Unitary circuits of finite depth and infinite width from quantum channels

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We introduce an approach to compute the spectra of reduced density matrices for local quantum unitary circuits of finite depth and infinite width. Suppose the time-evolved state under the circuit is a matrix-product state with bond dimension *D*; then the reduced density matrix of a half-infinite system has the same spectrum as an appropriate $D \times D$ matrix acting on an ancilla space. We show that reduced density matrices at different spatial cuts are related by quantum channels acting on the ancilla space. This quantum channel approach allows for efficient numerical evaluation of the entanglement spectrum and Rényi entropies and their spatial fluctuations at finite times in an infinite system. We benchmark our numerical method on random unitary circuits, where many analytic results are available, and also show how our approach analytically recovers the behavior of the kicked Ising model at the self-dual point. We study various properties of the spectra of the reduced density matrices and their spatial fluctuations in both the random and translation-invariant cases.

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I. INTRODUCTION

The dynamics of isolated quantum systems under generic unitary dynamics is one of the basic problems in many-body physics [\[1\]](#page-13-0); despite considerable recent work, many aspects of this problem are not fully understood. An isolated system, evolving under chaotic dynamics from an initial product state, becomes increasingly entangled over time. At sufficiently late times, any finite-size subsystem of an infinite system is well described by a thermal reduced density matrix, provided the system obeys the eigenstate thermalization hypothesis (ETH) [\[2–4\]](#page-13-0); this approach to a thermal local-density matrix is called "thermalization." There is considerable numerical evidence that generic many-body systems obey ETH [\[5\]](#page-13-0).

We are concerned with the dynamics of the reduced density matrix *before* the system has fully thermalized. To quantify the thermalization time scale more precisely, recall that the Rényi entropies of a subsystem *A* are defined in terms of the reduced density matrix ρ*^A* of *A* as

$$
S_A^{(n)}(t) = \frac{1}{1-n} \ln \text{tr}[\rho_A(t)^n],\tag{1}
$$

where *n* is called the Rényi index. The Rényi entropies fully characterize the spectrum of ρ*A*.

The general consensus $[6-10]$ $[6-10]$ is that—unless a system experiences many-body localization [\[11\]](#page-14-0)—the entropies initially obey $S_A^{(n)}(t) \sim v_n t$, increasing linearly with time with a growth rate *vⁿ* that depends on the Rényi index *n*. (However, recent results suggest that for $n > 1$ the growth is sublinear for generic initial states in the presence of conservation laws [\[12,13\]](#page-14-0).) The implication for the spectrum of the reduced density matrix is as follows. Parametrizing the eigenvalues of ρ_A in terms of an "entanglement energy" as $\lambda_i = e^{-\epsilon_i}$ [\[14,15\]](#page-14-0),

and introducing the "density of states" $\rho(\epsilon)$, we have

$$
S_A^{(n)} = \frac{1}{1-n} \ln \left[\int d\epsilon \, \varrho(\epsilon) e^{-n\epsilon} \right].
$$
 (2)

The behavior $S_A^{(n)}(t) \sim v_n t$ is then consistent with a density $\varrho(\epsilon)$ having the large deviation form

$$
\varrho(\epsilon) \sim \exp[t\pi(\epsilon/t)],\tag{3}
$$

for some function $\pi(\eta)$. In the saddle-point approximation we find

$$
v_n = \frac{S_A^{(n)}}{t} = \frac{\pi(\eta_n) - \eta_n}{1 - n},
$$
 (4)

where η_n is determined by $n = \pi'(\eta_n)$. The growth rates v_n are seen to be related to the function $p(\eta)$ describing the density of states of the entanglement spectrum by Legendre transformation. Note that the numerator in Eq. (4) vanishes at $n = 1$ due to the normalization of the density matrix, yielding a finite growth rate v_1 for the von Neumann entropy $S_A^{(1)}$. Rényi entropies with smaller *n* grow faster; the reduced density matrix for a subsystem becomes thermal when the slowest Rényi entropy, the so-called "min-entropy" $S_{\infty} \equiv \min_i(\epsilon_i)$ (i.e., the largest entanglement eigenvalue) has saturated.

