Zero-site DMRG and the optimal low-rank correction

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A zero-site density matrix renormalization algorithm (DMRG0) is proposed to minimize the energy of matrix product states (MPS). Instead of the site tensors themselves, the “message” tensors between neighbor sites are sequentially optimized, leading to a local minimization step that is independent of the physical dimension of the site. To avoid local minima, two new global perturbations based on the optimal low-rank correction to the current state are introduced. They are determined variationally as the MPS closest to the one-step correction of the Lanczos or Jacobi-Davidson eigensolver, respectively. These perturbations mainly decrease the energy and are free of hand-tuned parameters. Compared to existing single-site enrichment proposals, our approach gives similar convergence ratios per sweep while the computations are cheaper by construction. Our methods may be useful in systems with many physical degrees of freedom per lattice site, and we test them on the periodic Heisenberg spin chain for spins $S = 1, S = 3$ and $S = 5$.

I. INTRODUCTION

The exponential compression of the basis for a given portion $S$ of a quantum one-dimensional system using its density matrix is at the heart of the traditional DMRG algorithm [1,2]. The state wavefunction is expressed in terms of the single-site formulation considers a rank-3 tensor with dimensions $m \times d \times m$ determined by the site and its environment $E$. Conceptually, it should be possible to formulate the optimization of the wavefunction (which dominates the costs in DMRG) considering only a pure bipartition at a time, that is, asking for the optimal wavefunction expressed in terms of the $S$ and $E$ renormalized basis (both of size $m$). However, current computational schemes [3–9] explicitly include one or two sites and basis enrichment processes are mixed. For instance, the optimization and renormalization steps.

In the language of matrix product states (MPS) [6], the optimization is performed for a single- or two-site tensor at a time instead of a central zero-site (or “message”) tensor. The former could be motivated by the idea of expressing the state in a richer basis, optimizing the state out, and then renormalizing the basis again. In such an approach, the optimization and basis enrichment processes are mixed. For instance, the single-site formulation considers a rank-3 tensor with dimensions $m \times d \times m$ as the effective wavefunction. Some redundant information is present here, because only $m$ of the $md$ rows of the tensor (reshaped as an $md \times m$ matrix) are linearly independent. It would also be conceptually useful to separate the optimization and renormalization steps.

In this work, we propose an optimization step for a sequence of central zero-site rank-2 tensors $C_i$ of dimensions $m \times m$, depicted in Fig. 1 below. A priori, an objection to this approach is that it increases the chances that the DMRG is stuck in local minima. An important part of this work will then be to solve this in an efficient manner. This leads us to introduce an enrichment step based on the optimal low-rank correction to the global state. We will show that this method markedly increases the convergence of the algorithm and avoids metastable solutions.

The outline of the paper is as follows. In Sec. I, we present the zero-site DMRG and explain its basic properties. In Sec. II we introduce the enrichment step, with two approximate schemes (Lanczos and Jacobi-Davidson) to obtain the optimal low-rank correction. We also review the previous approaches [3–9], and establish the equivalence between [3] and [9]. Sec. II summarizes our algorithm, and Sec. V presents numerical results for the Heisenberg spin chain with spins $S = 1, S = 3$ and $S = 5$. Sec. VI contains our conclusions and perspectives.

II. ZERO-SITE DMRG

A quantum state $\psi$ written as a product of matrices [6]

$$|\psi\rangle = \sum_{\sigma_1...\sigma_L} M_{i_1}^{\sigma_1}...M_{i_L}^{\sigma_L} |\sigma_1...\sigma_L\rangle$$  

is called a matrix product state (MPS). Here, $\sigma_i$ (of dimension $d_i$ with $i = 1,2,...,L$) labels the physical degree of freedom of the site $i$ for a system of $L$ sites. $M_i$ is a rank-3 tensor of dimensions $m_{i-1} \times d_i \times m_i$. In practice, $m_i \leq m$ where $m$ is the MPS bond dimension, and $m_0 = m_L = 1$.

