{max}} = 1 \) is equivalent to a \( 2 \times 2 \) plaquette, \( r_{\text{max}} = 2 \) is equivalent to a \( 3 \times 3 \) plaquette, and \( r_{\text{max}} = 3 \) is equivalent to a \( 4 \times 4 \) plaquette. (ii) The computational effort: for a given number of spanned sites, our routines for bond correlators are 2, 3.5 and 5.5 times faster than for plaquette correlators respectively. (iii) The number of correlator elements: the number of correlator elements for a given number of sites spanned is far smaller for bond correlators. In addition to the reduction in computational effort, this makes it possible to span far more sites using bond correlators. The more fragmented structure of bond correlators also permits different minimization strategies as described in more detail in appendix B.

III. LONG RANGE CORRELATIONS AND CRITICAL BEHAVIOR: THE QUANTUM TRANSVERSE ISING MODEL ON A SQUARE LATTICE

As an ideal test case for examining different correlator types, we consider the TIM on an \( L \times L \) square lattice, described by the Hamiltonian

\[
H = -J \sum_{i,j=1}^{L} \left( \sigma_i^{[i,j]} \sigma_{i+1,j} + \sigma_i^{[i,j]} \sigma_{i,j+1} + g \sigma_i^{[i,j]} \right),
\]

where \( J > 0 \) is the coupling, \( i \) and \( j \) denote the lattice index in the two perpendicular directions, \( \sigma_i^{[i,j]} \) is the Pauli \( z(x, y) \) operator for spin \( (i, j) \), and \( g \) is the dimensionless transverse magnetic field. We consider this model because it is one of the archetypal systems that exhibits a quantum phase transition at a finite magnetic field. The system moves from an ordered ferromagnet in the \( z \) direction at \( g = 0 \) to a state disordered in the \( z \) direction and aligned in the \( x \) direction when \( g \gg 1 \). As the system approaches criticality at \( g = g_c \), the correlation length diverges, and the system becomes gapless, making a numerical description of the state challenging. The critical point in the thermodynamic limit, found using a finite-size scaling analysis, is \( g_c \approx 3.044 \), where the pseudo-critical point was defined using a careful extrapolation of the ratio of the energy gap between consecutive system sizes.

By applying the CPS ansatz to this system, we investigate its performance at describing the behavior of several important physical quantities as the transverse magnetic field is varied. We focus in particular at modeling the long-range correlations close to the pseudo-critical point, and compare with the many previous studies performed using other numerical and approximate analytical methods. This system is thus a highly useful benchmark of the method for demonstrating both the effectiveness and limitations of the CPS approach.

A. Energy

The energy and its derivative, as well as other observables, are calculated using the algorithm described in appendix B. Note that for all correlator types we restrict to only real parameters without any loss of generality, since for this model the ground state can be constructed using real, positive weights for all configurations. We first calculate the energy at a value of the magnetic field close to the critical point for two systems. The largest system that has been solved numerically exactly is a \( 6 \times 6 \) system and so provides a good comparison for how well the method is working. This system size has also been solved using TTNs so we also compare the accuracy of the CPS method to those results. The energy of the ground state found using different correlator types is shown in Table II. The error, defined as the
The difference between the energy estimated using CPS and the exact ground state energy, is also displayed. As expected, we find that the energy converges to the exact value for increasing correlator size. In comparison, for TTNs, around $10^7$ parameters are required to reduce the error to $2 \times 10^{-5}$ \cite{Note1}. We also apply the CPS method to much larger systems. In Table I, we show the minimized energy in a $31 \times 31$ system at $g = 3.05$, which is close to the pseudo-critical point.

We find that even though plaquette correlators have more elements, the energy convergence is more well-behaved than for bond correlators. In line with commonly known properties of non-convex optimization, it is likely that having a larger number of parameters is beneficial when there is a foliated energy landscape since it allows the optimization more freedom in finding the minimum energy state. Conversely constraining the number of parameters for such long range bonds tends to result in numerous local minima giving a more difficult minimization problem. For example in a previous work minimizing over a similar class of many-body states, it was found that the enforcement of certain symmetries increased the difficulty of finding a good state, since this amounted to cutting through the energy landscape of the parameter space, dividing it into separated minima\cite{Note1}. However, even though a larger span of sites is found to be needed to minimize to a given energy with bond correlators, the number of parameters required is still far smaller than for plaquettes substantially reducing the computational effort.