As well as the rates v_n , the time evolution is characterized by the butterfly velocity v_B at which local perturbations spread [\[8](#page-13-0)[–10,16–18\]](#page-14-0). Although the various velocities are generically separate, they coincide in exactly solvable models (such as random circuits in the limit of large local Hilbert space dimension or Clifford gates [\[8](#page-13-0)[–10\]](#page-14-0) and the self-dual kicked Ising model [\[19,20\]](#page-14-0)), so the entanglement spectrum evolves in a trivial way. Away from these nongeneric limits, little is known analytically about the entanglement spectrum. A few Rényi

FIG. 1. A depth $d = 4$ unitary circuit of the type considered in this work.

entropies can be explicitly computed by mapping the circuit dynamics to random classical partition functions [\[9,10,18\]](#page-14-0) but these mappings do not yield the full entanglement spectrum. The picture that emerges from numerical studies is, however, that the entanglement spectrum has nontrivial structure in generic systems, such as a bandwidth that widens linearly in time; this feature is absent in the exactly solvable limits [\[21\]](#page-14-0). However, this structure is not fully understood at present.

In the present work we develop and apply a numerical transfer-matrix approach to compute the structure of the entanglement spectrum for spatially infinite systems at early times. This approach allows us to access some aspects of entanglement for larger subsystems than were studied in previous work: for instance, we are able to compute the spatial fluctuations of entanglement in systems of 10 000 sites. Our approach works formally with infinite systems; we assume that the system of interest has been initialized in a product state and then subjected to a finite-depth quantum circuit (i.e., evolution for a finite time) consisting of on-site or nearest-neighbor quantum gates (see Fig. 1). We compute the spectrum of the reduced density matrix of a bipartition into two semi-infinite regions at an arbitrary point. After applying a quantum circuit of finite depth *t*, the reduced density matrix has rank q^{t-1} . We will see that the spectrum of the reduced density matrix can be interpreted as that of a $q^{t-1} \times q^{t-1}$ matrix *R* acting on an ancilla space. In addition, reduced density matrices across adjacent cuts are related by quantum channels that are straightforward to construct given the circuit. These quantum channels act as transfer matrices for the entanglement spectrum [\[22\]](#page-14-0).

The quantum-channel perspective is helpful for a number of reasons. First, using this method one can compute the largest few eigenvalues of the entanglement spectrum for circuits that are too deep to permit direct simulation. Second, as the transfer matrix acts on formally infinite systems, spatial fluctuations of entanglement can be directly studied. Third, standard methods from quantum optics such as the stochastic unraveling of quantum channels [\[23\]](#page-14-0) can be applied to simulate the dynamics of entanglement on larger scales than direct simulation permits. Finally, for translation-invariant initial states evolving under translation-invariant circuits, one can iterate the quantum channel until it converges, and thus extract the entanglement spectrum, free of finite-size effects, without incurring the computational overhead of time-evolving a large system.

These are the issues we explore in the present work. Our main results are as follows. First, we provide algorithms based on quantum channels for computing the entanglement spectrum and its low-energy tail, as well as for computing some Rényi entropies, by propagating the quantum channel in an ancilla space. For the exactly solvable model of Ref. [\[20\]](#page-14-0) we analytically demonstrate that the entanglement spectrum is trivial, using the properties of the associated quantum channel. We compute the behavior of the "low-energy" (large Schmidt rank) tail of the entanglement spectrum for random unitary circuits, random circuits with a conservation law, and translation-invariant integrable circuits, acting on various initial states. We compute the distributions of the purity and of the min-entropy; for random circuits we find that both the second Rényi entropy and the min-entropy follow Gaussian distributions at the accessible circuit depths (the purity, therefore, follows a log-normal distribution). For circuits with conservation laws or translation-invariant circuits, the nature of this low-entanglement-energy tail is sensitive to the fluctuations in the initial state. We compute the spatial correlations of entanglement, and find that their correlation length grows sublinearly in time. However, the correlation lengths are short at the accessible times and we are not able to identify a definite exponent.

This paper is organized as follows. In Sec. II we briefly review concepts such as unitary circuits, matrix-product states, and quantum channels, as they apply to the algorithms introduced here. In Sec. [III](#page-3-0) we describe how to construct quantum channels for unitary circuits, and estimate the complexity of various exact and approximate methods for extracting the entanglement spectrum (or some of its moments). In Sec. [IV](#page-6-0) we explicitly compute the transfer matrix for the self-dual kicked Ising model, and confirm that all the Rényi entropies coincide in this model, as they are known to [\[19,20\]](#page-14-0). In Sec. [V](#page-8-0) we present results for the entanglement spectrum, the distributions of purity and min-entropy, and the growth of spatial correlations in the entanglement, in random unitary circuits, and some variants of these. Finally Sec. [VI](#page-13-0) summarizes our results and discusses future directions.