An important property of the MPS is its gauge degree of freedom. For arbitrary invertible matrices $X_i$, the identity $L_i = X_i^{-1}X_i$ can be inserted between $M_i$ and $M_{i+1}$, effectively changing the matrices to $M_i' = X_{i-1}M_iX_i^{-1}$ while keeping the state invariant. It allows us to choose the so-called left (right) normalization for the matrices $M_i' = A_i^{\sigma}/(M_i'' = B_i^{\sigma})$ satisfying

$$\sum_\sigma (A_i^{\sigma})^\dagger A_i^{\sigma} = I_i \quad \text{or} \quad \sum_\sigma B_i^{\sigma}(B_i^{\sigma})^\dagger = I_{L-1}.$$  

We now introduce (as in [10]) the MPS zero-site canonical form at site $i$:

$$|\psi\rangle = \sum_{\sigma_1...\sigma_L} A_1^{\sigma_1}...A_i^{\sigma_i}C_iB_{i+1}^{\sigma_{i+1}}...B_L^{\sigma_L}|\sigma_1...\sigma_L\rangle,$$  

similar to the single-site canonical form where the central matrix $M_i''$ is decomposed as $M_i'' = A_i^{\sigma_i}C_i$. We illustrate this in Fig. 1. One advantage of this representation is its local expression for the square norm $\langle \psi | \psi \rangle = \text{tr}(C_i^\dagger C_i)$. We refer to $C_i$ as the message between tensors $-$ it contains the singular values and the entanglement of the bipartition in this case. In DMRG terminology, the products $A_1^{\sigma_1}...A_i^{\sigma_i}$ and $B_{i+1}^{\sigma_{i+1}}...B_L^{\sigma_L}$ represent the left and right renormalized basis for $S$ (system)
Figure 1. MPS representation of the state, with a bipartition (vertical dashed line) into $S$ and $E$. The message tensor $C_i$ is optimized at this step.

and $E$ (environment) respectively, and $C_i$ is the (strictly bipartite) wavefunction.

If the Hamiltonian is also written as a matrix product operator (MPO),

\[ \hat{H} = \sum_{\sigma_1...\sigma_L} W_{i}^{\sigma_1} \cdots W_{L}^{\sigma_L} |\sigma_1 \cdots \sigma_L\rangle \langle \sigma_1 \cdots \sigma_L|, \]

(4)

then the energy of a normalized state $\psi$ is

\[ E = \langle \psi | \hat{H} | \psi \rangle = C_i^\dagger \langle \partial_i \psi | \hat{H} | \partial_i \psi \rangle C_i \equiv C_i^\dagger H_{\psi \psi} C_i, \]

(5)

where $|\partial_i \psi\rangle$ is the derivative of $|\psi\rangle$ with respect to $C_i$ in $[i]$. The effective operator $H_{\psi \psi} = \langle \partial_i \psi | \hat{H} | \partial_i \psi \rangle$ does not depend on $C_i$, and can be calculated recursively using the transfer matrices for $\hat{H}$,

\[ H_{\psi \psi}^i = L_i R_i + 1. \]

Here, $L_i = L_{i-1} T_{A}^i$, $R_i = T_{B}^i R_i + 1$ with $L_0 = R_{L+1} = 1$, $T_{A}^i$ is the left transfer matrix $T_{A}^i = (A_i^\sigma)^\dagger W_i^{\sigma \sigma} A_i^\sigma$, and $T_{B}^i$ is the right transfer matrix $T_{B}^i = (B_i^\sigma)^\dagger W_i^{\sigma \sigma} B_i^\sigma$, for $i = 1, 2, \ldots, L$.

Our zero-site DMRG proposal (DMRG0) is to optimize one tensor $C_i$ at a time [1]. In a DMRG step, the position $i$ and the renormalized operators $L_i$, $R_i + 1$ are fixed while the wavefunction $C_i$ is updated. $E$ and $C_i$ are the lowest eigenvalue and eigenvector, respectively, of the effective Hamiltonian $H_{\psi \psi}$. An iterative eigensolver like Lanczos is used to diagonalize (6) starting from the previous $C_i$ until a given tolerance is reached. The position $i$ is then changed to $i+1$ in (3), performing a matrix decomposition of $M_{i+1}^\dagger = C_i B_i^\sigma = A_i^\sigma C_{i+1}$. An analogous step is performed for the change from $i$ to $i-1$.

