Tensor product variational formulation applied to pentagonal lattice

Michal Daniška and Andrej Gendiar

Institute of Physics, Slovak Academy of Sciences, SK-845 11, Bratislava, Slovakia

E-mail: michal.daniska@savba.sk and andrej.gendiar@savba.sk

Received 5 June 2015, revised 31 August 2015
Accepted for publication 14 September 2015
Published 7 October 2015

Abstract
The uniform two-dimensional variational tensor product state is applied to the transverse-field Ising, XY, and Heisenberg models on a regular hyperbolic lattice surface. The lattice is constructed by tessellation of the congruent pentagons with the fixed coordination number being four. As a benchmark, the three models are studied on the flat square lattice simultaneously. The mean-field-like universality of the Ising phase transition is observed in full agreement with its classical counterpart on the hyperbolic lattice. The tensor product ground state in the thermodynamic limit has an exceptional three-parameter solution. The variational ground-state energies of the spin models are calculated.

Keywords: tensor product state, quantum spin systems, non-Euclidean geometry, phase transition

(Some figures may appear in colour only in the online journal)

1. Introduction

The tensor product state (TPS) has been proved to be an appropriate ansatz for obtaining a ground-state of strongly correlated quantum systems. Numerous computational approaches and methods have been developed and successfully applied to one-dimensional (1D) and two-dimensional (2D) quantum systems [1–3]. The main purpose of this work is to address the TPS to 2D spin systems on a non-trivial lattice configuration, which is represented by an infinite hyperbolic surface. The lattice is made from the pentagonal tessellation and forms a negatively curved surface of the constant Gaussian curvature. Such a lattice geometry has not yet been considered for 2D quantum systems in the thermodynamic limit. Lacking information on constructing an appropriate numerical algorithm for the pentagonal lattice is given.

The adherence to the mean-field-like universality class was concluded in the 2D classical Ising systems on various types of hyperbolic lattices. The mean-field-like behavior of the
phase transition exponents is caused by the non-Euclidean geometry of the underlying lattices [4–6], and is not related to the analysis by the mean-field approximation. The classical spin systems have been analyzed by the corner transfer matrix renormalization group (CTMRG) method [7], which is a variant of the DMRG for 1D quantum systems [8]. Now, we extend our earlier studies, which dealt with the classical spin systems, to quantum spin systems on the curved pentagonal lattice surface.

We, therefore, propose an approximative scheme of the round-a-face TPS, which is related to the standard vertex type TPS with two-state auxiliary variables [1, 2]. Our main intention is to approximate the ground state of spin-$$\frac{1}{2}$$ models by a minimal number of variational parameters. Improvements of the numerical accuracy go beyond the scope of this work.

The paper is organized in the following way. Section 2 specifies the model Hamiltonians on the Euclidean and the non-Euclidean lattices. The variational approach is discussed with respect to the TPS approximation. A short description of the numerical algorithm and discussion of tensor symmetries are included within two subsections. The numerical results are analyzed in section 3, and we comment the results in section 4.

2. The model

We calculate the ground-state energy of the quantum Ising, XY, and Heisenberg models on two different lattices: the Euclidean square lattice and the hyperbolic pentagonal lattice. Figure 1 depicts the two lattice types constructed by tessellation of the congruent polygons: either by the squares ($$p = 4$$) shown on the left or by the pentagons ($$p = 5$$) on the right. The alternating gray-white color scheme of the polygons is chosen for improving the visibility only. The spin variables are located on the vertices of the regular polygons with the constant coordination number, which is equal to four in both cases. We study properties of the spin system in the thermodynamic limit, i.e., the number of the lattice vertices is infinite. The Hamiltonian $$\mathcal{H}$$ of the three models defined on the both lattices can be expressed in the following compact form.
where $G^{(p)}_k$ represents the local Hamiltonian of the $p$-sided polygon, the lattice is constructed from, and $k$ marks the position of the polygon on the lattice. The summation runs over all the positions of the polygons $k$. The polygon on the $k$th position is described by the ordered set of spin indices $k_1, k_2, \ldots, k_p$, see figure 2, where $k_i$ stands for the unique number which is assigned to the corresponding vertex within the labeling scheme of the lattice vertices. The local Hamiltonian has the expression

$$H = \sum_{k} G^{(p)}_k,$$

(1)

where $G^{(p)}_k$ represents the local Hamiltonian of the $p$-sided polygon, the lattice is constructed from, and $k$ marks the position of the polygon on the lattice. The summation runs over all the positions of the polygons $k$. The polygon on the $k$th position is described by the ordered set of spin indices $k_1, k_2, \ldots, k_p$, see figure 2, where $k_i$ stands for the unique number which is assigned to the corresponding vertex within the labeling scheme of the lattice vertices. The local Hamiltonian has the expression

$$G^{(p)}_k = -\frac{1}{2} \sum_{i=1}^{p} \left[ J_{xy}(S^x_{k_i}S^x_{k_{i+1}} + S^z_{k_i}S^z_{k_{i+1}}) + J_z S^z_{k_i}S^z_{k_{i+1}} + \frac{h}{4}(S^x_{k_i} + S^x_{k_{i+1}}) \right]$$