We also investigate how the choice of correlator affects the energy at different values of the transverse magnetic field $g$. Figure 3 shows the difference between the energy calculated using n. n. bond correlators (which give the highest estimated ground state energy) and the energy calculated using $4 \times 4$ plaquette correlators (which give the lowest estimated ground state energy), for different values of the transverse magnetic field $g$. There is not much difference between the two values for $|g - g_c| \gg 1$, but the energy difference increases when $g \approx g_c$. Far from criticality, when the correlations are expected to be short range, the energy can be calculated accurately using a small number of parameters, however as the correlation length increases a larger number of parameters are needed to describe the system accurately. The maximum energy difference is still small ($< 1\%$), however this can lead to large differences in the properties of the state due to the large number of low-lying excited states combined with a vanishing gap as $g_c$ is approached. This is a general problem with using a variational approach to describe critical systems. The investigation of these properties is described in the following sections, and we find that choosing an ansatz with a suitable structure helps to better describe some important physical prop-

<table>
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<th>Correlator type</th>
<th>No. of elements</th>
<th>Energy</th>
<th>Error</th>
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</thead>
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<td>-3.2348(4)</td>
<td>13 $\times 10^{-3}$</td>
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<td>2 $\times 10^{-3}$</td>
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<tr>
<td>$r^{\text{max}} = 3$</td>
<td>96</td>
<td>-3.2465(2)</td>
<td>1 $\times 10^{-3}$</td>
</tr>
<tr>
<td>$2 \times 2$ plaquettes</td>
<td>16</td>
<td>-3.2387(4)</td>
<td>9 $\times 10^{-3}$</td>
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<td>512</td>
<td>-3.2463(2)</td>
<td>1 $\times 10^{-3}$</td>
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<tr>
<td>$4 \times 4$ plaquettes</td>
<td>65,536</td>
<td>-3.24725(5)</td>
<td>2 $\times 10^{-5}$</td>
</tr>
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</table>

TABLE I. Energy per site in units of $J$ found using stochastic minimization for the two-dimensional TIM in a $6 \times 6$ system for different correlator types. Results are compared with exact results at $g = 3.05266$, which has been calculated to be the pseudo-critical point in a $6 \times 6$ system, with a ground state energy of $-3.2472744 \cdots$ (Ref. \textsuperscript{[55]}).

<table>
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<tr>
<td>$4 \times 4$ plaquettes</td>
<td>65,536</td>
<td>-3.23903(5)</td>
</tr>
</tbody>
</table>

TABLE II. Energy per site in units of $J$ found using stochastic minimization for the two-dimensional TIM in a $31 \times 31$ system for different correlator types at $g = 3.05$. 

FIG. 3. (Color online) Difference in energy between ground state calculated using nearest-neighbor bond correlators, and ground state calculated using $4 \times 4$ plaquette correlators. The dashed line indicates the critical magnetic field found by a finite-size scaling analysis (Ref. \textsuperscript{[55]}), and the dotted line indicates the peak of the energy difference at $g = 3.17$ — this is approximately midway between the pseudo-critical points found with nearest-neighbor bond and $4 \times 4$ plaquette correlators.
FIG. 4. (Color online) Absolute magnetization $\langle |\sigma_z| \rangle$ as a function of transverse magnetic field $g$ in an $L \times L$ system, for the ground state found using $2 \times 2$ plaquette correlators. The pseudo-critical point occurs at $g \approx 3.15$. Inset: $\langle |\sigma_z| \rangle$ at constant $g > g_c$ as a function of $L$, both axes having a logarithmic scale, with a fit of the form $\langle |\sigma_z| \rangle \propto L^{-b}$. The data was fitted with $b = 1.0$.

We investigate the position of the pseudo-critical point in the system by examining the local order parameters. Specifically, we calculate both the transverse magnetization $\langle \sigma_x \rangle$, and the absolute magnetization $\langle |\sigma_z| \rangle$, defined as the expectation of the absolute value of the average of all $L \times L$ spins, i.e. it quantifies how well the spins are aligned with one another (N.B. $\langle \sigma_z \rangle$ is zero due to the global $\mathbb{Z}_2$ symmetry of the system). Figure 4 shows the results for the absolute magnetization $\langle |\sigma_z| \rangle$ as a function of transverse magnetic field using $2 \times 2$ plaquette correlators for a number of $L \times L$ system sizes. The error bars for each point, where the error is given as the standard deviation of the $G$ different bins (see appendix [3]), are smaller than the marker for the data point.

As expected, we find a sharp drop in the magnetization at around $g = 3.05$ and the change in magnetization becomes steeper as the system size increases. We also find that the small non-zero magnetization for $g > g_c$ decreases as the system size increases: for $L = 51$, the magnetization at $g = 4$ is $\langle |\sigma_z| \rangle \approx 0.022$. The inset in Fig. 4 shows the magnetization as a function of lattice size for different fixed values of $g > g_c$, with a fit to a power law decay of the magnetization, i.e. $\langle |\sigma_z| \rangle_{g > g_c} = aL^{-b}$. The data fit an inverse scaling with system size, indicating that the magnetization decays to zero for $g > g_c$ in an infinite system. While the behavior found for $2 \times 2$ correlators is not quantitatively precise (the pseudo-critical point occurs at $g \approx 3.15$ rather than at $g = 3.044$), it is simple and offers a computationally affordable means of approximately locating a critical point once the appropriate local order parameters signifying it are known.

