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Efficient variational diagonalization of fully many-body localized Hamiltonians

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We introduce a variational unitary matrix product operator (VUMPO) based variational method that approximately finds *all* the eigenstates of fully many-body localized (fMBL) one-dimensional Hamiltonians. The computational cost of the variational optimization scales linearly with system size for a fixed depth of the UTN ansatz. We demonstrate the usefulness of our approach by considering the Heisenberg chain in a strongly disordered magnetic field for which we compare the approximation to exact diagonalization results.

Introduction: The phenomenon of many-body localization (MBL) generalizes Anderson localization [1] (AL) to interacting systems [2–4]. The pioneering work by Basko, Aleiner and Altshuler [4] showed that the Anderson localized phase is perturbatively stable to small interactions. Quantum manybody systems in the MBL phase have the characterizing property that they do not thermalize [5] and allow to stabilize quantum and topological order which would otherwise melt [6–8]. Following a global quantum quench, MBL phases a show a logarithmic growth of entanglement as a function of time [9–12].

In the Anderson problem the many-body Fock/Slater states constructed from the single particle states have two important features. First, they exhibit an economical description-L single particle states for a system of size L are sufficient to construct all 2^L many-body states. Second, all many-body states exhibit an area law [13-16] for the entanglement entropy stemming from the localized nature of the constituent single particle states. Naturally, attention has focused on what happens to these two properties in the MBL regime. It was noted early on [17] that many-body eigenstates in the MBL regime would have only local entanglement and thus obey the area law. Subsequently numerical studies examined the behavior of the entanglement entropy in detail and demonstrated the area law as well as deviations due to rare regions and states [7, 18, 19]. In another set of papers [20-23] the phenomenology of MBL systems was traced to an emergent set of L commuting local integrals of motion (often called "l-bits") which are believed to exist in fMBL systems-i.e. systems in which all many-body eigenstates are localized.

These two developments invite a natural closure in which the full set of 2^L many-body eigenstates are explicitly constructed from O(L) local ingredients, at least approximately. The well known connection of the area law to matrix-product state (MPS) / tensor network representations of many-body states [24–26] suggests that the latter are the correct language in which to carry out this program. The program has two components: showing that such a compact representation exists and providing a recipe for finding it without recourse to a knowledge of the exact eigenstates, potentially rendering a much larger range of system sizes computationally tractable.

In an important development, two groups have addressed the existence problem. Building on earlier work [27], Pekker and Clark (PC) [28] have shown that the unitary operators that exactly diagonalize fMBL systems can be represented by matrix products operators (MPOs) [25, 29] of bond dimensions that appear to grow very slowly with system size [30] ---in contrast to delocalized systems where the dimension grows exponentially with system size. The slow growth that they do observe is presumably due to rare many-body resonances/Griffiths effects; in its absence, the MPOs would yield the sought after O(L) local description of the full spectrum. Parallel work [31] argued for the congruent result that the presence of local integrals of motion implies the existence of a single "spectral tensor network" that efficiently represents the entire spectrum of energy eigenstates in the fMBL phase. These developments however have not led to an algorithm for finding a compact representation directly without having to diagonalize the full system at a cost that scales exponentially with system size.

In this paper we propose an approach to directly and efficiently find an approximate compact representation of the diagonalizing unitary by using a variational unitary matrix product operator (VUMPO) ansatz. To this end, we construct a cost function whose minimum yields the exact unitary and, hence, the *entire* set of 2^L exact eigenstates of a system of Lqubits. We show that for a fixed "depth" of the approximate \tilde{U} , optimizing the cost-function in d = 1 can be performed at a computational cost that is only *linear* in system size which, in theory, allows us to access system sizes far beyond those possible by ED.

Tensor network notation: For one dimensional systems, an MPS representation of a quantum state living in a basis spanned by L qubits takes the form

$$|\psi\rangle = \sum_{\{\sigma\}} \sum_{0 \le \gamma_i < D} B_{\gamma_1}^{[1]\sigma_1} B_{\gamma_1\gamma_2}^{[2]\sigma_2} \cdots B_{\gamma_{L-1}}^{[L]\sigma_L} |\sigma_1 \cdots \sigma_L\rangle, \quad (1)$$

whereas an MPO representation of an operator in the same Hilbert space takes the form

$$O = \sum_{\substack{0 \le \gamma_i < D\\ \{\boldsymbol{\sigma}\}, \{\boldsymbol{\tau}\}}} A_{\gamma_1}^{[1]\sigma_1, \tau_1} \cdots A_{\gamma_{L-1}}^{[L]\sigma_L, \tau_L} |\sigma_1 \cdots \sigma_L\rangle \langle \tau_1 \cdots \tau_L |,$$

$$(2)$$

where $\sigma_i, \tau_i \in \{\uparrow, \downarrow\}$ and we use a compact notation in which $\boldsymbol{\sigma} = \sigma_1, \sigma_2, \cdots, \sigma_L$ denotes the 2^L states (analogous for $\boldsymbol{\tau}$). Figure 1 shows a pictorial representation of these objects.