(2)

where $S^x_{k_i}, S^y_{k_i}, S^z_{k_i}$ are the Pauli operators, and the spin indices obey the cyclic condition $k_{p+1} \equiv k_1$. The $x$-component of the external magnetic field is described by the variable $h$, and the spin couplings $J_{xy}$ and $J_z$ specify the three models. In particular, $J_{xy} = -J_z = 1$, and $h = 0$ describe the Heisenberg model, $J_{xy} = 1$ and $J_z = h = 0$ specify the XY model, whereas $J_{xy} = 0, J_z = 1$ with an arbitrary $h$ lead to the transverse field Ising model (TFIM)\[9\]. Since the ferromagnetic ordering of the models leads to a simpler TPS formulation, we have opted for the positive coupling, $J_{xy} = 1$, and we consider dimensionless units throughout the entire work. The constant prefactors $\frac{1}{2}$ and $\frac{h}{4}$ reflect the sharing of the spin couplings and the magnetic field, respectively, if the Hamiltonian is formed by the polygonal tessellation in equation (1).

Our objective is to obtain the ground-state of the system

$$\left| \Phi_p \right> = \lim_{N \to \infty} \sum_{\sigma_1, \sigma_2, \ldots, \sigma_N} \Phi^{(1)}_{\sigma_1 \sigma_2 \cdots \sigma_N} \left| \sigma_1 \sigma_2 \cdots \sigma_N \right>$$

(3)

in the thermodynamic limit by a variational minimization of the ground-state energy normalized per bond

---

\[1\] The Hamiltonian energy spectrum remains identical if $J_{xy} = \pm 1$ provided that $J_z = 0$ or $-1$ (for $h = 0$) because the lattice is bipartite. Note, that generalization to the antiferromagnetic ordering is trivial for the square geometry, however, the pentagonal lattice geometry would result in strong frustration for all of the models.
For instance, there are 32 base spin configurations for the pentagons, which can be represented in the arrow notation as \( \{ \uparrow \uparrow \downarrow \downarrow \downarrow \} \), \( \{ \downarrow \downarrow \downarrow \downarrow \uparrow \} \), \( \{ \downarrow \downarrow \downarrow \uparrow \uparrow \uparrow \} \), \( \{ \uparrow \uparrow \downarrow \uparrow \uparrow \} \). Thus, the approximative ground state in the form of the polygonal TPS \(^2\) \cite{1} has the following form in the thermodynamic limit

\[
\Psi_p = \lim_{N \to \infty} \sum_{\sigma_1, \sigma_2, \ldots, \sigma_N} \prod_{k} W_p(\{ \sigma_k \}) | \sigma_1 \sigma_2 \ldots \sigma_N \rangle ,
\]

where the sum runs over the \( 2^N \) base spin states. Since the TPS \( \Psi_p \) has the product structure of the identical tensors \( W_p \), the variational problem in equation (4) is in the thermodynamic limit equivalent to the minimization of the local energy of an arbitrary bond in the lattice center (to avoid boundary effects)

\[
E_p = \min_{\psi_p} \frac{2}{p} \left( \langle \psi_p | G_p^{(0)} | \psi_p \rangle \right) > E''_p ,
\]

where \( \ell \) is the index of a polygon containing the selected central bond and the factor \( 2/p \) reflects that each polygon contains \( p \) bonds shared with neighboring polygons. Moreover, the product structure of \( \Psi_p \) enables us to express the denominator

\[
\langle \psi_p | G_p^{(0)} | \psi_p \rangle = \sum_{\sigma, \sigma'} \prod_{\ell} W_p^\sigma(\{ \sigma' \}) (G_p^{(0)})_{\sigma', \sigma} | \sigma' \rangle | \sigma \rangle \equiv \mathcal{D}(W_p(\{ \sigma \}))
\]

and the numerator

\[
\langle \psi_p | G_p^{(0)} | \psi_p \rangle = \sum_{\sigma, \sigma'} \prod_{\ell} W_p^\sigma(\{ \sigma' \}) (G_p^{(0)})_{\sigma', \sigma} | \sigma' \rangle | \sigma \rangle \equiv \mathcal{N}(W_p(\{ \sigma \}))
\]

as sole functions of the tensor elements \( W_p(\{ \sigma \}) \), where we removed the subscript \( k \) due to the uniform TPS. Here, \( (G_p^{(0)})_{\sigma', \sigma} \) stands for the corresponding matrix element of the local Hamiltonian \( G_p^{(0)} \), \( \delta_{\sigma, \sigma'} \) is the Kronecker symbol, and \( \{ k \} \setminus \{ \ell \} \) denotes the set of all polygon indices except for the index \( \ell \).