As described in section III A, we find that larger correlators are better able to minimize the energy. Figure 5 shows how the behavior of the local order parameter $\langle |\sigma_z| \rangle$ and the transverse magnetization $\langle \sigma_x \rangle$ depend on the correlator type and size. The position of $g_c$, the critical point found by a finite-size scaling analysis, is also indicated. The plots of the absolute magnetization $\langle |\sigma_z| \rangle$ shown in Figures 5(a) and (b) show that as the correlator size increases, the calculated pseudo-critical point moves closer to $g_c$ for both plaquette and bond correlators. The transverse magnetization (shown in Figures 5(c) and (d)) displays the expected ‘knee’ at the pseudo-critical point, which again moves closer to $g_c$ as the correlator size increases. The magnetization close to $g_c$ has not completely converged for the larger plaquette correlators in our calculations; the difference in $\langle |\sigma_z| \rangle$ for the $4 \times 4$ plaquette correlator and the $3 \times 3$ plaquette correlator is 0.003. Comparing this with the convergence found using TTNs, for a $10 \times 10$ system, the difference in the transverse magnetization for $\sim 10^6$ parameters and $\sim 10^7$ parameters is around 0.002. We see some scatter in the behavior of both the absolute and transverse magnetization calculated using bond correlators, owing to the more difficult
minimization, which means that the convergence cannot be accurately determined. However, despite this noise it is clear from our results that bond correlators display a pseudo-critical point much closer to that found in Ref. 35 for $r_{\text{max}} \geq 5$ as indicated by both the absolute and transverse magnetization. This suggests that bond correlators are capturing the critical behavior of the local order parameters in the TIM better. It is perhaps surprising that the energy density is better described by plaquette correlators, while the behavior of the local order parameter is qualitatively better with bond correlators. We next study the performance of plaquette and bond correlators at describing the long-range correlations, to provide some further insights.

C. Long-range correlations

The success of the MPS and PEPS-type tensor network approach is intimately connected with the decay of correlations in the system. When the correlations are short range, the tensor network representation is likely to be able to model the system well, with a number of parameters exponentially smaller than the Hilbert space size. When the long-range correlations approach polynomial decay, however, it can be difficult to model the system accurately using these methods, and a larger number of parameters are required to describe the entanglement in the system. Conversely, the success of MERA and TTNs is based on the logarithmic rather than linear scaling of the distance in the network between two points a given distance apart, and this ability to describe long range correlations can dramatically aid performance.

To investigate how the success of the CPS description is related to the ability to describe long range correlations we calculate the connected two point correlation function given by

$$C_{x,y} = \langle \sigma_z^{[0,0]} \sigma_z^{[x,y]} \rangle - \langle |\sigma_z^{[0,0]}| \rangle \langle |\sigma_z^{[x,y]}| \rangle,$$

for a wave function that has been minimized at different values of the transverse magnetic field and for different correlator types. The connected two point correlation function $C_{x,y}$ describes the probability of two spins on separated sites being aligned with one another, and can only be non-zero if entanglement exists between these two sites in the underlying ground state. When the system is close to the pseudo-critical point, it possesses its maximum correlation length $\xi$, and the system is the hardest to simulate numerically.

We calculate the average connected two-point correlation function in a $31 \times 31$ system at $g = 3.05$ found using (a) n. n. correlators and (b) bond correlators with $r_{\text{max}} = 8$. (c) Correlation length as a function of $g$ for different correlator types, bond correlators are open circles, plaquette correlators are closed circles. See Fig. 3 for full legends. The dashed line indicates the critical magnetic field found by a finite-size scaling analysis (Ref. 35).