As discussed in Sec. I, the main problem to solve in this approach is how to avoid local minima. To this end, we will now present a new enrichment method based on the optimal low-rank correction.

### III. ENRICHMENT VIA OPTIMAL LOW-RANK CORRECTION

The MPS ansatz is highly non-linear in its parameters, the matrices $M_P$. Despite the success of the DMRG proposal to optimize one tensor at a time, there is the danger of being trapped in local minima, especially for single-site effective wavefunction approaches. For our DMRG0, in principle we expect an even worse situation; we will analyze this in examples below in Sec. V. The development of an efficient space enrichment method is then central to the success of DMRG0.

The space enrichment methods [3, 8, 9] are local, i.e. they enrich only one site-tensor at a time. They are based on the application of renormalized operators living on $S$ (of dimension $md$) to the single-site wavefunction of dimensions $md \times m$. This introduces the possibility that the renormalization from $md$ to $m$ changes the wavefunction. We review these approaches in Sec. III A establishing the formal equivalence between [3] and [9]. These ideas, however, are not directly applicable to our DMRG0 because the effective wavefunction $C_i$ is a full-rank $m \times m$ matrix with entries on $S$ and $E$ basis, both of size $m$. In Sec. III B we present our new global proposal for the optimal correction, and two approximate schemes for obtaining it.

#### A. Previous approaches and equivalence

To simplify the explanation, let us focus on the decimation step in DMRG. Given a bipartition $\{S,E\}$ equipped with their respective bases, a state $|\psi\rangle$ corresponds to a matrix $M$. The reduced density matrix for $S$ is $\rho = MM^\dagger$. In the original DMRG [1, 2], $\rho$ is diagonalized $\rho = UDU^\dagger$, its eigenvalues $D$ are truncated to $\bar{D}$ containing the $m$ largest values, and $U$ is truncated to $\bar{U}$ containing the corresponding $m$ eigenvectors, that is, $\rho \approx \bar{U} \bar{D} \bar{U}^\dagger$. In this approximation, the operators in $S$ are renormalized according to $\bar{O} = \bar{U}^\dagger O \bar{U}$, and the states according to $\bar{\psi} = \bar{U}^\dagger \psi$. In particular, $|\psi\rangle$ transforms as $\bar{M} = \bar{U}^\dagger M$.

The first approach to enrich the space was introduced by S. White in [3]. Anticipating later incorporations of relevant states in the environment basis, the density matrix of $S$ is perturbed using the Hamiltonian terms living in $S$,

\[ \bar{\rho} = \rho + \beta^2 L_\gamma \rho (L_\gamma)^\dagger. \]

(7)

This replaces $\rho$ by $\bar{\rho}$ in the decimation step above. Here, the number $\beta^2$ is a small weight tuned by hand, and $L_\gamma$, $R_\gamma$ are the renormalized operators of $S$ ($E$) appearing in the Hamiltonian $\hat{H} = \sum_\gamma L_\gamma \otimes R_\gamma$ for the given bipartition $\{S,E\}$. The resulting density matrix $\bar{\rho}$ is renormalized to keep the largest $m$ eigenvalues.

The second approach [3], as part of the alternating minimal energy (AMEn) algorithm, enriches the space by directly enlarging the wavefunction $M$ to $\bar{M} = (M P)$, where $I$, $0$ are the appropriate identity and null matrices, respectively. Ref. [8] uses $\bar{M}$ to grow the subsystem $S$ basis. In

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1 The original DMRG considers a two-site tensor as the effective wavefunction.
the next step, this allows to choose a richer state by changing the initially vanishing components in (8). Choosing \( P \) as the single-site wavefunction of the (approximate) residual \( \langle H - E \rangle |\psi\rangle \), with \( E = \langle \psi | H | \psi \rangle \), this method guarantees convergence to the global minima [11].