FIG. 1. (a) Schematic representation of an MPS representation of a state $|\psi\rangle$. (b) Variational ansatz for the unitary U that encodes *all* eigenstates of a fully many-body localized Hamiltonian. The local unitaries $u_{[n]}^{[m]}$ are parametrized as $u_{[n]}^{[m]} = e^{iS_{[n]}^{[m]}}$ with real symmetric matrices $S_{[n]}^{[m]}$, $n = 1 \dots L - 1$ and $m = 1 \dots N_{\text{layer}}$. (c) Two dimensional generalization of the unitary network.

The MPSs/MPOs are represented by rank three/four tensors $B^{[i]}/A^{[i]}$ on each site *i* (except the first and last tensors which are rank two/three); the external leg(s) σ_i , τ_i refer to the physical spin indices whereas the γ_i are the internal virtual indices that are contracted. Each $B^{[i]\sigma_i}/A^{[i]\sigma_i\tau_i}$ is a D^2 dimensional matrix where *D* is the bond-dimension of the matrix. For two-dimensional systems, the concept of MPS can be extended to projected entangled pair states (PEPS) [32, 33] where the matrices on each site are generalized to higher rank tensors.

Method: We now introduce the VUMPO ansatz and an algorithm to numerically optimize it. Let us assume a onedimensional system and that H is an fMBL Hamiltonian defined on an L-site chain of spin 1/2 operators. It is our goal to find a unitary approximation \tilde{U} of the unitary that diagonalizes the Hamiltonian such that the 2^L eigenstates of H are given by $|\psi_{\tau}\rangle \approx \sum_{\{\sigma\}} \tilde{U}_{\tau}^{\sigma} |\sigma\rangle$. In the parlance of Refs. [20, 21], the physical basis operators σ_i are the "p-bits" wheras the τ_i are the local "l-bits". Each eigenstate is labeled by the occupation of l-bits $\tau = \{\uparrow \uparrow \downarrow \cdots \uparrow\}$, and is obtained by acting with the MPO representation of U on the product state $|\tau\rangle$. In this language of MPOs, it is clear how the 2^L MB eigenstates are constructed from the L matrices $A^{[i]\tau_i}$; further, if the bond-dimension of the matrices scales as O(1) with the system size, the eigenstates are only locally entangled in the p-bit basis and a description of the full eigenbasis in terms of O(L) local ingredients is possible.



FIG. 2. Comparison of the exact energy levels (blue lines) with the ones found by the variational optimization (red lines) for W = 8 and L = 8 as a function of the number of layers of two-site gates. The right panel shows a zoom of some energy levels at the bottom and in the center of the spectrum.

The approximation \tilde{U} is found by numerically minimizing the cost functional

$$f(\{A^{[n]}\}) = \sum_{\{\tau\}} \langle \psi_{\tau} | H^2 | \psi_{\tau} \rangle - \langle \psi_{\tau} | H | \psi_{\tau} \rangle^2 \ge 0, \quad (3)$$

with $\langle \psi_{\tau} | \psi_{\tau'} \rangle = \delta_{\tau,\tau'}$. The cost function is the variance of the energy summed over all approximate MB eigenstates. Naively, one might expect the time to evaluate the cost function Eq. (3) to scale exponentially with the system size L as the sum is performed over 2^L MB eigenstates. However, remarkably, the computation can be performed in a time scaling *linearly* with system size [29]! For example, the term $\sum_{\{\tau\}} \langle \psi_{\tau} | H | \psi_{\tau} \rangle^2$ can be evaluated by "doubling" the degrees of freedom and defining a state $|\phi\rangle = \sum_{\{\tau\}} |\psi_{\tau}\rangle |\psi_{\tau}\rangle |\tau\rangle$. With this notation we find that $\sum_{\{\tau\}} \langle \psi_{\tau} | H | \psi_{\tau} \rangle^2 = \langle \phi | H \otimes H \otimes \mathbb{1} | \phi \rangle$. This expectation value can be efficiently evaluated using the MPO formalism and the most expensive part of the evaluation scales, for a given Hamiltonian in MPO form, as $\propto LD^5$ (see supplemental materials for details). One can now iteratively minimize fby locally optimizing each $A^{[n]}$ using, for example, the conjugate gradient algorithm. In general, an MPO compression of a unitary operator will not strictly respect unitarity. To get a valid positive-definite cost function in these cases, we need to add a Lagrange multiplier to enforce unitarity. In practice, this leads to unstable optimizations.

