Solving frustrated quantum many-particle models with convolutional neural networks

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(Dated: September 27, 2018)

Recently, there has been significant progress in solving quantum many-particle problem via machine learning based on the restricted Boltzmann machine. However, it is still highly challenging to solve frustrated models via machine learning, which has not been demonstrated so far. In this work, we design a brand new convolutional neural network (CNN) to solve such quantum many-particle problems. We demonstrate, for the first time, of solving the highly frustrated spin-1/2 J1-J2 antiferromagnetic Heisenberg model on square lattice via CNN. The energy per site achieved by the CNN is even better than previous string-bond-state calculations. Our work therefore opens up a new routine to solve challenging frustrated quantum many-particle problems using machine learning.

PACS numbers:

I. INTRODUCTION

The successes of machine learning in image recognition,1 playing Go,2,3 etc., stimulate interests in using machine learning techniques to solve physics problems, such as designing optical experiments,4 processing signals in finding gravity waves,5,6 quantum teleportation photon spot recognition,7 topological invariants,8,13 inferring Hamiltonian solutions,14,15 classifying quantum states based on topological invariants,16 classifying spin configurations into phases,17,18 solving ground states of Bose-Hubbard model19 and so on.

Recently machine learning has been applied to study quantum many-particle problems.19,20 This posed serious challenges, because it requires much higher precision than traditional machine learning problems. It has been demonstrated that by unsupervised learning, the restricted Boltzmann machine (RBM) can solve the ground states of transverse-field Ising models and anti-ferromagnetic Heisenberg models in rather high precision.20 The ground states of those models obey the Marshall-Peierls sign rule (MPSR)21 where the wave functions can be represented by positive numbers, suitable for RBM with real parameters.

The representation ability of RBM for some advanced states has been investigated, and often connects to another state-of-the-art method for many-particle problems, namely the tensor network state (TNS) method.22 The equivalence between RBM and TNS has been investigated in Refs.20,23,24 It has been suggested that RBM can represent the quantum states beyond area law.20 Furthermore, the extension of RBM, namely deep Boltzmann machine (DBM), can be transformed into tensor networks, and DBM can be a general representation of quantum many-particle states.25

In the community of machine learning, it is believed that convolutional neural networks (CNN) have shown more efficiency compared to the neural networks within dense connections.26 Recent works try to bridge the state-of-the-art convolutional neural network to the tensor network states.27 It has been proposed that the information reuse introduced by overlapping convolution filters can enhance the state representation ability. The convolutional neural network has shown effectiveness in finding the critical points in phase transitions of Ising models28 which infers the network’s valid state representation ability.

However, so far solving frustrated quantum many-particle models on large two-dimensional lattices is still challenging for neural networks. The ability to solve the frustrated model is a milestone for neural networks, because the non-frustrated models can be efficiently solved to extremely high accuracy by quantum Monte Carlo (QMC) methods. In this work, we built a CNN to solve the ground states of two-dimensional frustrated quantum spin models, more specifically the antiferromagnetic spin-1/2 J1-J2 Heisenberg model on square lattices. This model is highly frustrated, and considered as one of the most interesting and challenging spin models.29,30 The network is built using the elements from state-of-the-art neural networks. By design, the network associates the many-body wave functions with high-order spin correlation functions. We name such CNN as the convolutional quantum neural state (CQNS). We solve the ground states at J2 = 0 and J2/J1 = 0.5 and it is shown that CQNS can achieve the ground energies lower than those obtained by the previous string bond state (SBS) calculations.30,31 We further compare the spin correlation functions, focusing on the convolution filters’ size effects. We show that if the size of the filter is too small, CQNS can not capture the correct long-range spin correlations, which leads to poor energy precisions. However, for systems with the correlation length shorter than the system size, it is efficient to use relative smaller convolution filters. The CQNS is a general method which can be easily applied to study other quantum many-particle problems.
outputs the wave function coefficient \( w(S) \), which is the products of all the final output neurons. Figure 1(a) depicts the network structure of using one convolution filter, which generates one chain of output neurons. When \( M \) convolution filters are used, there will be \( M \) chains of output neurons. To correlate different filters, we sum all the output neurons on the same site generated from all filters, as Fig. 1(b) denotes. The wave function coefficient \( w(S) \) is obtained by taking the products of all the neurons in the final chain.

One can easily extend the above one-dimensional neural network to higher dimensions. Figure 1(c) depicts a two-dimensional CQNS with a \( K \times K \) convolution filter, with \( K = 3 \). After convolution layer, a max-pooling is performed on adjacent four neurons followed by a transposed convolution restores the dimensions. The wave function coefficient \( w(S) \) is the product of all the neurons on the final output plane.