II. BACKGROUND

A. Quantum circuits

In this work we consider systems that evolve under the application of discrete local unitary gates tiled in the pattern shown in Fig. 1. Our approach applies both to Floquet systems in which the gates are applied periodically in time, and to random circuits where each gate is drawn independently and randomly. As with the TEBD algorithm [\[24\]](#page-14-0), it can be extended to continuous time evolution under strictly local Hamiltonians by discretizing the time evolution, e.g., through a Suzuki-Trotter decomposition.

B. Matrix product states

In this section we introduce some background material on matrix product states (MPS's), which will form an essential part of the following development. More detailed expositions may be found in Refs. [\[25–27\]](#page-14-0).

We consider a quantum system described by *N* identical subsystems with finite Hilbert space dimension *q*. The (pure) quantum states of the system are therefore defined by vectors in the Hilbert space,

$$
\mathcal{H}_N \equiv \overbrace{\mathbb{C}^q \otimes \cdots \otimes \mathbb{C}^q}^{N \text{ times}}.
$$
 (5)

A basis of orthonormal product states has the form

$$
|s_{1:N}\rangle = |s_1\rangle_q \otimes |s_2\rangle_q \cdots \otimes |s_N\rangle_q, \qquad (6)
$$

where $|s\rangle_q$ $s = 1, \ldots q$ is an orthonormal basis for \mathbb{C}^q , and we have introduced the sequence notation $s_{1:N} \equiv s_1, s_2, \ldots s_N$. By taking components of a vector $|\Psi\rangle \in \mathcal{H}$,

$$
\Psi_{s_{1:N}} = \langle s_{1:N} | \Psi \rangle \,, \tag{7}
$$

 $|\Psi\rangle$ can be regarded as a rank-*N* tensor with components $\Psi_{s_1 \ldots s_n}$. An MPS is a tensor of the form

$$
\Psi_{s_{1:N}} = A_{s_1}^{(1)} A_{s_2}^{(2)} \cdots A_{s_N}^{(N)},\tag{8}
$$

where $A_s^{(1)}, \ldots, A_s^{(N)} \in \mathbb{C}^{D_j \times D_{j+1}}$ are matrices, and the numbers D_j $j = 1, \ldots N + 1$ are known as the *bond dimensions*, with $D_1 = D_{N+1} = 1$. Thus the product Eq. (8) has the form "row vector, product of matrices, column vector," and yields a complex number for each sequence s_1 : s_N .

An arbitrary vector $|\Psi\rangle$ may be approximated by an MPS with an error that decreases as $D \equiv \max_j D_j$ increases. We will see that the state arising from applying a unitary circuit to a product state is exactly given by an MPS with $D = q^{d-1}$, where *d* is the depth of the circuit.

A graphical notation for MPS proves to be extremely convenient, and is discussed extensively in Ref. [\[26\]](#page-14-0). In this notation tensors—such as the matrices or vectors $A^{(j)}$ —are represented as boxes, with the number of lines or edges entering a box indicating the number of indices the object bears. An edge joining two vertices indicates the (pairwise) contraction of an index. Thus the MPS in Eq. (8) is denoted

Ψ^s1:*^N* = A(1) s1 A(2) s2 ... A(N) sN

Here, $A^{(2)}$ has three lines attached because the collection of matrices $A_s^{(2)} \in \mathbb{C}^{D \times D}$ may be regarded as rank-3 tensor $A^{(2)} \in$ $\mathbb{C}^{D \times D \times q}$. The vertical leg represents the indices s_i that live in the "physical space" while the horizontal lines represent indices in the "bond space" (which we shall also refer to as the "ancilla space").

The squared norm of a vector $|\Psi\rangle$ may be calculated by contracting all physical indices between $\Psi_{s_{1:N}}$ and $\bar{\Psi}_{s_{1:N}}$. This has the graphical representation:

$$
\langle \Psi | \Psi \rangle = \sum_{s_{1:N}} \bar{\Psi}_{s_{1:N}} \Psi_{s_{1:N}} = \underbrace{\begin{pmatrix} A^{(1)} & A^{(2)} & A^{(N)} \\ \hline A^{(1)} & A^{(2)} & A^{(N)} \end{pmatrix}}_{\text{(9)}}
$$

More generally, the reduced density matrix for the leftmost *n* subsystems that arises from a pure state by tracing over the remaining subsystems is denoted

$$
\rho_{s_{1:n},s'_{1:n}} \equiv \sum_{s_{n+1:N}} \bar{\Psi}_{s_{1:n}s_{n+1:N}} \Psi_{s'_{1:n}s_{n+1:N}} =
$$

 (10)

where for simplicity we denote $A^{(j)}$ by *j* and $\overline{A}^{(j)}$ by \overline{j} . The above expressions may also be written

where the Hermitian matrices $R^{(j)} \in \mathbb{C}^{D_{j+1} \times D_{j+1}}$ are defined by

$$
R^{(j-1)} = \sum_{s} A_s^{(j)} R^{(j)} A_s^{(j)\dagger}, \quad j = n+1, \dots N, \qquad (12)
$$

and $R^{(N)} = 1$.