Consequently, the minimization over the set of variational parameters \( \Phi_{p, \sigma_1 \ldots \sigma_N} \) in equation (4) is replaced by a much simpler problem

\(^2\) The non-physical (auxiliary) states in the TPS language are represented as states with two degrees of freedom only. Such an approximation enhances the mean-field behavior around the criticality and is a compromise to make the calculations feasible numerically due to the exponentially increasing complexity of the hyperbolic lattice structure.
where we minimize over $2^p$ tensor elements $W_p(\{\sigma\})$ only. This set can be further reduced if additional constraints are taken into account. The translational and rotational symmetries shrink them to three variational parameters, provided that the ground state $|\Psi_0\rangle$ is evaluated at zero magnetic field, i.e., no symmetry-breaking mechanism is present in the system. The calculation of the numerator $\mathcal{N}(W_p(\{\sigma\}))$ and the denominator $\mathcal{D}(W_p(\{\sigma\}))$ in equation (9) is carried out separately by means of a numerical algorithm described below.

Thus defined TPS approach offers a new perspective of estimating the 2D quantum systems on hyperbolic surfaces. Such a significant approximation is sufficient only for the case of $p = 5$ since we have observed a weak entanglement and non-critical behavior of the classical Ising model on variety of hyperbolic lattices in our earlier studies [4–6]. An exponential decay of the density matrix spectra and the correlation function result in the phase transition, where the correlation length $\xi \lesssim 1$ is finite, reaching its maximal value at the phase transition [17]. For this reason, we apply the current scheme of TPS as it can be a sufficient approximation for the ground-state properties of the models on the pentagonal lattice. (However, the square lattice requires to consider a larger dimension in the tensors $W_4$, see [13] and [16].)

2.1. The TPFV algorithm

We refer to the algorithm as the tensor product variational formulation (TPVF) and it consists of two parts. The first one evaluates the ratio in equation (9) by applying the CTMRG method [7] separately to the numerator and the denominator for a given set of the tensor elements $W_p(\{\sigma\})$. The second part contains a multi-dimensional minimizer, the Nelder–Mead simplex algorithm [10–12], which uses the first part to search for the optimized set of the tensor elements $W_p(\{\sigma\})$, which minimize the ratio in equation (9). The minimizer starts from an initial simplex in the space of free variational parameters, one vertex of which is specified by the initial tensor elements $W_p(\{\sigma\})$. The simplex undergoes an iterative sequence of size changes and moves towards lower energies and stops if the energy in equation (9) converged.

The CTMRG was originally developed to study 2D classical spin systems. Primarily, it evaluates the partition function (preferably, in the thermodynamic limit) as the configuration sum of the tensor product of the Boltzmann weights $W_b$. The central idea of this study is to apply CTMRG to quantum systems by replacing the concept of the Boltzmann weight $W_b$ from the classical statistical mechanics by the tensors $W_p$. In order to do this, let us introduce a double-layer tensor $Z_p$ with the tensor elements

Figure 3. The double-layer tensor structure of $Z_4(\{\sigma'_k, \sigma_k\})$ on the left and $Z_5(\{\sigma'_k, \sigma_k\})$ on the right.
Notice that there are $22p$ double-layer base spin configurations. Figure 3 graphically depicts the double-layer tensors $Z_p$ at the position $k$, where each shaded polygonal area represents $W_p$. Thus, in the language of the classical statistical mechanics, the general expression for the denominator $W_p$ in equation (7) corresponds to a tensor product object, which is equivalent to the partition function of a (non-physical) classical Hamiltonian given by the product of the tensors $Z_p$.

This generalization of the Boltzmann weight enters the CTMRG algorithm for both the lattice geometries and the consequent numerical calculation yields the denominator $W_p$ for the given set of the tensor elements $W_p$ according to equation (7). A similar approach can be also used to determine the numerator $N(W_p)$, as it differs from $D(W_p)$ only by the additional double-layer structure at the central position $\ell$ containing the local Hamiltonian $G^{(p)}$. The detailed survey of the CTMRG algorithm is explained in [4, 5, 7] for both the square and the pentagonal lattices. We point out only the most relevant part of the CTMRG algorithm in the following.