To reveal how the CQNS works, we use the \( L=4 \) spin chain as an example. Considering a convolution filter with length \( K = 3 \) and one additional spin padding in both the head and the tail of the spin chain, the convolution operation can be written in the matrix form:

\[
\begin{bmatrix}
  p_1 \\
p_2 \\
p_3 \\
p_4
\end{bmatrix} = \mathbf{b} + \begin{bmatrix}
w_1 & w_2 & w_3 & 0 & 0 & 0 \\
w_0 & w_1 & w_2 & w_3 & 0 & 0 \\
0 & 0 & w_1 & w_2 & w_3 & 0 \\
0 & 0 & 0 & w_1 & w_2 & w_3
\end{bmatrix} \begin{bmatrix}s_2 \\
s_1 \\
x_1 \\
x_2
\end{bmatrix},
\]

where \( S=[s_4, s_1, s_2, s_3, s_4, s_1]^\top \) is the input spin configuration and \( \mathbf{b} \) is the bias vector of the convolution filter. In our cases, the input number of each site is \( s_1=\pm 1 \). The output neurons after convolution are the input neurons of the max-pooling, i.e.,

\[
\begin{bmatrix}
g_1 \\
g_2
\end{bmatrix} = \begin{bmatrix} \text{max}(p_1, p_2) \\
\text{max}(p_3, p_4)
\end{bmatrix}.
\]

We then perform transposed convolution,

\[
\begin{bmatrix}h_1 \\
h_2 \\
h_3 \\
h_4
\end{bmatrix} = \begin{bmatrix}d_1 & 0 \\
d_2 & 0 \\
0 & d_1 \\
0 & d_2
\end{bmatrix} \begin{bmatrix}g_1 \\
g_2
\end{bmatrix},
\]

to restore the original size of the lattice.

Equations (3) describe the transformations between the input spins and the output neurons when using one convolution filter. Using \( M \) convolution filters lead to \( M \) output chains, as shown in Fig. 1(b). The final output chain is the direct summation of the output chains from all convolution filters, i.e.,

\[
\begin{bmatrix}h_1 \\
h_2 \\
h_3 \\
h_4
\end{bmatrix} = \sum_{m=1}^{M} \begin{bmatrix}d_1^{(m)} & g_1^{(m)} \\
d_2^{(m)} & g_2^{(m)} \\
d_1^{(m)} & g_2^{(m)} \\
d_2^{(m)} & g_2^{(m)}
\end{bmatrix}.
\]

The final wave function coefficient is the product of all neurons in the final output chain, i.e., \( w(S) = h_1 h_2 h_3 h_4 \).
From the above equations, it is easy to see that for a given spin configuration \( S \) of a \( N \) sites system, \( h_i \) is linear combination of \( s_k \) plus a constant, i.e., \( h_i = A_{1,k}s_1 + A_{2,k}s_2 + \cdots + A_{N,k}s_N + c_i \), where \( A_{i,k} \) is the coefficient of spin \( s_k \) in \( h_i \), and \( c_i \) is a number. However, due to the max-pooling, the coefficients \( A_{i,k} \) are spin configuration dependent. Therefore we have,

\[
w(S) = \sum_{n_1, \cdots, n_N} g(n_1, \cdots, n_N; s_1, \cdots, s_N) s_1^{n_1} \cdots s_N^{n_N},
\]

where \( n_1 + n_2 + \cdots + n_N \leq N \), and \( g(n_1, \cdots, n_N; s_1, \cdots, s_N) \) are given by CQNS. We note that \( g \) functions are also spin configuration dependent, because of the max-pooling. Equation 5 associates the many-body wave functions with the high order spin correlation functions. This is crucial for the success of CQNS that are very different from traditional CNN, in which the nonlinearity of the neural networks is usually introduced by the activation functions.

It is easy to show that CQNS can exactly represent the classical Néel state, which has alternating spin up and spin down in the lattice, and is the ground state of the classical antiferromagnetic Heisenberg model, using a single convolution filter. The filter kernel has the form \([0, 0, 0, 5]\), and biases are taken to be zero. Figure 2 depicts the intermediate neurons after convolution and max-pooling when the input state is a Néel state. The output neurons after max-pooling are all positive, thus the wave function coefficient \( w(S) > 0 \). It is easy to verify that at least one of the output neurons after max-pooling is zero if the input state is not the Néel state, and therefore \( w(S) = 0 \). For a quantum antiferromagnetic Heisenberg model more filters are needed to represent the ground state. As an example, for the spin chain of \( L = 4 \), it is easy to verify that the ground state can be exactly represented by the CQNS with four filters, which are \([-0.5, 0.5, 0.5], [0.5, 0.5, -0.5], [-0.5, -0.5, 0.5] \) and \([0.5, 0, 0, 0, 5]\) and biases for the filters are -1.5, -1.5, -0.5 and 0 respectively.