FIG. 6. (Color online) The connected two-point correlation function plotted using a logarithmic color scale in a $31 \times 31$ system at $g = 3.05$ found using (a) n. n. correlators and (b) bond correlators with $r_{\text{max}} = 8$. (c) Correlation length as a function of $g$ for different correlator types, bond correlators are open circles, plaquette correlators are closed circles. See Fig. 3 for full legends. The dashed line indicates the critical magnetic field found by a finite-size scaling analysis (Ref. 35).
FIG. 7. (Color online) Correlators used in triangular and hexagonal lattice geometries. (a) Four-site plaquette correlators on a triangular lattice. (b) n. n. bond correlators on a triangular lattice. (c) \( r_{\text{max}} = 1 \) bond correlators on a triangular lattice. (d) Six-site plaquette correlators on a hexagonal lattice.

better at capturing the long-range correlations despite having a larger energy estimate than plaquettes. Note that it is not uncommon for a state with a higher energy to represent other properties of the ground state more accurately than that with the lower energy estimate. For this reason it is important that the choice of ansatz should try to reflect some expected underlying properties of the ground state and that the results, beyond just energy, need to be carefully examined (as in our work). The energy of a given state depends entirely on short-range correlations. Plaquette correlators have a large number of parameters to describe the short-range interactions, while the comparatively small number of parameters for bond correlators limit the accuracy to which it can describe the energy. However the structure of bond correlators allows an efficient description of any prevalent long-range correlations since it provides direct bonds between more distant lattice sites than can be reached with plaquette correlators. Although the calculated correlation length is still not more than two lattice sites close to the pseudo-critical point, it is for this reason that bond correlators are better able to describe the longer ranged properties of the Ising ground state, as found in this investigation.

IV. DIFFERENT GEOMETRIES: ISING MODEL ON TRIANGULAR AND HEXAGONAL LATTICES

To generalize the above conclusions to different lattice geometries, we also apply the CPS ansatz and minimization method to triangular and hexagonal lattices using the correlator types illustrated in Fig. 7. The behavior of the system is expected to be qualitatively the same for the square lattice, with the critical magnetic field shifted due to the different coordination numbers. For hexagonal and triangular lattices, finite-size scaling analysis predicts \( g_c = 2.13 \) and \( g_c = 4.77 \) respectively. We calculate the absolute magnetization as a function of transverse magnetic field for the smaller plaquette correlators, for different lattice sizes, and the results are shown in Fig. 8. We see the expected behavior, with the pseudo-critical point shifted to a higher magnetic field for small plaquette correlators, and find that the value of the absolute magnetization at \( g \gg g_c \) decreases for increasing lattice size \( L \).

As with the square lattice, we calculate energy close to the pseudo-critical point, and we find that plaquette correlators give the lowest energy. For example, for a triangular lattice with 400 sites at \( g = 4.77 \) the energy found using \( 4 \times 4 \) plaquettes is \(-4.9899(1)\) J, while with bonds having \( r_{\text{max}} = 5 \) the minimum energy is found to be \(-4.9895(2)\) J. Also as in square lattices we find that despite this, bond correlators seem to be better for determining the critical behavior of the system. Figure 9(a) shows the absolute magnetization as a function of transverse magnetic field for different correlator types. As with the square lattice, the longer range of the bond correlators predict a pseudo-critical point that is closer to the critical point (although once again more scatter is evident). The correlation length as a function of transverse magnetic field in shown in Fig. 9(b). Again, we find that the largest correlation length for bond correlators, and the position of the pseudo-critical point is consistent with that predicted by the local order parameter.

The numerical calculations indicate that the results found for the TIM are not dependent on the lattice geometry chosen. For all the geometries investigated it is found that the longer ranged bond correlators predict a
V. ANTIFERROMAGNETIC ORDER: THE HEISENBERG MODEL

To examine how plaquette and bond correlators compare when describing very different physics, we also investigate the AFHM on a square lattice. This system is more challenging to describe than the TIM, as it exhibits richer behavior owing to its symmetry and antiferromagnetism. The antiferromagnetic spin-$\frac{1}{2}$ Heisenberg model is described by the Hamiltonian

$$H_0 = J \sum_{i=1,j=1}^{L} S^{[i,j]} \cdot S^{[i+1,j]} + S^{[i,j]} \cdot S^{[i,j+1]},$$

where $S = \frac{1}{2}(\sigma_x, \sigma_y, \sigma_z)$ is the spin-$\frac{1}{2}$ operator. It corresponds to the half-filled band limit of the Hubbard model, and is assumed to describe the antiferromagnetic undoped insulator La$_2$CuO$_4$ as well as other undoped copper oxide materials.

For a pair of spins, the ground state of $H_0$ is simply a spin singlet. However, due to the monogamy of entanglement, each spin cannot be in a singlet state with more than one neighbor, and the ground state for $N$ spins has to satisfy both local minimization and global translational symmetries giving a highly complicated superposition state. As a result of this, it is predicted that long-range order arises. This is characterized by the two-point correlation function for the total spin $\langle S^z \rangle$ and the staggered magnetization, which is reduced from the Néel, or classical, value by quantum fluctuations.