The third approach [9], as part of the DMRG3S algorithm, also uses the subspace expansion technique, and is based on a perturbation of the form

\[
P = P_t M \equiv \beta \left( L_1 M \ L_2 M \ \ldots \ L_p M \right).
\]

It makes a singular value decomposition (SVD) \( \tilde{M} = U_2 \tilde{S} \tilde{V}^\dagger \), followed by a truncation \( \tilde{M} \approx \tilde{U}_2 \tilde{S} \tilde{V}^\dagger \) containing the largest \( m \) singular values of \( s \). Because some reordering of \( \tilde{M} \) takes place during this truncation, the original state is modified/enriched. The basis of \( S \) is rotated with \( \tilde{U}_2 \) and the new state is

\[
\mathcal{R} = \tilde{S} \tilde{V}^\dagger \cdot \begin{pmatrix} I \\ 0 \end{pmatrix}.
\]

Results comparable to [3] are obtained, at a lower computational cost.

Let us discuss the connection between [3] and [9]. For this, we note that the density matrix calculated from \( \tilde{M} \approx \tilde{U}_2 \tilde{S} \tilde{V}^\dagger \) is the same as [7]. Therefore the enrichment steps are equivalent in the \( S \) subsystem. In more detail, since \( \tilde{\rho} = \tilde{M} \cdot \tilde{M}^\dagger \), the diagonalization of \( \tilde{\rho} = U_2 D U_2^\dagger \) can be extracted from the SVD of \( \tilde{M} = U_2 \tilde{S} \tilde{V}^\dagger \) where \( D = \tilde{s}^2 \). The state \( \tilde{M} \) obtained after truncation using \( \tilde{\rho} = U_2^\dagger D U_2 \) is

\[
\tilde{M} = U_2^\dagger M = U_2^\dagger \left( M \ P \right) \cdot \begin{pmatrix} I \\ 0 \end{pmatrix} \\
= U_2^\dagger \tilde{M} \cdot \begin{pmatrix} I \\ 0 \end{pmatrix} \approx U_2^\dagger U_2 \tilde{S} \tilde{V}^\dagger \cdot \begin{pmatrix} I \\ 0 \end{pmatrix} \\
\mathcal{R},
\]

which means that both approaches are equivalent also for the wavefunction within the truncation error. They differ, however, in the representation, and this is responsible for the difference in computational time.

B. Optimal correction

A natural way to enlarge our variational space while keeping the computation tractable is to add a new state \( \psi_\alpha \psi + \beta \bar{\psi} \), \( \alpha, \beta \in \mathbb{R} \),

\[
\alpha \psi + \beta \bar{\psi} : \alpha, \beta \in \mathbb{R}, \tag{12}
\]

where the current (normalized) state \( \psi \) is fixed and \( \bar{\psi} \) is the perturbation. We enrich the space by adding the new direction \( \bar{\psi} \), and determine the coefficients \( \alpha, \beta \) by the Rayleigh-Ritz method. That is, the pair \( \langle \alpha, \beta \rangle \) is the lowest eigenvector of the \( 2 \times 2 \) matrix:

\[
H_{\alpha \beta} = \langle \alpha | \hat{H} | \beta \rangle, \ a, b \in \{ \psi, \bar{\psi} \}.
\]

We note that the alternating linear scheme “ALS(t+z)” method [11], developed to solve linear systems, also enriches the approximate solution \( t \) by adding the residual \( z \).

We propose to obtain the perturbation \( \psi \) by extremizing the energy,

\[
\langle \delta \psi | \left[ \alpha P \hat{H} | \psi \rangle + \beta P (\hat{H} - \lambda) P | \psi \rangle \right] = 0,
\]

Here \( \lambda \) is a Lagrange multiplier coming from the normalization condition. The projector \( P = 1 - |\psi \rangle \langle \psi| \), satisfying \( P \psi = 0 \) and \( P \bar{\psi} \neq \bar{\psi} \), implements the orthogonality condition \( \langle \psi | \psi \rangle = 0 \).