The key to a stable optimization protocol turns on restricting our algorithm to the manifold of strictly unitary MPOs of a given bond-dimension. To achieve this, we parameterize the VUMPO as a finite depth circuit of two-site unitaries as shown in Fig. 1(b). This Ansatz incorporates two important properties: (i) The VUMPO is unitary for all parameters and (ii) it is local for any finite N_{layer} . We use a single unitary to obtain all eigenstates, but readers will note the obvious connection to the quantum computational notion [34]. Finally, we note that we can rewrite the unitary network as a strictly unitary MPO with bond dimension $D \leq 2^{2N_{\text{layer}}}$, where N_{layer} is the number of layers of two-site gates [35]. However, this step is not necessary and we can evaluate the cost function by directly contracting the unitaries circuit which, in fact, gives a considerable speed up for the systems we consider here. In fact, using the locality of the unitary circuit, the cost function can be evaluated locally and thus it is, in principle, possible to generalize the approach to higher dimensions, e.g., to the two-dimensional square lattice as shown in Fig. 1(c).

The algorithm to find the VUMPO is then similar in spirit to the density matrix renormalization group (DMRG) method [36], except instead of finding the lowest energy state, we minimize the cost function Eq. (3) by sweeping through the local unitaries:

(i) Initialize the local unitaries $u_{[n]}^{[m]} = e^{iS_{[n]}^{[m]}}$ by choosing random symmetric matrices $S_{[n]}^{[m]}$, where $n = 1, 2, \dots L$ and $m = 1, 2, \dots N_{\text{layer}}$.

(ii) Locally minimize the cost function by varying the elements of a given $S_{[n]}^{[m]}$ by using, e.g., a conjugate gradient method.

(iii) Update the network and repeat the previous step for the next unitary.

(iv) Continue the sweeping procedure by minimizing the local unitaries successively until convergence. A full sweep across all the unitaries has to scale as O(L).

We find that the number of steps needed for convergence appears to be approximately independent of L. This gives an overall scaling of the algorithm as $O(LD^5) \sim O(Le^{N_{\text{layer}}})$. Once the algorithm has converged, the VUMPO can be used to obtain all the eigenstates of the system, and to efficiently compute observables using the MPS formalism.

Results: We consider the Heisenberg model with random *z*-directed magnetic fields:

$$H = J \sum_{n} \vec{S}_{n} \cdot \vec{S}_{n+1} - \sum_{n} h_{n} S_{n}^{z}.$$
 (4)

where $\vec{S_n}$ are spin 1/2 operators and the fields h_n are drawn randomly from the interval [-W, W] and we set J = 1. This model has been studied extensively in the context of MBL and several numerical studies strongly suggest that H is fMBL for $W \gtrsim 3.5$ [17, 19].

Energy Spectrum: We begin by comparing the energies obtained using the VUMPO approach with the exact spectrum (full diagonalization). The converged results for W = 8 and L = 8 with different numbers of layers N_{layer} are shown in Fig. 2. For $N_{\text{layer}} = 0$, the VUMPO is the identity (i.e., no variational parameters) and the resulting approximate eigenstates are simple product states of the form $|\sigma_1\rangle|\sigma_2\rangle \dots |\sigma_L\rangle$ with $\sigma_n = \uparrow, \downarrow$. The overall bandwidth in this case agrees relatively well with the exact results because W is the dominant energy scale in the problem. However, as shown in the zoomed in plots, the deviation of individual energy levels is relatively large compared to the mean-level spacing because



FIG. 3. Mean variance of the energy as a function of system size for different number of layers for W = 8. Inset: Mean variance as function of W for a fixed L = 8.



FIG. 4. Comparison of the exact spectral function $A(\omega)$ (black dots) with those obtained using different approximations (see text for details) for L = 10 and W = 8. Spectra are shown using a Lorentzian broadening with an imaginary part of $\epsilon = 0.1$. Inset: Same data with W = 16.

the product states completely neglect local quantum fluctuations which are present in the exact eigenstates. Increasing $N_{\rm layer}$ strongly improves the agreement between the exact and approximate energy levels as the network successively adds entanglement over longer distances.

Next we turn to the mean variance of the energy, which is simply the disorder averaged cost function Eq. (3) divided by 2^L . Figure 3 shows this quantity disorder averaged over 50 realizations as a function of system size for different fixed N_{layer} . We observe a linear increase of the mean variance with system size, and find that the slope decreases as N_{layer} is increased. This tells us that for a given N_{layer} our approximate eigenstates involve a constant error per unit length which decreases as N_{layer} is increased. When reducing the disorder strength W below the MBL transition, the MPO approximation, which is based on the locality of the eigenstates, breaks down (see inset of Fig. 3)