The AFHM has also already been treated using plaquette type correlators, but here we extend that description by comparing them with bond correlators, and examine which correlator type better captures the expected long-range order. The results obtained here are also compared with those found using SSE, which give the current best estimate. This system has been treated using a resonating-valence bond (RVB) picture, by considering states that are superpositions of all possible coverings of singlets on the sites. The latest study gives estimates of the energy and staggered magnetization that agree well with those found using SSE.

We restrict to configurations with an equal number of up and down spins, and move between configurations by flipping a pair of opposing spins (so that the net magnetization is unchanged). This can be done since the ground state exists in the spin sector having $\sum_{i,j} \sigma^z_{i,j} = 0$. We assume real, positive correlators and use the Marshall-sign rule to determine the correct sign for each configuration. Otherwise we apply the same minimization routine as used above for the TIM. The ground state energy was found for different plaquette types and lattice sizes, and the results for a $14 \times 14$ lattice are shown in Table III.

For plaquette correlators, the energies obtained are consistent with previous calculations. We see that plaquette correlators are able to minimize the energy much better than bond correlators, and that the discrepancy between the correlator types is larger than with the TIM. For example, for $L = 14$, even $3 \times 3$ correlators are better able to minimize the ground state energy than bonds with $r_{\text{max}} = 7$, where bonds are long enough to couple every pair of sites on the lattice, and where the number of parameters is comparable. Thus the relative behavior of bond correlators with respect to plaquette correlators is worse than with the TIM. This is likely to be as a result of a more complicated energy landscape for
the AFHM than for the TIM, such that the fragmented structure imposed by the bond correlators makes it difficult to describe the ground state. Consistent with the RVB picture, it therefore appears that having a large number of local parameters is more important to accurately describe the ground state of this model than to provide longer-ranged parameters. This is further highlighted by the fact that the energy estimated using bond correlators with \( r_{\text{max}} = 4 \) is the same (within error) as the energy estimated with \( r_{\text{max}} = 7 \), so the improvement with increasing bond size is seen to saturate.

As with the TIM, we also investigate how well the correlations are described by the different correlator types by calculating the staggered magnetization. The staggered magnetization has been previously calculated using a variety of tensor network methods.\(^{8,46–48}\) It has been found to be challenging to determine accurately, with the estimated value in general being larger than the value found using SSE.\(^{44}\)

<table>
<thead>
<tr>
<th>Correlator type</th>
<th>No. of elements</th>
<th>Energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>( r_{\text{max}} = 1 )</td>
<td>16</td>
<td>-0.6560(3)</td>
</tr>
<tr>
<td>( r_{\text{max}} = 2 )</td>
<td>48</td>
<td>-0.6627(3)</td>
</tr>
<tr>
<td>( r_{\text{max}} = 3 )</td>
<td>96</td>
<td>-0.6641(2)</td>
</tr>
<tr>
<td>( r_{\text{max}} = 4 )</td>
<td>160</td>
<td>-0.6646(1)</td>
</tr>
<tr>
<td>( r_{\text{max}} = 5 )</td>
<td>240</td>
<td>-0.6648(3)</td>
</tr>
<tr>
<td>( r_{\text{max}} = 6 )</td>
<td>336</td>
<td>-0.6647(2)</td>
</tr>
<tr>
<td>( r_{\text{max}} = 7 )</td>
<td>504</td>
<td>-0.6649(3)</td>
</tr>
<tr>
<td>2 \times 2 plaquettes</td>
<td>16</td>
<td>-0.6567(3)</td>
</tr>
<tr>
<td>3 \times 3 plaquettes</td>
<td>512</td>
<td>-0.6662(2)</td>
</tr>
<tr>
<td>4 \times 4 plaquettes</td>
<td>65,536</td>
<td>-0.6685(4)</td>
</tr>
</tbody>
</table>

TABLE III. Energy per site \( E/J \) found using stochastic minimization for the AFHM in a 14 × 14 system for different correlator types. The result from using a SSE is \( E/J = -0.670222(7) \) (Ref. 44).

FIG. 10. (Color online) Staggered magnetization found using different plaquette types against the inverse of the lattice size \( L \). (a) shows the staggered magnetization \( M_2^z = C_{L/2,L/2} \) and (b) shows the staggered magnetization \( M_2^z(L) = \frac{1}{L^2} \sum_{i,j} (-1)^{i+j} C_{i,j} \). In both plots the black dashed line denotes the staggered magnetization in the thermodynamic limit found using SSE (Ref. 44).