The MPS representation of $|\Psi\rangle$ has a redundancy sometimes referred to as "gauge freedom." For a set X_i $j =$ $1, \ldots N-1$ of invertible matrices, the transformation

Canonical forms

$$
A_s^{(1)} \to A_s^{(1)} X_1^{-1},
$$

\n
$$
A_s^{(j)} \to X_{j-1} A_s^{(j)} X_j^{-1}, \quad j = 2, \dots N - 1,
$$

\n
$$
A_s^{(N)} \to X_{N-1} A_s^{(N)}
$$
\n(13)

leaves $\Psi_{s_{1:N}}$ unchanged. Further conditions may be imposed to reduce this redundancy [\[25\]](#page-14-0). We will be concerned with matrices in the *left canonical* form, satisfying the condition

$$
\sum_{s} A_{s}^{(j)\dagger} A_{s}^{(j)} = \mathbb{1}_{D_{j+1}}
$$
 (14)

$$
\begin{pmatrix}\nA^{(j)} \\
\vdots \\
\overline{A^{(j)}}\n\end{pmatrix} = \begin{pmatrix}\n\cdot & & & (15)\n\end{pmatrix}
$$

Note that $D_2 = q$ is necessary for $A^{(1)}$ to be placed in the left canonical form, but this places no restriction on the state.

Analogously, matrices in the *right* canonical form satisfy

$$
\sum_{s} A_{s}^{(j)} A_{s}^{(j)\dagger} = \mathbb{1}_{D_{j}}
$$
(16)

$$
\overline{A_{j}^{(j)}} = \overline{A_{j}}
$$
(17)

(requiring $D_N = q$).

One benefit of the canonical forms is that they may be contracted "automatically." For example, in terms of an MPS in the right canonical form the reduced density matrix $\rho_{s_{1:n}, s'_{1:n}}$ in Eq. (10) takes the form

$$
\rho_{s_{1:n},s'_{1:n}} = \begin{array}{c}\n1 \\
\downarrow \\
s_1 \\
s'_1 \\
\hline\n\vdots \\
s_n\n\end{array} \cdots \begin{array}{c}\n\downarrow \\
s_n \\
s'_n \\
\hline\n\vdots \\
s_n\n\end{array} \qquad (18)
$$

A second benefit—which is more relevant for us—is that the spectrum of the reduced density matrix $\rho_{s_{1:n}, s'_{1:n}}$ coincides with the spectrum of $R^{(n)}$ for an MPS in the *left* canonical form. This may be seen by introducing the spectral representation

$$
R^{(n)} = \sum_{\alpha} \lambda_{\alpha} r_{\alpha} r_{\alpha}^{\dagger}, \qquad (19)
$$

in terms of the eigenvalues λ_{α} and eigenvectors r_{α} of $R^{(n)}$. Upon substitution into Eq. (11) this yields a spectral representation for $\rho_{s_{1:n}, s'_{1:n}}$,

$$
\rho_{s_{1:n}, s'_{1:n}} = \sum_{\alpha} \lambda_{\alpha} \begin{array}{ccc} \boxed{1} & & \boxed{n} & \boxed{r_{\alpha}} \\ \hline s_1^{\dagger} & \cdots & s_n^{\dagger} & \\ \boxed{1} & & \boxed{n} & \boxed{\bar{r}_{\alpha}} \end{array}, \qquad (20)
$$

in terms of vectors in \mathcal{H}_n that are orthonormal by the left canonical condition [\[25\]](#page-14-0).

A more direct way of seeing that the spectra of $R^{(n)}$ and ρ_A coincide, for MPS's in the left canonical form, is shown in Fig. 2. The moments of the spectrum are given by traces over matrix products of the form ρ_A^k ; one can use the left canonical property [\(14\)](#page-2-0) to eliminate the "boxes" pairwise and arrive at the result tr $\rho_A^k = \text{tr}[(R^{(n)})^k]$. Since all moments coincide, ρ_A and $R^{(n)}$ must have the same spectrum.

FIG. 2. Left: the reduced density matrix ρ_A can be expressed as a matrix product operator involving matrices in left canonical form (squares), and a matrix R that comes from the complement, \overline{A} . Rényi entropies are proportional to tr ρ_A^n ; when the MPS is in left canonical form, tr $\rho_A^n = \text{tr } R^n$. Right: application of this idea to a unitary circuit. The matrix-product state $|\psi(t)\rangle$ is constructed by cutting diagonally through the circuit (solid lines); the leftover gates in the shaded region form the matrix *R*.