The concept of CTMRG is built on corner transfer tensors $C_j^{(n)}$ and transfer tensors $T_j^{(n)}$, where $j = 1, 2, ..., p$. Each of the (corner) transfer tensors is composed of the polygon representing tensors $Z_p$ via recurrence relations indexed by the iteration step $n$ (as specified later). Further details of the construction of the transfer tensors are given in [4, 7]. As a result, the (corner) transfer tensors represent a specific lattice sector formed by the corresponding polygons. The square and the pentagonal lattices are constructed from a central polygon surrounded by the alternating sectors represented by the tensors $C_j^{(n)}$ and $T_j^{(n)}$. The central polygon is represented by the tensor $Z_p$ or $W_p G^{(p)} W_p$ in the structure of the denominator $D(W_p)$ and the numerator $N(W_p)$, respectively. Figure 4 illustrates the situation...
for the case of $D(W_p(\{\sigma\}))$. Consequently, the relations (7) and (8) for the denominator $D(W_p(\{\sigma\}))$ and the numerator $\tilde{N}(W_p(\{\sigma\}))$, if formulated in the CTMRG language of the (corner) transfer tensors, take the form

$$D(W_p(\{\sigma\})) = \lim_{n \to \infty} \text{Tr} \left( Z_p \prod_{j=1}^p C_j^{(n)} T_j^{(n)} \right),$$

(11)

$$\tilde{N}(W_p(\{\sigma\})) = \lim_{n \to \infty} \text{Tr} \left( W_p G_j^{(n)} W_p \prod_{j=1}^p c_j^{(n)} T_j^{(n)} \right).$$

(12)

The tensors $C_j^{(n)}$ and $T_j^{(n)}$ expand their sizes iteratively, following the recurrence relations [4, 5]

$$C_j^{(n)} = Z_p T_{j-1}^{(n-1)} \left[ C_j^{(n-1)} T_j^{(n-1)} \right]^{p-3},$$

(13)

$$T_j^{(n)} = Z_p T_{j-1}^{(n-1)} \left[ c_j^{(n-1)} T_j^{(n-1)} \right]^{p-4},$$

(14)

where the transfer tensors satisfy the cyclic condition $T_j^{(n)} = T_{p+j}^{(n)}$. The infinite TPS geometry is built up gradually by increasing the iteration step $n = 2, \ldots, \infty$, which induces an exponential increase of the degrees of freedom of the (corner) transfer tensors. The calculations are kept numerically feasible by means of the renormalization group procedure, which integrates out the least probable configurations in the tensors determined by the reduced density matrix [7, 8].

The recurrence relations are initialized in the first iteration step, $n = 1$ by construction of the tensors $C_j^{(n=1)}$ and $T_j^{(n=1)}$ from the tensor $Z_p$ for all $j = 1, 2, \ldots, p$. If the spin variables $\sigma_i$ are explicitly included in the tensors, we have

$$C_j^{(1)}(\{\sigma_k, \sigma_k, \sigma_k, \sigma_k, \sigma_k\}) = \sum_{\sigma_i \sigma_{i4}^{\sigma_{k4}}} Z_s(\{\sigma_i\}),$$

$$T_j^{(1)}(\{\sigma_k, \sigma_k, \sigma_k, \sigma_k, \sigma_k, \sigma_k\}) = \sum_{\sigma_i \sigma_i} Z_s(\{\sigma_i\}).$$

(15)

### Table 1. The full spin configurations for $p = 4$ sorted by $\theta_i$. The last column lists the reduced parameter set $a_i$ with $\ell = \min(j, p-j)$.

| $j$ | $W_4(\theta_j)$ | $\theta_j \equiv \{\sigma_k, \sigma_k, \sigma_k, \sigma_k\}$ | $a_i$ |
|-----|-----------------|-------------------------------------------------|-------|
| 0   | $W_4(\theta_0)$ | $\{\uparrow \downarrow \downarrow \downarrow\}$ | $a_0$ |
| 1   | $W_4(\theta_1)$ | $\{\uparrow \uparrow \uparrow \uparrow\}$ | $a_1$ |
| 2   | $W_4(\theta_2)$ | $\{\uparrow \uparrow \uparrow \uparrow\}$ | $a_2$ |
| 2'  | $W_4(\theta_{2'})$ | $\{\uparrow \uparrow \uparrow \uparrow\}$ | $a_2'$ |
| 3   | $W_4(\theta_3)$ | $\{\uparrow \uparrow \uparrow \uparrow\}$ | $a_3$ |
| 4   | $W_4(\theta_4)$ | $\{\uparrow \uparrow \uparrow \uparrow\}$ | $a_4$ |
 equivalents. The prime symbol is used for such con
decreases down to four parameters for the Hamiltonian models.