An alternative definition for the staggered magnetization is\(^{49}\)

\[
M_1^z(L) = C_{L/2,L/2} = \langle S^{[0,0]} \cdot S^{[L/2,L/2]} \rangle. \tag{6}
\]

An alternative definition for the staggered magnetization in the thermodynamic limit \( M(\infty) \). The scaling behavior of \( M_{1,2} \) has been predicted using chiral perturbation theory and spin-wave theory, and is given by \( M_{1,2}^2(L) = M(\infty)^2 + h_{1,2} L + \cdots \).\(^{14,15,45,51}\)

We calculate the two-point correlation function, and similar to a previous work\(^{13\text{a}}\) it is found that the \( z \)-component of the correlations departs from the \( x,y \)-component of the correlations — the \( z \) correlations tend to zero while the \( x,y \) correlations remain finite. This is likely to be due to the preferential treatment of the \( z \) axis in the Monte Carlo method and inherent in the CPS correlator elements, i.e. the chosen basis. We also find that very small changes in the estimated ground state energy can lead to large differences in the estimated staggered magnetization. The results of these calculations are shown in Fig. 10. In contrast to the results in previous sections where bond correlators performed better, we find that plaquette correlators provide a better estimate of the long-range order, with an extrapolated value of \( M_1(\infty) \approx 0.33(1) \) and \( M_2(\infty) = 0.322(7) \) found using \( 4 \times 4 \) plaquette correlators closest to the value \( M(\infty) = 0.3070(3) \) found using stochastic series expansion.\(^{14\text{a}}\)

VI. SUMMARY AND OUTLOOK

We have investigated the performance of the most common forms of correlator product states for describing the

\[ M_2^z(L) = \frac{1}{L^2} \sum_{i,j} (-1)^{i+j} C_{i,j}. \]
The system sizes reached here are comparable with experimental cold-atom lattice systems, and the ability to efficiently model spin systems is useful for rapidly progressing cold-atom simulations of quantum magnetism. A general result for the optimal correlator structure for a given system is not straightforward to deduce, but our findings still provide useful insights for applications to condensed matter systems. For example, we would expect plaquette correlators to be the optimal choice for the $J_1 - J_2$ model studied in Ref. 22 and for the Fermi-Hubbard model. However, we may expect bond correlators to perform better for the Bose-Hubbard model, where the choice of the local Fock basis may allow a good description of the ground state with a smaller number of local parameters, or in spin-systems which are highly anisotropic like the TIM. In addition, the Laughlin wave function has an exact CPS description using bond correlators.

It may be that an optimal CPS ansatz consists of a hybrid of bond and plaquette correlators, such as strings of sites (in a similar set up to string bond states) linking all sites on the lattice, or extended rectangular plaquette correlators. This would provide a larger number of local parameters while still allowing the correlators to have a long range. In further work, the CPS description will be applied to other systems, such as the Bose-Hubbard model in the presence of an artificial magnetic field, in the regime where the fractional quantum Hall states are predicted to occur.

VII. ACKNOWLEDGEMENTS

SA thanks Vlatko Vedral and Nigel Cooper for useful discussions. This work was supported by the UK EPSRC through project EP/J010529/1 and by the ESF program EuroQUAM (EPSRC grant EP/E041612/1). SRC and DJ thank the National Research Foundation and the Ministry of Education of Singapore for support.

Appendix A: Exact and efficiently samplable tensor network states

For the case of n. n. bond correlators, the resulting CPS is in fact equivalent to an MPS with internal dimension $\chi$ equal to the physical dimension $d$. Note that this equivalence does not hold in the reverse direction, i.e. not all MPS with $\chi = d$ are equivalent to a CPS. Correlator product states can be thought of as a special class of matrix product states that can be factorized using the copy tensor (also known as the diagonal in category theory, the COPY-gate or the COPY-dot in circuits). The copy tensor copies a chosen set of physical basis states e.g. $|x\rangle$ where $x = \uparrow, \downarrow$, and subsequently breaks up into disconnected states. This is essential for such states to be efficiently and exactly sampled.

This is illustrated in Fig. 11(a), where the copy tensor is represented using a black dot, while the large circle represents a matrix formed of the correlator elements. The copy tensor has one input (the physical leg) and multiple outputs (two for the case of bond correlators in one dimension). This network is equivalent to an MPS, where each matrix in the network can be formed from the correlators and copy tensors. The copy tensor acts
by taking the value of the physical leg in the chosen basis and copying it to all its legs. As shown in Fig. 11(b), this acts to break up the bonds between the matrices while ensuring that neighboring matrices are contracted according to the same physical index on a pair of virtual legs. The result of this is a product of scalars, shown in Fig. 11(c), which are the correlators for the given input configuration used.

In two dimensions, correlator product states can also be considered to be equivalent to a special case of PEPS — in this case the copy tensor copies the physical index to four different correlator matrices. String bond states are another special factorization of PEPS where the contractible components are themselves MPS, rather than a correlator matrix. States such as these can then be sampled exactly-efficiently, as they consist of product of tensors describing a 2×2 plaquette for each plaquette. Efficient sampling is again possible due to the copy tensor.

systems that do not obey an area law.