C. Quantum channels

The above formalism has a natural interpretation in terms of quantum channels, or completely positive trace preserving (CPTP) maps $[25]$. In this interpretation the matrices $R^{(j)}$ are regarded as a sequence of density matrices that represent mixed states in the bond space. The definition of $R^{(j)}$ given in Eq. [\(12\)](#page-2-0) guarantees that the maps from one $R^{(j)}$ to the next are completely positive (Choi's theorem), while the left canonical condition, Eq. [\(14\)](#page-2-0), ensures that they are trace preserving. Therefore, contracting a physical leg (i.e., moving the entanglement cut in real space) amounts to applying a CPTP map to the ancilla. In this context the $A_s^{(j)}$ are known as Kraus operators.

From now on we assume without loss of generality that all $D_j = D$, which may be achieved by padding the matrices with zeros. Square matrices $A_s^{(j)} \in \mathbb{C}^{D \times D}$ satisfying the left canonical condition may be parametrized in terms of unitary matrices $U_j \in \mathcal{U}(qD)$ as

$$
A_{s,ab}^{(j)} = \langle s|_q \langle a|_D U_j |0\rangle_q |b\rangle_D. \tag{21}
$$

Physically, this corresponds to the amplitude for the following process: prepare the physical subsystem *j* in a fixed state $|0\rangle_a$, and then act on the state $|0\rangle_q |b\rangle_p$ of the subsystem *j* and ancilla with a unitary, arriving in state $|s\rangle_q |a\rangle_p$. While any CPTP map may be presented in this form [\[28\]](#page-14-0), we will see that this is precisely how quantum channels arise in the case of unitary circuits.

III. QUANTUM CHANNELS FROM UNITARY CIRCUITS

This section is organized as follows. We first discuss how to slice up a unitary circuit into an MPS where all matrices are in left canonical form. This immediately gives us a quantum channel that propagates the ancilla-space density matrix *R* in the spatial direction. We then discuss two controlled but approximate ways of implementing the quantum channel for larger subsystems: first, an approach for computing the entanglement spectrum based on approximating the ancillaspace density matrix *R* as a lower-rank object, and second, an approach for computing the purity, specifically, by a stochastic unraveling of the quantum channel (i.e., by sampling "quan-tum trajectories" in ancilla space [\[23\]](#page-14-0)).

A. From unitary circuits to canonical-form MPS's

We now introduce the main idea behind our approach. A planar unitary circuit that starts from a product state may be presented as an MPS by slicing it into strips in an arbitrary way. Each slice *j* is associated with q^2 matrices $A_{s_1,s_2}^{(j)}$ indexed by two physical indices. The dimension of the ancilla is $D =$ *q*^{*d*−1}. For a general decomposition of a circuit, the resulting matrices $A_{s_1,s_2}^{(j)}$ will not be in the appropriate canonical form, so the spectra of $R^{(j)}$ and ρ_A will not coincide. An MPS in left canonical form *is* obtained for slices along the southwest to northeast diagonal,

A graphical proof is straightforward, since

The unitarity of the constituent blocks

$$
\begin{vmatrix}\nb'_1 & b'_2 & b'_1 & b'_2 \\
\hline\n\frac{U_1^{\dagger}}{U_1} & & & \\
b_1 & b_2 & b_1 & b_2\n\end{vmatrix} = \delta_{b_1, b'_1} \delta_{b_2, b'_2}
$$
 (24)

guarantees that the left canonical condition

$$
\sum_{s_1, s_2} A_{s_1, s_2}^{(j)\dagger} A_{s_1, s_2}^{(j)} = \mathbb{1}_D
$$
 (25)

is satisfied. Choosing the southeast to northwest diagonal gives an MPS in the right canonical form.

B. Applying the quantum channel

We have shown that a unitary circuit may be represented as an MPS in the left canonical form. This observation may be used to efficiently compute the ancilla density matrices $R^{(j)}$, whose spectrum coincides with that of the reduced density matrix $\rho_{s_1, j, s'_1, j'}$. We start with an arbitrary ancilla density matrix R_0 , and apply the channel until convergence. In practice we find that convergence is fast, and the ensemble converges to its stationary form after ∼*t* iterations, where *t* is the depth of the circuit. This rate does not seem to depend appreciably on the initial conditions. The convergence of the channel is related to edge effects of the entanglement spectrum as one approaches a boundary. At time *t*, regions in the interior of the system, for which $L > t$, will not see the edges by causality, so their entanglement spectra should not have edge effects. Beyond this point, we do not have a detailed understanding of the convergence properties of these quantum channels; developing this is an interesting topic for future work.