As mentioned earlier, let the polygon base spin con
ctions for the pairs

\[
\begin{array}{c}
W_0(\theta_0) \\
W_1(\theta_1) \\
W_2(\theta_2) \\
W_2(\theta_2') \\
W_3(\theta_3) \\
W_3(\theta_3') \\
W_4(\theta_4) \\
W_5(\theta_5)
\end{array}
\]

\[
\begin{array}{c}
\{\downarrow \downarrow \downarrow \downarrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \} \\
\{\uparrow \uparrow \uparrow \uparrow \}
\end{array}
\]

\[
\begin{array}{c}
a_0 \\
a_1 \\
a_2 \\
a_2' \\
a_2 \\
a_2' \\
a_1 \\
a_0
\end{array}
\]

\[
\begin{array}{c}
\theta_{fi} \\
\theta_{fi}(p-j) \\
\theta_{fi}(p-j)
\end{array}
\]

2.2. The tensor symmetries

As mentioned earlier, let the polygon base spin configuration \( \{ a_1 \} \equiv (\sigma_1, a_2, \ldots, a_{n_p} ) \) be given in
the spin-arrow notation. Let function \( \theta_{fi} \) return all such configurations for which the integer
number \( j \) counts the number of the spins aligned upward as listed in tables 1 and 2. Each line
in the tables contains such spin configurations, which are identical with respect to the rota-
tional symmetry operations of the \( p \)-sided polygon. We make a difference between the spin
configurations \( \theta_{fi} \) and \( \theta_{fi}' \) when \( j=2 \) (or \( j=3 \) for the case of \( p=5 \)) as they are not rotationally
equivalent. The prime symbol is used for such configuration (and its rotations), in which
either the two \( (j=2) \) or three \( (j=3) \) spins, aligned upward in each polygon, cannot be
grouped together (or equivalently, if the alternating spin alignment is maximal).

As a consequence, we count six or eight distinguishable configurations \( \theta_0, \theta_1, \theta_2, \theta_3, \theta_4, \theta_5 \) for \( p=4 \)
and \( p=5 \), respectively. These configurations determine the number of the variational
parameters \( \mathcal{W}_p(\theta_{fi}) \) used in the calculation of the TFIM. If, however, the spontaneous sym-
metry-breaking does not affect the solution, the total number of the variational parameters
decreases down to four parameters for the Hamiltonian models (for both lattices \( p \)). Therefore,
the spin-inversion symmetry reduces the variational parameters due to the absence of the
preferred direction of the spontaneous magnetization. In particular, the spin configuration
probability for the pairs \( \theta_0 \), \( \theta_1 \), \( \theta_2 \), and \( \theta_{p-1} \), etc becomes identical, which can be formally
generalized into the equations

\[
\begin{align*}
\mathcal{W}_p(\theta_{fi}) &= \mathcal{W}_p(\theta_{p-j}) = a_{\min(j,p-j)}, \\
\mathcal{W}_p(\theta_{fi}') &= \mathcal{W}_p(\theta_{p-j}) = 1.
\end{align*}
\]

There are the three equations in the upper expression defining the new variational parameters
\( a_0, a_1, a_2 \), and one equation in the lower expression for \( a_2' \), which has already been eliminated
from the set of the free parameters by putting \( a_2' \equiv 1 \) being the normalization condition in \( \mathcal{W}_p \).

Hence, the three free variational parameters suffice to approximate the ground-state wave
function of the models with no spontaneous symmetry-breaking phases. For the same reason,
if we consider the system without the spin-inversion symmetry, there are either five or seven
free variational parameters for \( p=4 \) or \( p=5 \), respectively, excluding the normalization
parameter \( \mathcal{W}_p(\theta_{fi}) \equiv 1 \).

The number of the operations performed by the implementation of the Nelder–Mead
algorithm we used scales linearly with the number of the free variational parameters [11]. Of
course, increasing the number of the parameters prolongs the computational time, and may
encounter numerical instability caused by trapping the system in a local minimum of the
energy, rather than approaching to the correct global minimum, which corresponds to $E^{(p)}_0(p)$. On the other hand, a faster Nelder–Mead optimization enables to improve the accuracy by increasing the number of the CTMRG states kept (the effective block spin states [7, 8]) and make TPVF more efficient.

3. Numerical results

As we have mentioned earlier, we consider two types of the lattices. The Euclidean one is defined on the regular square lattice characterized by $p = 4$, which serves as a reference for the hyperbolic pentagonal lattice with $p = 5$ (see figure 1), which is of our main interest. First, we start analyzing the ground-state properties of the TFIM in details, where the number of the variational parameters $W_p(\theta_j)$ is either five (for $p = 4$) or seven (for $p = 5$), respectively, excluding the normalization parameter $W_1(p) \equiv 1$. We analyze the TFIM and its phase transition by the following three ways: (1) by evaluating the optimized parameters $W_p(\theta_j)$, (2) by the expectation value of the magnetization constructed from the ground state, and (3) by the ground-state energy per bond $E^{(p)}_0(p)$ and its second derivative. Second, we reduce the number of the free variational parameters down to three (if all symmetries are considered) and calculate the ground-state energies of the XY and Heisenberg models.