Spectral Functions: To examine the quality of our approx-

imated eigenstates, we use the VUMPO ansatz to obtain the infinite-temperature spectral function

$$A(\omega) = \frac{1}{2^L} \sum_{\{\tau_1\},\{\tau_2\}} |\langle \tau_1 | S_{L/2}^z | \tau_2 \rangle|^2 \delta(\omega - E_{\tau_1} + E_{\tau_2}).$$
(5)

Spectral functions can again be efficiently evaluated using matrix-product techniques and it is also possible to efficiently target different energy densities by considering finitetemperature spectral functions [29, 37]. Figure 4 compares $A(\omega)$ obtained using the VUMPO approach for L = 10 with different disorder strengths and $N_{\text{laver}} = 0, 1, 2$ with the exact results. The large peak at $\omega = 0$ reflects the strongly localized nature of the eigenstates, i.e., the eigenstates of H are close to being eigenstates of local S^z operators. It is interesting to compare the peaks at $\omega > 0$ which are due to local fluctuations in the eigenstates. Clearly, $N_{\text{layer}} = 0$ does not show any features because the VUMPO is diagonal in S^z . When additional layers of unitaries are taken into account, the peak structure of $A(\omega)$ is well approximated. The agreement in both the frequencies and the intensities rapidly improve with increasing N_{laver} , and the results match almost perfectly for W = 16.

Comments on accuracy: We have presented some evidence above for the accuracy of the VUMPO obtained by our method. It remains to establish more precise theorems on what values of N_{laver} it would take to calculate various physical quantities to a specified accuracy. In a step in that direction, PC have looked at the bond dimension D needed to ensure that the smallest singular value in the Schmidt decomposition across any cut in U is less than some fixed ϵ . They found a slow growth of the D_{\min} needed to achieve a desired ϵ with L. In the absence of rare resonances or Griffiths regions, D_{\min} would presumably saturate at a fixed O(1)value for a fixed error density independent of system size implying that we would need only O(1) layers to represent the entire spectrum to the desired accuracy. As it is, with the resonances/Griffiths regions present, D_{\min} is expected to grow as poly(L) whence N_{laver} will grow logarithmically. Let us return to our spectral function computation above but this time we first obtain the exact $2^L \times 2^L$ dimensional unitary that diagonalizes H and then compress it to an MPO of a given bond dimension D (following the prescription outlined in [28]). As seen in Fig. 4 (labeled ED MPO), when compressing $U_{\rm PC}$ to D = 16 (which can exactly represent our $N_{\text{layer}} = 2$ results), the spectral functions $A(\omega)$ are reproduced in a comparable accuracy as in our VUMPO approach.

Summary and discussion: We have introduced an algorithm to find a variational unitary MPO that approximately diagonalizes fully many-body localized Hamiltonians. Our method finds an approximation to all 2^L eigenstates of the Hamiltonian in a time that remarkably scales only linearly with system size! We have benchmarked the method by comparing the results to exact diagonalization for small systems and studied the scaling of the mean variance as a function of system size. For a Heisenberg model in a strongly disordered

field we find good qualitative and quantitative agreement of the obtained energies and spectral functions for a fixed N_{layer} and, importantly, rapid improvement with increasing N_{layer} . With this work we have provided a proof of principle that we can efficiently (i.e, polynomially in system size) perform a variational calculation that provides a complete diagonalization of fMBL systems. As the VUMPO encodes the entire set of eigenstates for fMBL Hamiltonians, many relevant observables such as spectral functions and conductivities can be evaluated efficiently at zero and finite temperatures.

A few comments are in order. First, it is intuitively clear that our VUMPOs should capture most of the structure of the eigenfunctions, or equivalently 1-bits, out to a fixed "lightcone" radius, set by N_{laver} . In terms of the dynamics this should allow accurate inclusion of local excitations on the same length scale and via the recently discussed connection between the energy and size of many-body resonances [38] down to a related frequency scale. Indeed, this feature can be effectively used to study different "slices" of the response function as more layers are added. For example, Figure 4 shows that the exact solution in the case of W = 8 shows certain features at lower frequencies which are absent in the variational solution. Second, for a given VUMPO, one can construct[39] a family of parent Hamiltonians $H = U^{\dagger} H^{\text{diag}} U$ with the same eigenstates by picking different energy distributions for diagonal Hamiltonians in the "l-bit" basis, H^{diag} .

We can visualize many possible avenues for improving our method. Using standard methods [42], a projection onto sectors with fixed magnetization is easily implemented. It is possible to choose the same number of two-qubit gates in a different architecture [40, 41] to get a softer cutoff on the entanglement. More ambitiously we could optimize over *both* the architecture of the unitary network, and the particular gates used. Of course the most desired improvement would be to run at $N_{\text{layer}} \gg 2$ which is currently stymied by the exponential scaling of the cost function. As the diagrams to be contracted now start resembling 2D tensor-network graphs, algorithms from this field could presumably be used to improve the scaling of contraction times.

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