Recall that the $R^{(j)}$ are defined by

$$
R^{(j-1)} = \sum_{s_1, s_2} A^{(j)}_{s_1, s_2} R^{(j)} A^{(j)\dagger}_{s_1, s_2}, \quad j = n+1, \dots N. \tag{26}
$$

After cutting along the SW-NE diagonal, $R_{a_{1:d-1}, a'_{1:d-1}}^{(j-1)}$ has the graphical representation

This expression may be simplified somewhat by noting that the topmost unitary $(U_4$ in the above example) may be eliminated to give

The algorithm for applying the quantum channel is therefore as follows:

(1) Trace over the first index of $R^{(j)}$,

$$
R_{a_{1:D},a'_{1:D}}^{(j)} \to \sum_{a} R_{aa_{1:D-1},aa'_{1:D-1}}^{(j)}.
$$
 (29)

This reduces the number of indices of $R^{(j)}$ to 2(*t* − 2), or *q*^{2(*t*−2)} components.

(2) Apply the unitaries U_{t-1} and U_{t-1}^{\dagger} . This increases the rank of the resulting tensor back to $2(d - 1)$.

(3) Continue applying unitaries from the "middle out" for $j = t - 2, \ldots 2.$

(4) Apply U_1 and U_1^{\dagger} , with the outer indices fixed.

This process involves $O(t)$ steps of matrix multiplication, where the matrices are of size $O(q^t)$. Thus if one directly applies the channel to a density matrix, the overall complexity is $O(tq^{2(t-1)})$. This is better than the naive $O(tq^{3(t-1)})$ because each of the unitaries that make up the channel is a sparse

matrix. A Python implementation of the algorithm is available [\[29\]](#page-14-0).

Since our quantum channel is constructed from a diagonal cut through the unitary circuit, there will be edge effects in a rectangular circuit of finite width. Our approach is well suited to *infinite* width circuits: the channel is applied repeatedly to a random initial density matrix until a steady-state density matrix is approached (for translationally invariant circuits) or a stationary distribution (for random circuits). This typically occurs on the scale of a number of steps roughly equal to the depth of the circuit.

C. Low-rank approximation of $R^{(n)}$

Away from fine-tuned points (see Sec. [IV\)](#page-6-0), finite-depth local unitary circuits give rise to entanglement spectra that are very broad; thus, the vast majority of the eigenstates of ρ_A are close to zero and do not contribute to Rényi entropies with $n \geq 1$ [\[21\]](#page-14-0). This observation is implicit in the fact that different S_n have different growth rates [\[18\]](#page-14-0). This fact allows us to propagate $R^{(n)}$ with negligible error using far fewer than *q^{t-1}* basis states. We proceed as follows. We approximate

$$
R^{(n)} \approx \sum_{k=1}^{K} \lambda_k |k\rangle\langle k|,\tag{30}
$$

where λ_k are the *K* largest eigenvalues of *R* and $|k\rangle$ are the associated eigenvectors. We renormalize all the eigenvalues to preserve the trace. We now evolve each $|k\rangle$ under each "leg" of the quantum channel. This evolution is efficient because each of the unitaries is a very sparse matrix. At the end of this process we have the expression

$$
R^{(n+1)} \approx \sum_{k=1}^{K} \sum_{i=1}^{q^2} \lambda_k |\phi_{ik}\rangle \langle \phi_{ik}|,
$$
 (31)

where $|\phi_{ik}\rangle$ are not mutually orthogonal or normalized, but nevertheless span a Kq^2 -dimensional space. Equation (31) is the ancilla density matrix that would result from one step of the quantum channel applied to the approximate density matrix (30). It is a legitimate density matrix, since Eq. (30) was. Now we can repeat this process by approximating $R^{(n+1)}$ with its top *K* eigenvectors, renormalizing, propagating, and so on.

When K is sufficiently small, the complexity of evolving the *K* top eigenvectors scales as $O(Ktq^{t-1})$, since the unitary gates are individually sparse matrices. The diagonalization step scales as $O(K^3)$, meanwhile. Benchmarking our results against exact diagonalization at small sizes (and against exact results for random unitary circuits at arbitrary sizes) we find that keeping ≤ 100 eigenvectors suffices to capture the quantities that are of interest here—mainly, the purity and min-entropy, and their fluctuations.