Eq. (14) is the exact condition that determines the optimal correction. It can be cast as an inhomogeneous eigenvalue problem of the form \( A \tilde{x} = \tilde{\lambda} \tilde{x} + \tilde{b} \) for the Hermitian matrix \( A = P \hat{H} P, \tilde{b} = -P \hat{H} | \psi \rangle, \tilde{x} = \beta / \alpha \ P | \psi \rangle \). See e.g. [12] for methods to solve such problems in linear algebra.

We now present two approximate iterative schemes for solving (14): they are motivated by considering a small correction to the current state \( \psi \), but are more broadly applicable. A small correction \( \tilde{\beta} \ll \alpha \) gives rise to a residual (or Lanczos) scheme, while approximating \( \tilde{\lambda} \approx E = \langle \psi | \hat{H} | \psi \rangle \) obtains a Jacobi-Davidson scheme. Both perform very well, and their strengths and weaknesses are inherited from the respective original methods. The Lanczos iteration is fast and straightforward to implement as a direct computation; it is sensitive to the loss of orthogonality, it is problematic when there are (exact or approximate) degenerate states, and it is hard to apply to the interior eigenvalues due to the shift-and-invert mechanism required. On the other hand, the Jacobi-Davidson iteration is determined by a more involved inverse problem, which in turn must be solved iteratively; it is sensitive to the use of preconditioning, but it is very powerful even for degenerate states or interior eigenstates.

1. Lanczos correction

If \( \tilde{\beta} \ll \alpha = \sqrt{1 - \tilde{\beta}^2} \approx 1 - \frac{1}{2} \tilde{\beta}^2 \), we approximate

\[
\langle \delta \psi | \left[ P \hat{H} | \psi \rangle - \lambda \beta P | \psi \rangle \right] = 0,
\]

recalling that at this stage \( \lambda \) is unknown. Eq. (15) implies that \( |\bar{\psi} \rangle \propto |\psi \rangle \langle \psi| \hat{H} | \psi \rangle = (\hat{H} - E) | \psi \rangle \).

The perturbation is then determined by a global residual calculation (16), which is similar to both the ALS(t+z) and the AMEn algorithms.

In our zero-site DMRG framework, numerical experiments show that the above correction (16) implemented as a global correction after each DMRG sweep works well, see for instance Fig. 1. We will choose \( \tilde{m} = m \), but note that one can take even \( \tilde{m} = 2m \), while keeping the global computational cost governed by the optimization step. See Secs. IV and V for more details.

The self-consistency of the residual (16) as an approximate solution to the optimal correction (14) could be checked by direct substitution. However, this is not of our concern here, since the goal is to enrich the space using some well-motivated perturbation.
2. Jacobi-Davidson correction

Eventually, if the residual correction \(16\) becomes insufficient, the following method can be applied. It can be verified that \(\lambda\) represents the energy of the new state \(\alpha \psi + \beta \tilde{\psi}\) when Eq. [14] is solved exactly. Motivated by the Jacobi-Davidson algorithm, we take \(\lambda \approx E = \langle \psi | \hat{H} | \psi \rangle\) in [14], obtaining

\[
|\tilde{\psi}| \propto -[P(\hat{H} - \lambda)P]^{-1} P\hat{H}|\psi|.
\]

(17)

We keep the symbol \(\lambda\) because in our calculation scheme some energy better than \(E\) is usually available. The approximate solution of a linear system like \(12\) has a well established algorithm in the DMRG community, see for instance the calculation of Green’s function response of \(13-15\). In our case, some additional remarks concerning the presence of the projector \(P\) are needed. Ignoring the normalization of \(|\psi|\), and retaking the variational principle \(14\), we obtain

\[
\langle \delta \tilde{\psi} | \hat{H} - \lambda | \psi \rangle = \langle \delta \tilde{\psi} | \hat{H} - \lambda | \psi \rangle - \langle \delta \tilde{\psi} | \hat{H} - \lambda | \psi \rangle + \langle \delta \tilde{\psi} | (E - \lambda) | \psi \rangle = -\langle \delta \tilde{\psi} | P\hat{H}|\psi| \rangle.
\]