In our numerical calculations we keep at most 20 effective block spin states [7], which is sufficient to reach more than six-digit convergence in $\mathcal{N}(W_p(\{\sigma\}))$ and $\mathcal{D}(W_p(\{\sigma\}))$. Notice that the numerical accuracy is mainly given by the uniform TPS approximation. Further improvement of the numerical accuracy requires to consider a non-uniform TPS (i.e. violation of the translational lattice symmetry), and a gradual expansion of the tensor order by implementation of auxiliary variables [13] is necessary. This is, however, beyond the scope of our interest.
The Nelder–Mead algorithm has been tested for various initial conditions, most of them leading to the identical results. Typically, we started with a simplex, where we set all the coordinates of one of its vertices to unity (the vertex coordinates represent the free variational parameters $W_p(\theta_j)$). The initial simplex size was set to $0.1^{10, 11}$.

Figure 5 and 6 illustrate behavior of the optimized variational parameters $W_p(\theta_j)$ in the TFIM with respect to the external magnetic field $h$ for the particular lattice geometry $p$. It is evident that above a certain magnetic field (depicted by the vertical dotted lines), specific pairs of the variational parameters collapse onto identical values, which yields four single curves in this region for both the lattices. These pairs are exactly those coupled by the system of equations (16), i.e. those representing spin configurations which are equivalent with respect to spin inversion. The magnetic field, at which the collapse causes a singular behavior of $W_p(\theta_j)$, corresponds to the quantum phase transition of TFIM at the magnetic field $h^{(c)}$, which we analyze below. In the ordered phase at $h < h^{(c)}$, the distinct optimized values of the coupled parameters $W_p(\theta_j)$, as specified in tables 1 and 2, reflect the existence of the spontaneous symmetry-breaking in the TFIM for both the lattice types. In the disordered phase at $h \geq h^{(c)}$, the four-parameter description coincides with the free variational parameters $a_0$, $a_1$, $a_2$, and the normalization parameter $a_2'$. This confirms the relevance of the additional symmetries in such systems, where the spontaneous symmetry-breaking mechanism is not present, such as in the XY and Heisenberg systems at the zero magnetic field.

Having evaluated the optimized free variational parameters, we can easily reconstruct the approximative eigenstate $|\psi_p\rangle$ as the TPS and apply it for the evaluation of the spontaneous magnetization

$$
\langle S_p^z \rangle = \frac{\langle \psi_p | S_p^z | \psi_p \rangle}{\langle \psi_p | \psi_p \rangle},
$$

(17)
where $\alpha = x$ or $z$. The expectation value of the spin operator $S_i^{\alpha}$ is evaluated in the central part of the lattice in order to suppress all boundary effects. Here, $S_z^p$ denotes the order parameter of TFIM and specifies the quantum phase transition at the phase transition field. The resulting dependence of magnetization $S_z^p$ and $S_x^p$ with respect to the magnetic field $h$ is shown in figure 7. The quantum phase transition is reflected by the singular behavior of all magnetization profiles.

Analyzing the TFIM, we calculated $h_c^{(4)} = 3.158$ for the Euclidean square geometry and $h_c^{(5)} = 3.264$ for the hyperbolic pentagonal lattice. The most relevant value of the critical magnetic field for the TFIM on the 2D Euclidean lattice (as obtained by a recent tensor RG algorithm) yields $h_c^{TRG} = 3.0439$ [16]. Since the TPS is built up by the tensors $W_4$ of the too low dimension, the long-range order correlations are excluded [6], and the mean-field-like dependence near the phase transition is necessarily observed. The expectation value of the magnetization $S_z^p$ obeys the scaling relation

$$\langle S_z^p(h) \rangle \propto (h_c^p - h)^{\beta_p}.$$  \hfill (18)

The mean-field exponent $\beta_p = \frac{1}{2}$ is observed regardless of the lattice geometry. As a reference, the numerical TRG analysis gives correct $\beta_4^{TRG} = 0.3295$ on the square lattice (free of any mean-field approximations), which is also in agreement with Monte Carlo simulations [16]. The inset of figure 7 displays the magnetization squared, which confirms the mean-field-like behavior for both $p$ by its linear dependence on $h$ if approaching the phase transition. The blue dashed lines serve as guides for the eye in order to enhance the linearity.

A more detailed analysis of the TPVF approximation near the phase transition can be visualized by evaluating the effective (field dependent) exponent $\beta_{cl}^p(h)$, which converges to $\beta_p$ when approaching the phase transition field $h_c^p$.
Figure 8. The detailed dependence of the inverse effective magnetic exponent on the magnetic field in the logarithmic form. The inset shows the effective exponent at wider magnetic field scale. The blue dashed line estimates behavior of the correct effective exponent for the Euclidean lattice.