D. Trajectory approach for computing the purity

In this section we describe an approach based on mapping *vectors* in the ancilla space rather than density matrices. Formally, this method is equivalent to the trajectory approach developed to analyze master equations in quantum optics [\[23\]](#page-14-0). Applying unitaries to a vector is an $O(dq^{(d-1)})$ operation because the unitaries are sparse. While such an approach is evidently attractive, we will see that there is a trade-off in terms of the number of times the matrices $A_{s_1,s_2}^{(j)}$ must be applied.

The ancilla density matrix can be defined in terms of averages over trajectories in the physical indices in the following way. Starting from the Kraus form,

$$
R^{(j)} = \sum_{s} A_s^{(j)} R^{(j-1)} A_s^{(j)\dagger}, \quad j = n+1, \dots N; \tag{32}
$$

in the case of unitary circuits the index *s* is a composite, $s = (s_1, s_2)$, and we have changed the indexing of slices so that indices increase going right to left. We see that *L* updates correspond to summing over trajectories in the physical indices of length *L*

$$
R^{(L)} = \sum_{s_1:s_L \in \{\mathbb{Z}_q\}^L} A_{s_L}^{(L)} \cdots A_{s_1}^{(1)} R^{(0)} A_{s_1}^{(1)\dagger} \cdots A_{s_L}^{(L)\dagger}.
$$
 (33)

If we start from a pure state $R^{(0)} = |\psi_0\rangle \langle \psi_0|$ we can write this as an average over trajectories with uniform distribution,

$$
R^{(L)} = q^L \mathop{\mathbb{E}}_{s_{1:L} \sim \text{uniform}} \left[\left| \tilde{\psi}_{s_{1:L}} \right\rangle \left\langle \tilde{\psi}_{s_{1:L}} \right| \right],\tag{34}
$$

where the vectors $|\psi_{s_1} \rangle$ are defined as

$$
\left| \tilde{\psi}_{s_{1:L}} \right\rangle = A_{s_L}^{(L)} \cdots A_{s_1}^{(1)} \left| \psi_0 \right\rangle. \tag{35}
$$

These vectors are unnormalized. The normalization factors

$$
p(s_1 : s_L) \equiv \langle \tilde{\psi}_{s_{1:L}} | \tilde{\psi}_{s_{1:L}} \rangle \tag{36}
$$

are a normalized probability distribution over trajectories by virtue of the left canonical condition Eq. [\(14\)](#page-2-0). Denoting the normalized vectors as $|\psi_{s+1}\rangle$ we can express the ancilla density matrix as

$$
R^{(L)} = \mathop{\mathbb{E}}_{s_{1:N} \sim p(\cdot)} [|\psi_{s_{1:L}}\rangle \langle \psi_{s_{1:L}}|]. \tag{37}
$$

As a trajectory increases in length, the normalization factors are updated according to

$$
p(s_1 : s_L) = p(s_1 : s_{L-1}) \langle \psi_{L-1} | A_{s_L}^{(L) \dagger} A_{s_L}^{(L)} | \psi_{L-1} \rangle, \qquad (38)
$$

so that the second factor may be interpreted as a conditional probability

$$
p(s_L|s_{1:L-1}) = \langle \psi_{L-1} | A_{s_L}^{(L)\dagger} A_{s_L}^{(L)} | \psi_{L-1} \rangle, \qquad (39)
$$

Equation (37) expresses the ancilla density matrix in terms of vectors, but it requires an average over trajectories. The downside of this approach is that evaluating *R* will require roughly γ^{-1} trajectories, where γ is the purity of a halfinfinite region, which sets the approximate rank of the reduced density matrix. However, this approach lends itself to parallelization while the channel based approach does not.

For a translationally invariant system, and assuming this random process is ergodic, we can substitute an average over the length of single long trajectory in the $L \to \infty$ limit,

$$
R = \lim_{L \to \infty} \frac{1}{L} \sum_{l=1}^{L} |\psi_{s_{1:l}}\rangle \langle \psi_{s_{1:l}}| \,. \tag{40}
$$

In practice, long runs and multiple trajectories are used.

 $O(D)$ evaluation of matrix products is not much use if we still need $O(D^3)$ evaluation of the spectrum of $R^{(L)}$ or $O(D^2)$ evaluation of the purity. However, we can access the purity without dealing with $R^{(L)}$ directly using

$$
\gamma_L = \text{tr}[R^{(L)2}] = \mathop{\mathbb{E}}_{s_{1:N}, t_{1:N} \sim p(\cdot)} |\langle \psi_{t_{1:L}} | \psi_{s_{1:L}} \rangle|^2, \tag{41}
$$

which follows from Eq. (37) . This formula expresses the purity as the average fidelity over pairs of trajectories. In a high-purity state the ancilla vectors stay close to each other as they evolve over different trajectories, whereas in a highly entangled state different trajectories explore different regions of ancilla space.