As dictated by DMRG, we fix the canonical position \(i\) to find one tensor \((\tilde{C}_i\) of \(|\tilde{\psi}\)\) at a time. In this context, the relation \(|\tilde{\psi}| \equiv |\partial_i \tilde{C}_i\rangle\) implies \(|\partial_i \rangle \equiv |\partial_i \psi\rangle \delta \tilde{C}_i\), and we have the following equation for \(\tilde{C}_i\):

\[
\begin{aligned}
&\{ H_{\psi\tilde{\psi}} | e_i^O \} \langle e_i^H | - | c_i^O \langle e_i^H | c_i^O \rangle \\
&\quad + (E + \lambda) | c_i^O \langle e_i^H | - \lambda \} | \tilde{c}_i = | c_i^H | - E | c_i^O \}
\end{aligned}
\]

(18)

where \(|\chi\rangle\) corresponds to the matrix \(X\) treated as a vector, \(c_i^H = \langle \partial_i \tilde{\psi} | \hat{H} \psi \rangle\), and \(c_i^O = \langle \partial_i \psi \rangle\).

An iterative algorithm can be used to solve \(18\) starting from the previous \(\tilde{C}_i\). For instance, we can apply the generalized minimal residual method (GMRES \(12\)), with computational cost similar to the Lanczos diagonalization of \(6\).

The Jacobi-Davidson correction appears to be more expensive than the residual correction; it would be interesting to perform a comparison of the convergence ratios of both methods.

IV. ALGORITHM

We summarize the previous results in the following algorithm.

Let us call a sweep to the sequence of positions from \(i = 0\) to \(i = L\) (sweeping right) followed by its reverse form \(i = L\) down to \(i = 0\) (sweeping left).

(i) Initialize \(\psi\) with random matrices and canonicalize them to the position \(i = 0\). Set the current error \(\epsilon = 1\) (arbitrarily large).

(ii) Make a standard sweep for \(\langle \psi | \hat{H} | \psi \rangle\) calculating the ground state \(\psi\) using tolerance \(\sim 0.1\epsilon\) to diagonalize each \(H_{\psi\psi}\) in Eq. \(6\).

(iii) Set \(\tilde{\psi} = (H - E)\psi\); starting from the exact MPO-MPS product apply the zip-up algorithm \(16\) to compress \(\tilde{\psi}\) to bond dimension \(\tilde{m}\).

(iv) Update \(\psi\) using the compression of \(\alpha \psi + \beta \tilde{\psi}\) to bond dimension \(m\). Set \(e = E - \lambda_1\) where \(\lambda_1\) is the first eigenvalue of the \(2 \times 2\) matrix \(H\) in \(13\). Since \(\langle \psi | \psi \rangle \neq 0\) the overlap matrix \(O = (a|b)\) with \(a, b \in \{\psi, \tilde{\psi}\}\) should be taken into account, yielding a generalized eigenvalue problem \(Hx = \lambda O \vec{x}\) with Hermitian positive-definite \(2 \times 2\) matrix \(O\).

Steps (ii-iv) are repeated until the energy \(E\) (or \(\lambda_1\)) does not change. Step (ii) is the zero-site DMRG (DMRG0), while (iii-iv) are the basis enrichment steps, in this case, based on the Lanczos (DMRG0-L) or Jacobi-Davidson (DMRG0-JD) correction. Optionally, \(n\) successive perturbations can be applied; the respective algorithms are denoted by DMRG0-Ln and DMRG0-JDn.

As usual, the diagonalization (ii) is the most time-consuming part; its cost per site scales as \(O(2m^3 wK)\), where \(w\) is the MPO bond dimension and \(K\) is the number of eigen-solver iterations. Notice the absence of the physical site dimension \(d\) in this cost. To control the number \(K\) we need both a good starting point (already provided) and we should avoid iterations far beyond the renormalization error of the MPS. Step (iv) provides an appropriate error quantity \(\epsilon\) to ask for during the diagonalization, keeping \(K\) in the order of few tens during the entire calculation. For comparison, the single-site scheme scales as \(O(2m^3 dK)\) with a typical larger value for \(K\) because the local problem is \(d\) times bigger.