### III. Numerical Investigations on \( J_1-J_2 \) Model

We benchmark the CQNS on the two-dimensional spin-\( 1/2 \) \( J_1-J_2 \) Heisenberg model,

\[
H = \sum_{\langle i,j \rangle} J_1 s_i \cdot s_j + J_2 \sum_{\langle (i,j) \rangle} s_i \cdot s_j ,
\]

(6)

where \( \langle i,j \rangle \) denotes the nearest neighbour sites and \( \langle (i,j) \rangle \) denotes the next nearest neighbour sites. We assume both \( J_1, J_2 > 0 \), and take \( J_1 = 1 \) in all calculations in this work. The \( J_1-J_2 \) model is one of the most interesting and challenging quantum spin models due to the strong frustrated interactions, and the ground state near \( J_2/J_1 = 0.5 \) is still under intensive debate. In this work, we only focus on the ground state energies of the model.

The total energy of the system can be calculated as:

\[
E = \langle \Psi | H | \Psi \rangle = \frac{1}{2} \sum_h w_h \sum_S w_S H_{h,S},
\]

(7)

where \( H_{h,S} = \langle S | H | S' \rangle \). Our goal is to minimize the total energy, which can be achieved by the stochastic gradient method, where the energy \( E \) and the gradients \( G \) are calculated via Monte Carlo sampling over spin configurations,

\[
E = \langle E_S \rangle, \quad G = \langle O_S E_S \rangle - \langle E_S \rangle \langle O_S \rangle,
\]

(8)

where \( E_S = \sum_{S'} \frac{w(S')}{w(S)} H_{S',S} \) and \( O_S = \frac{1}{w(S)} \frac{\partial w(S)}{\partial a_i} \). Here \( a_i \) are all the parameters in the CQNS.

However, the neural networks are highly nonlinear, therefore the system may be easily trapped in some local minima via the stochastic gradient descending method. To overcome this difficulty, we optimize CQNS via a recently developed replica-exchange molecular dynamics method, which has been successfully applied to optimize the SBS one type of TNS. We map the quantum many-particle problem to a classical mechanical problem, in which we treat the parameters of CQNS as the generalized coordinates of the system. We optimize the energy of the virtual system using a replica exchange molecular dynamics method. By exchanging the system configurations among higher and lower temperatures, it can explore large phase space and therefore effectively avoid being stuck in the local minima. Details of the method are presented in Ref. 31.

In our simulations, we use 56 temperatures. Initially the temperatures distribute exponentially between the highest \( 1/\beta_0 = 10^{-3} \) and lowest \( 1/\beta_55 = 10^{-5} \) temperatures. The MD step length is set to \( \Delta t = 0.01 \). For each temperature we start with random weights and
For number dramatically improve the energy. For example, for filter energy much. On the other hand, increasing filter size can increasing the number of filters does not improve the decreases to -0.6522. However for the small filters, fur-
ters is -0.6461. The energy can be improved by increasing
obtained by CQNS. The energy for
Carlo whereas the dashed line is the result of quantum Monte
E
the ground state energy per site obtained by the SBS,
sometric states (EPS) and QMC are also shown in the dotted line (-0.6581), the dash-dotted line (-0.6699) and the dashed line (-0.6715), respectively.

We study $J_1$-$J_2$ model on a $10 \times 10$ two-dimensional square lattice, with periodic boundary conditions. Figure 3(a) depicts the numerical results for $J_2=0$ as functions of the number of filters $M$ and size of filters $K$. The reference ground energies obtained by the string-bond-state (SBS), entangled-plaquette-state (EPS) and the energies used for replica exchange configurations. For each MD step, we sample 5000 spin
configurations. For each MD step, we sample 5000 spin
configurations. The energies for
replica exchange are averaged over 200 MD steps.

We study $J_1$-$J_2$ model on a $10 \times 10$ two-dimensional square lattice, with periodic boundary conditions. Figure 3(a) depicts the numerical results for $J_2=0$ and $J_2=0.5$, with different number of filters and filter sizes. The filter sizes are taken to be $K \times K$. The obtained ground state energies are summarized in Table I.