Appendix B: Numerical methods

1. Calculating expectation values

Expectation values of the energy and other operators are determined from a CPS by using Monte Carlo importance sampling. The energy \( E \) is given by

\[
E = \langle \psi | H | \psi \rangle = \frac{\sum_s W^*(s') \langle s'| H | s \rangle W(s)}{\sum_s |W(s)|^2}, \tag{B1}
\]

where we assume that the wave function is not normalized, as is generally the case when using CPS. The energy can be expressed as a weighted sum \( E = \sum_s P(s) E(s) \) which is as a sum over configurations of the product of the local energy \( E(s) \) and the probability \( P(s) \), given by

\[
E(s) = \sum_{s'} W^*(s') \langle s'| H | s \rangle, P(s) = \frac{|W(s)|^2}{\sum_s |W(s)|^2}. \tag{B2}
\]

By replacing the Hamiltonian \( H \) with any operator \( O \) its expectation value \( \langle \psi | O | \psi \rangle \) can be similarly computed. For example, for the TIM described in section III, calculation of the interaction energy terms \( \sigma_z^{[i,j]} \) is straightforward since the operators are diagonal: the local energy \( E_{zz}(s) \) is given by

\[
E_{zz}(s) = - \sum_{i=1,j=1}^L (s^{[i,j]} s^{[i+1,j]} + s^{[i,j]} s^{[i,j+1]}), \tag{B3}
\]

where \( s^{[i,j]} = \pm 1 \). In a translationally invariant system, the sum over all sites can be dispensed with and the exchange energy simply taken with respect to the first (or any other) spin and its neighbors.

To calculate the transverse energy terms \( g_{ij}^{[i,j]} \) we use \( \sigma_x = \sigma_+ + \sigma_- \). The only non-zero terms of the energy estimator \( E(s) = \sum_{s'} W^*(s') \langle s'| H | s \rangle \), are those for which \( s' \) differs from \( s \) by a single spin-flip, so the local transverse energy is given by

\[
E_2(s) = - \sum_{i=1,j=1}^L \frac{W(s'_{ij})}{W(s)}. \tag{B4}
\]

where \( s'_{ij} \) denotes that the spin on site \((i,j)\) has been flipped. The absolute magnetization \( \langle |\sigma_z| \rangle \) of the ground state can also be calculated. Note that the expectation value of the magnetization \( \langle \sigma_z \rangle \) is always be zero, since the global \( \mathbb{Z}_2 \) symmetry of the system means the ground state forms equal superpositions of configurations with all spins flipped. However, the absolute magnetization quantifies how well the spins are aligned with
one another, and is found by summing the local absolute magnetization

$$|\sigma_z(s)| = \frac{1}{L^2} \left| \sum_{i=1}^{L} s_i \right|.$$  

The probability of a given configuration is never explicitly calculated using equation (B2). Instead, the configurations are visited according to importance sampling, calculated acceptance probability $|W(s')/W(s)|^2$. There are a total of $F \times N$ spin flips per bin, where we call $F$ the number of sweeps per bin, and $N$ is the number of spins (since we visit each spin sequentially in a given sweep). As is typical, we also first perform a few warm-up sweeps that do not contribute to the estimates to ensure that the random walk through configuration space starts in an equilibrium position, and also sample the local expectation value only every $s^{th}$ sweep, where $s$ is the sample rate $\sim 10$, to allow a larger portion of the configuration space to be visited and to ensure that the configurations that are sampled are independent from one another.

At no point during the calculation is knowledge of the normalization of the wave function needed. The most time-consuming step in the calculation is determining the correlator fraction only the correlators that represent the number of bins is increased as $G = G_0 k$ per iteration to slow the cooling rate, where typically $F_0$ and $G_0 \sim 10$. The step size is reduced per iteration as $\delta = \delta_0 k^{-\eta}$. We find best results with $\eta$ between 0.75 and 0.9. After an initial run with a relatively large step size to get close to the minimum, the resulting correlators are then used as a starting point for a new run of iterations (i.e. $k$ is reset) where $F_0$ and $G_0$ remain unchanged, but $\delta_0$ is decreased. Depending on the system parameters this can reduce the energy further, while for other regimes (i.e. away from criticality) the energy has already converged after the first run.