Concerning our enrichment proposal, the cost of the Lanczos correction is similar to that of the subspace expansion \(9\) in DMRG3S. It is dominated by the SVD compression of a \(mw \times md\) matrix, which scales as \(O(m^3 w^2 d^2)\). The cost of compressing the sum of two MPSs is negligible \(O(8m^3 d)\). White’s density matrix perturbation \(7\) costs \(O(2m^3 w^3 d^3)\), but it can be reduced to that of DMRG3S if the equivalence of the approaches is taken into account.

The Jacobi-Davidson correction \(18\) is more expensive than the Lanczos one because the GMRES solver required \(K\) iterations, scaling as \(O(2m^3 wK)\), similar to the diagonalization step (ii). However, the missing factor \(d\) can be used to compensate the greater cost of the single-site diagonalization. The advantage is the splitting into smaller problems, which typically decreases \(K\).

Step (iii) can be replaced by \(\tilde{\psi} = \hat{H}\psi\), which mathematically brings us to the same new state Eq. \(12\). We find some cases where, starting from the compression of the exact \(\tilde{\psi} = (H - E)\psi\), the sweeping \(\langle \psi | \hat{H} | \psi \rangle\) setting \(\tilde{\psi} = H\psi\) improves the energy of the compression at step (iv).
V. RESULTS

We benchmark our algorithm with a standard test problem \[^3\] [\[^8\] [\[^9\]]\]: the \(S = 1\) Heisenberg spin chain with \(L = 100\) sites and periodic boundary conditions,

\[
H = \sum_{i=1}^{L} S_i \cdot S_{i+1}.
\]  

(19)

The site dimension corresponds to \(d = 2S + 1 = 3\).

Fig. 2 shows the convergence of the energy \(E\) using DMRG0 and DMRG3S. The top panel presents the methods without enrichment. As expected, DMRG0 gets stuck in a considerably greater energy than DMRG3S does for the same bond dimension \(m\). This is because DMRG0 updates only \(m^2\) parameters for each position \(i\), compared to DMRG3S which updates \(m^2d\). In fact, the updates of DMRG0 do not cover the number of parameters per site \(m^2d\) of the MPS ansatz.

On the other hand, the observed convergence ratios at the bottom panel of Fig. 2 for DMRG0-L and DMRG-JD2 are surprising. Particularly, DMRG0-L makes only \(m^2\) updates per site at step (ii) enriched by a cheap direct residual calculation in step (iii). DMRG-JD2 would cost in principle like DMRG0 but would not cover the number of parameters per site \(m^2d\) of the MPS ansatz.

One of the important features of our method is that the optimization step (ii) does not depend on \(d\), which opens up the possibility of analyzing systems with large \(d\). Although we postpone a more detailed study of this aspect to a future work, let us briefly present results for the Heisenberg model \(^{19}\) with \(S = 3\), namely \(d = 2S + 1 = 7\), and \(S = 5\), i.e. \(d = 11\). The results are shown in Fig. 3. We find that both the Lanczos and the Jacobi-Davidson corrections in DMRG0 outperform the precision of DMRG3S.

VI. CONCLUSIONS AND PERSPECTIVES

In this work, we have presented the zero-site DMRG, a new algorithm to find MPS ground states, with the feature that the local optimization does not depend on the site dimension. We have also proposed a new space enrichment method that avoids local minima and speeds up the convergence ratios to the level of state-of-the-art single-site algorithms. Conceptually, the local optimization of the wavefunction and the en- richment become separate steps.

Both the DMRG0 and the enrichment methods (Lanczos and Jacobi-Davidson) open up the possibility of several develop- ments and extensions. Since the optimization approach is independent of \(d\) (the site physical dimension), DMRG0 could be well-suited to analyze systems with large \(d\). This limit is interesting both theoretically as well as for its applications, such as in the Kondo lattice, dimensional reductions on cylinders, holographic models, etc.

It would also be interesting to investigate in more detail the Lanczos and Jacobi-Davidson methods that we introduced. More nontrivial combinations of these approaches are possible. The Jacobi-Davidson corrections can also be applied to
obtain excited states. The global enrichment could be replaced by a sequential local enrichment similar to [8, 11]. It would also be important to apply this to the single-site scheme.

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