Figure 9. The ground-state energy of the TFIM with respect to the magnetic field $h$. The small difference between the two energies is plotted in the inset.

\[
\beta_p = \lim_{h \to h_c^{(p)}} \beta_{\text{eff}}^{(p)}(h) = \lim_{h \to h_c^{(p)}} \frac{\partial \ln \langle S^z_0(h) \rangle}{\partial \ln (h_c^{(p)} - h)}
\]

as plotted in figure 8. The effective exponent obviously converges to the mean-field exponent $\beta_p = \frac{1}{2}$ for both lattice types if the phase transition field is approached from the ordered phase, i.e., if $\ln(h_c^{(p)} - h) \to -\infty$. The inset shows the same dependence on larger scales. The critical exponent on the square lattice (the black curve for $p = 4$) starts deviating at around $h > 2.0$ from the expected exponent (estimated by the blue dashed curve), which is known to converge to $\beta_{\text{TRG}} = 0.3295$ [16].
The ground-state energies normalized per bond $E_p^0$ given by equation (9) are depicted in figure 9. There is a tiny difference in both of the energies. Therefore, we plotted the energy difference $E_p^0 - E_p^{MC}$ in the inset (which has no physical meaning) in order to point out the difference. The two vertical lines in the inset correspond to the phase transitions (for reference only). The magnetic susceptibility
\[
\chi_p = -\frac{\partial^2 E_p^0}{\partial h^2}
\] (20)
is an independent physical quantity, which characterizes the phase transition. Figure 10 shows the functional dependence of the susceptibility on the magnetic field. The calculation of $\chi_p$ requires a precise data for the ground-state energy because of performing the second derivative numerically. The shape of the non-diverging discontinuity in $\chi_p$ at the phase transition magnetic field $h_p^{(p)}$ is the typical consequence of the mean-field universality.

The XY and the Heisenberg models are critical (at $p = 4$) at zero magnetic field. Therefore, the full tensor symmetries are considered, see tables 1 and 2, resulting in the three free variational parameters $a_0$, $a_1$, and $a_2$ which is sufficient to obtain the ground-state energy.
per bond $E_0^{(p)}$. Table 3 summarizes thus calculated optimized free variational tensor parameters and the energies. The comparison of $E_0^{(p)}$ on the square lattice with the Monte Carlo results by Sandvik [14, 15] serves as an estimate of the exact ground-state energy $E_0^{(5)}$ for the XY and the Heisenberg models on the hyperbolic pentagonal lattice, which has not been determined yet.

4. Conclusions

We have applied the TPVF algorithm to the calculation of the quantum systems (transverse field Ising, XY, and Heisenberg models) on the pentagonal hyperbolic lattice with the constant negative Gaussian curvature and the coordination number fixed to four. As the reference lattice, the Euclidean square lattice was taken. We have used the RG-based algorithm, which approximates the ground state as the product of identical tensors representing the congruent polygons; either squares ($p = 4$) or pentagons ($p = 5$) for the Euclidean or hyperbolic lattices, respectively. We have analyzed the symmetries in the tensors. The uniform TPS reduces the infinite number of the variational parameters in the thermodynamic limit down to six (for $p = 4$) or eight (for $p = 5$) for the Ising model in the presence of the transverse magnetic field. If no spontaneous symmetry-breaking mechanism is present, the number of the free variational parameters approximating the tensor product ground state further shrinks down being three only. We have analyzed the phase transition of the Ising model by (1) the numerical calculation of the optimized variational parameters, (2) the expectation value of the spin polarization, (3) the phase transition magnetic exponent, (4) ground-state energy and the magnetic susceptibility. The mean-field-like universality class was observed in the vicinity of the phase transition magnetic fields $h_{0}^{(p)}$. The ground-state energy per bond $E_0^{(p)}$ has been evaluated for the XY model and the Heisenberg model for $p = 4$ and $p = 5$.