To achieve convergence, $F$, the number of sweeps in a bin is increased every iteration, and the step size $\delta(k)$ is reduced. For every iteration $k$, $F$ is given by $F_0 k$, and the number of bins is increased as $G = G_0 k$ per iteration to slow the cooling rate, where typically $F_0$ and $G_0 \sim 10$. The step size is reduced per iteration as $\delta = \delta_0 k^{-\eta}$. We find best results with $\eta$ between 0.75 and 0.9. After an initial run with a relatively large step size to get close to the minimum, the resulting correlators are then used as a starting point for a new run of iterations (i.e. $k$ is reset) where $F_0$ and $G_0$ remain unchanged, but $\delta_0$ is decreased. Depending on the system parameters this can reduce the energy further, while for other regimes (i.e. away from criticality) the energy has already converged after the first run. After the minimization is complete, the procedure is repeated for a single iteration with zero step size and large $F$ and $G$, to obtain an accurate estimate of the expectation values.

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2. Energy minimization

The correlator elements that minimize the energy are found using a stochastic minimization method which requires only the sign of the first derivative of the energy with respect to the correlator elements. Specifically, the first derivative is given by

$$\frac{\partial E}{\partial C_{s(i)}^{(i)}} = 2(\Delta_{s(i)}^{(i)}(E) - \langle \Delta_{s(i)}^{(i)} \rangle(E)),$$  

where $\Delta_{s(i)}^{(i)}(E)$ is given by

$$\Delta_{s(i)}^{(i)}(s) = \frac{1}{W(s)} \frac{\partial W(s)}{\partial C_{s(i)}^{(i)}}.$$  

This is trivial to compute, since $W(s)$ is simply a product of the different correlators $C$. If each correlator is only used once $\Delta_{s(i)}^{(i)}(s) = 1/C_{s(i)}^{(i)}$, whereas if the same correlator is used for multiple sites (e.g. in a translationally invariant system) then $\Delta_{s(i)}^{(i)}(s) = b^{(i)} / C_{s(i)}^{(i)}$, where $b^{(i)}$ is the number of times the correlator $C_{s(i)}^{(i)}$ appears in the product for the correlator amplitude for configuration $s$.

Following the procedure in Ref. 33, the expectation value of the derivative is calculated in the same way that other operators are calculated for $F$ sweeps in a bin, and the derivatives for all correlators are calculated simultaneously. After this every correlator is updated according to

$$C_{s(i)}^{(i)} \rightarrow C_{s(i)}^{(i)} - r \delta(k) \text{sign} \left( \frac{\partial E}{\partial C_{s(i)}^{(i)}} \right)^*$,$$  

where $r$ is a random number between 0 and 1, and $\delta(k)$ is the step-size for a given iteration $k$. Unlike the Newton method, the second derivative is not required, considerably simplifying the calculation. This method has been found to give better convergence than the Newton method, since it avoids the sizable statistical errors in the second derivative and also because it does not exactly follow the direction of steepest decent which can be more efficient.

To achieve convergence, $F$, the number of sweeps in a bin is increased every iteration, and the step size $\delta(k)$ is reduced. For every iteration $k$, $F$ is given by $F_0 k$, and the number of bins is increased as $G = G_0 k$ per iteration to slow the cooling rate, where typically $F_0$ and $G_0 \sim 10$. The step size is reduced per iteration as $\delta = \delta_0 k^{-\eta}$. We find best results with $\eta$ between 0.75 and 0.9. After an initial run with a relatively large step size to get close to the minimum, the resulting correlators are then used as a starting point for a new run of iterations (i.e. $k$ is reset) where $F_0$ and $G_0$ remain unchanged, but $\delta_0$ is decreased. Depending on the system parameters this can reduce the energy further, while for other regimes (i.e. away from criticality) the energy has already converged after the first run. After the minimization is complete, the procedure is repeated for a single iteration with zero step size and large $F$ and $G$, to obtain an accurate estimate of the expectation values.
FIG. 13. (Color online) Energy during minimization of the plaquette correlators for the two-dimensional TIM in a 31 by 31 system with \( g = 3.05 \) (see section III). The initial minimization routine is performed with \( \delta = 0.02 \) for 25 iterations, followed by a further round of minimization with \( \delta = 0.005 \) for 30 iterations (although not shown, for 4 \( \times \) 4 plaquette correlators the further round of minimization has 40 iterations).

For bond correlators, the best results are found when the step size \( \delta \) is a function of bond length as well as iteration number \( k \). For each bond in the correlator, we define the radius \( r \) as the maximum displacement along one of the lattice indices, and scale the step size as \( \delta(r, k) = \delta(k)r^{-0.75} \). For the first round of minimization we minimize the bonds for a given radius separately in sequence, while in the second round of iterations all correlator elements are updated simultaneously.


34 S. Sachdev, Quantum phase transitions, 1st ed. (Cambridge University Press, 2000).


36 The pseudo-critical point in a finite-size anomaly, where evidence of the critical point in the thermodynamic limit can be observed in a finite-sized system.