The energies per site for $J_2=0$ calculated with different number of filters and filter sizes are shown in Table I. We also show the total energies calculated from other methods as comparisons. The dotted line denotes the ground state energy per site obtained by the SBS, $E = -0.6581_{23}$ and the dash-dotted line denotes the result of the entangled plaquette states $E = -0.6699_{24}$ whereas the dashed line is the result of quantum Monte Carlo $E = -0.6715_{25}$. The solid dots are the results obtained by CQNS. The energy for $K = 3$ and $M=128$ filters is -0.6461. The energy can be improved by increasing the number of filters. When using 512 filters, the energy decreases to -0.6522. However for the small filters, further increasing the number of filters does not improve the energy much. On the other hand, increasing filter size can dramatically improve the energy. For example, for filter number $M=128$, the energy is -0.6536 for $K = 5$, -0.6596 for $K = 7$ and -0.6649 for $K = 9$. As shown in the figure, when $K = 9$, the energy per site converges faster than using smaller filters, the energies for $M=128$, 192 and 256 filters are -0.6649, -0.6659 and -0.6665, respectively. Based on the numerical results, CQNS with filter size of $K=7$, 9 can exceed SBS. From the numerical results, it is revealed that larger filter number and larger filter size lead to more accurate ground states, however increasing filter size is more efficient than increasing filter number.

At $J_2 = 0.5$, the model is highly frustrated, where QMC suffers from sign problem, and therefore is much more difficult to calculate. In Fig. 3(b), we show the calculated energies with different filter size $K$ and number of filters $M$. The ground state energy per site calculated by SBS is -0.4705, which is shown in dotted line as a comparison. According to the figure, the energy has limited improvement from $M=128$ to 256 filters. However increasing filter size can significantly improve the energy. For $M=128$, the energy is -0.4455 for $K=3$, -0.4627 for $K=5$ and -0.4736 for $K=9$. For $M=256$, the energy per site is -0.4475 for $K = 3$, -0.4649 for $K = 5$ and -0.4715 for $K = 7$. CQNS can exceeds SBS with filter size $K \geq 7$.

We further calculate the spin correlation functions, and the results are shown in Fig. 3(c) and (d) for $J_2=0$ and $J_2=0.5$ respectively. The error bars are smaller than the width of the solid lines and therefore not shown. The spin correlation functions $C(r)$ are calculated as,

$$C(r) = \frac{1}{2L^2} \sum_{i,j} (s_{i+r,j} s_{i,j} + s_{i,j} s_{i+r,j}). \quad (9)$$
The calculated results show that the correlation functions obey power-law decay. We fit $C(r)$ as $C(r) = C_0 + \alpha r^{-\gamma}$. We plot the correlation functions $G(r) = \alpha r^{-\gamma}$, i.e., the constants have been subtracted. Because of the periodic boundary condition, the maximal distance is $r = L/2$. The calculated exponents $\gamma$ for different $K$ are listed in Table II. From these results, one may see that using small filters greatly underestimate the decay length of the spin correlation functions, and therefore can not represent the ground state very well. Increase filter size may describe the correlation functions more accurately and therefore can not represent the long-range correlation by multiple convolution layers. Otherwise the neural network can not capture the long quantum correlations.

The CQNS is expected to work well for quantum systems with short correlation lengths, e.g., gapped systems. On the other hand, to simulate very large systems, which are much larger than their correlation lengths $\xi$, the CQNS is also advantageous. One may use a suitable filter size $K$, which is larger than $\xi$, but still much less than the system sizes, therefore the number of tunable parameters in CQNS can be greatly reduced.

### IV. CONCLUSIONS AND OUTLOOKS

We design a convolutional quantum neural network, namely CQNS, to study quantum many-particle problems. By design, CQNS associates the many-body wave functions with high-order spin correlation functions, which is crucial for its success. We use CQNS to study the spin-1/2 $J_1$-$J_2$ Heisenberg model which is strongly frustrated and challenging to solve. We have obtained fairly accurate ground state energies that are even better than previous SBS calculations. The CQNS is a general and flexible method that can be easily applied to different Hamiltonian, with various boundary conditions (open, periodic and cylindrical, etc.) and in different dimensions. It therefore provides a new powerful tool to study the long-standing quantum many-particle problems.

The CQNS used in this paper is as simple as three layers. Using deep neural networks may further enhance the state representation ability, as it is possible to capture the long-range correlation by multiple convolution layers with smaller filters in each layer. It recently has been proposed that deep CNN may efficiently represent the many-particle states that are even beyond area law. Designing a practical deep CQNS is an interesting and promising route to follow.

**Acknowledgments**

The authors appreciate insightful discussions with Shao-Jun Dong and Chao Wang. The construction and computation of neural networks in this paper is performed by the mainstream deep learning framework PyTorch. The numerical calculations have been done on the USTC HPC facilities. This work was supported by the National Natural Science Foundation of China (Grant No. 11674306, 61590932), National key R&D program (No. 2016YFA0301300, 2016YFA0317300).

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