Our earlier studies of the classical spin lattice models on various types of the hyperbolic surfaces [4–6] exhibited a couple of interesting features, which are important to mention here. The classical Ising model on the pentagonal lattice results in a higher phase transition temperature if compared to the Euclidean square lattice [5]. Analogously, the TFIM model in this work also exhibits the identical feature, i.e., $h_{c}^{(5)} > h_{c}^{(4)}$. Classical spin systems on any hyperbolic lattice belong to the mean-field-like universality class in accord with the current study. The mean-field-like behavior of the exponents in the classical systems originates from the hyperbolic lattice geometry, not from the numerical CTMRG method; the CTMRG accurately reproduces the critical exponents on the 2D Euclidean lattices [4–6]. The low dimension of the tensors in the TPS approximation of the round-a-face type suppresses the quantum long-range correlations on the square lattice ($p = 4$) near the criticality, where we obtained $h_{c}^{(4)}$ deviating by 3.7% if compared to the recent numerical analysis [16]. The discrepancy is the consequence of the low dimension of the tensor $W_4$ appearing in the ground-state approximation, which limits the numerical accuracy if approaching the criticality, and it results in the mean-field exponent $\beta = 1/2$. Therefore, it is essential to distinguish between the mean-field approximation of the method used and the mean-field-like behavior caused by the hyperbolic lattice geometry. That means that we have dealt with both of the ‘mean-field’ aspects in the current study. The physical reasoning of the mean-field-like behavior caused by the hyperbolic lattice comes from the exceedance of the critical dimensionality (being $d_{c} = 3$ or $d_{c} = 4$, respectively, for the classical or quantum systems with the nearest-neighbor spin couplings). Notice that the Hausdorff dimension of the pentagonal lattice is infinite.
Strictly speaking, classical spin systems exhibit the phase transitions exclusively in the center of the infinite hyperbolic lattices because of strong boundary effects. The ratio between the number of the spins on the boundary with respect to the number of the spin on the remaining lattice area inside is \( \approx 2.73 \) for the pentagonal lattice. For this reason, the hyperbolic lattice becomes significantly sensitive to the boundary effects \([18]\). On the other hand, the boundary effects on the Euclidean lattices play no role in the thermodynamic limit when the lattice size is expanded to infinity. If calculating the correlation length \( \xi \) on the pentagonal lattice (the distance between two spins measured along a geodesics) for the classical Ising model, no divergence at the phase transition was present, i.e., \( 0 < \xi \lesssim 1 \) \([17]\). Moreover, the correlation function was found to decay exponentially even at the phase transition, which is the direct consequence of the finiteness of the correlation length at the phase transition (and is related to an analogous exponential decay of the density matrix spectra) \([6]\). In that sense, the phase transition on pentagonal hyperbolic surface is non-critical, and we use the term critical for the spin models on the square (Euclidean) lattice only since it is related to the divergence of the correlation length by definition. The identical non-critical behavior for \( p = 5 \) has also been confirmed in this work (for instance, by evaluating the non-divergent magnetic susceptibility at \( h^{(p)} \)). It is highly non-trivial to evaluate the correlation length unambiguously as both of the 'mean-field' aspects are closely related and almost impossible to be clearly separated or distinguished.

An exponentially fast decay of the reduced density matrix spectra has been observed for the classical systems on hyperbolic surfaces \([6, 17]\). Similar behavior for the quantum systems had to be taken into account in the current work. For all the above-mentioned reasons, we conjecture the TPFV analysis of the models on the hyperbolic pentagonal lattice is more accurate than on the Euclidean ones. It is caused by the mean-field-like behavior (the mean-field-like universality) as the consequence of the pentagonal hyperbolic lattice with the infinite Hausdorff dimension, which has been studied by the numerical improved mean-field approximation. Our preliminary results of the identical quantum Hamiltonian on different hyperbolic lattices (to be published elsewhere) support our claims and are in complete agreement with the conjectures we have made for the classical spin systems.

Acknowledgments

We thank Tomotoshi Nishino and Frank Verstraete for valuable discussions. This work was supported by the grants QIMABOS APVV-0808-12, VEGA-2/0130/15, and EU project SIQS No. 600645.

References

[1] Orus R 2014 Ann. Phys., NY 349 117
[2] Schollwöck U 2011 Ann. Phys., NY 326 96
[3] Schollwöck U 2005 Rev. Mod. Phys. 77 259
[4] Ueda K, Kremar R, Gendiar A and Nishino T 2007 J. Phys. Soc. Japan 76 084004
[5] Kremar R, Gendiar A, Ueda K and Nishino T 2008 J. Phys. A: Math. Theor. 41 125001
[6] Gendiar A, Kremar R, Andergassen S, Daniška M and Nishino T 2012 Phys. Rev. E 86 021105
[7] Nishino T 1996 J. Phys. Soc. Japan 65 891
[8] White S R 1996 Phys. Rev. B 48 10385
[9] Nishio Y, Maeshima N, Gendiar A and Nishino T 2004 arXiv:cond-mat/0401115
[10] http://gnu.org/software/gsl/
[11] Galassi M et al 2009 GNU Scientific Library Reference Manual 3rd edn (Network Theory) ISBN 0954612078
[12] Nelder J A and Mead R 1965 A simplex method for function minimization *Comput. J.* 7 308–13
[13] Gendiar A, Maeshima N and Nishino T 2003 *Prog. Theor. Phys.* 110 691
[14] Sandvik A W and Hamer C J 1999 *Phys. Rev.* B 60 6588
[15] Sandvik A W 1997 *Phys. Rev.* B 56 11678
[16] Xie Z Y et al 2012 *Phys. Rev.* B 86 045139
[17] Iharagi T, Gendiar A, Ueda H and Nishino T 2010 *J. Phys. Soc. Japan* 79 104001
[18] Baxter R J 1982 *Exactly Solved Models in Statistical Mechanics* (London: Academic)