Note that the expectations discussed in this section are unrelated to any random variables that may form part of the specification of the circuit. Evaluating the average purity, for example, would require an additional average of Eq. (41) over these variables.

IV. EXACTLY SOLVABLE EXAMPLE: SELF-DUAL KICKED ISING MODEL

In order to illustrate the utility of the formalism introduced in the previous section, we now turn to an example of a unitary circuit in which the entanglement spectrum can be determined analytically. This is the kicked Ising model at the self-dual point discussed in two recent papers [\[19,20\]](#page-14-0). The kicked Ising model describes the evolution of a system of *L* spin-1/2 subsystems (qubits) for an integer time *t* by the unitary operator $(U_{\text{KI}})^t$, where $U_{\text{KI}} = K I_h$ is composed of the two unitaries

$$
I_{\mathbf{h}} = e^{-iH_{\rm I}[\mathbf{h}]}, \quad K = e^{-iH_{\rm K}}, \tag{42}
$$

where

$$
H_{\rm I}[\mathbf{h}] = \sum_{j=1}^{L} [JZ_j Z_{j+1} + h_j Z_j],\tag{43}
$$

$$
H_{\mathcal{K}} = b \sum_{j=1}^{L} X_j,
$$
\n⁽⁴⁴⁾

and (X_i, Y_i, Z_i) are the Pauli matrices for spin *j. H*_I[**h**] is the classical Ising model with arbitrary longitudinal fields h_i , while H_K describes a transverse field.

In Ref. [\[20\]](#page-14-0) the growth of the entanglement entropies was found exactly for some particular initial product states at the special "self-dual" values

$$
|J| = |b| = \frac{\pi}{4}.
$$
 (45)

For a region *A* of size *N*, the authors found that when starting from an arbitrary product state in the Z_i basis the Rényi entropies [\(1\)](#page-0-0) need to be *exactly* given by

$$
\lim_{L \to \infty} S_A^{(n)}(t) = \min(2t - 2, N)\ln 2,\tag{46}
$$

independent of Rényi index *n*. The interpretation in terms of the entanglement spectrum is striking: there are 2min(2*t*−2,*N*) eigenvalues equal to 2[−] min(2*t*−2,*N*) and the rest are zero.

We now show how our quantum channel approach may be used to derive the corresponding result for the case of a semiinfinite interval,

$$
\lim_{L \to \infty} S_A^{(n)}(t) = (t - 1)\ln 2.
$$
 (47)

We can present the unitary $(U_{\text{KI}})^t$ as a unitary circuit of depth *t*, where the layers alternate between unitaries operating between spins $2j - 1$ and $2j$, and between spins 2*j* and $2j + 1$. A variety of decompositions are available. For reasons that will become clear, we choose the following:

where the one-qubit (K) and two-qubit (I) gates have the form

$$
\mathcal{K} = \exp[-ibX],\tag{49}
$$

$$
\mathcal{I} = \exp[-iJZ_1Z_2 - i(h_1Z_1 + h_2Z_2)/2].
$$
 (50)

In the *Z* basis, the elements of U_{12} are

$$
(U_{12})_{ab,cd} = -\frac{\sin 2b}{2} \exp(-iJ[ab+cd] - i\tilde{J}[ac+bd])
$$

× exp (-ih₁[a + c]/2 - ih₂[b + d]/2), (51)

where *a*, *b*, *c*, *d* \in {1, -1} and

$$
\tilde{J} = -\frac{\pi}{4} - \frac{i}{2} \ln \tan b. \tag{52}
$$

The matrix $(U_{12})_{ab,cd}$ is unitary, but the matrix \tilde{U}_{12} with elements $(\tilde{U})_{ab,cd} = (U_{12})_{ac,bd}$ is not *except* at the self-dual points, Eq. [\(45\)](#page-6-0). The unitarity of U_{12} has the graphical representation

$$
\begin{pmatrix}\n a \\
 b\n\end{pmatrix}\n\begin{pmatrix}\n b \\
 c\n\end{pmatrix}\n\begin{pmatrix}\n a \\
 b\n\end{pmatrix}\n=\n\begin{pmatrix}\n b \\
 b'\n\end{pmatrix}\n=\n\delta_{aa'}\delta_{bb'}.
$$
\n(53)\n
\n
$$
\begin{pmatrix}\n d' \\
 a'\n\end{pmatrix}
$$

The unitarity of U_{12} has the interesting consequence that the SW-NE MPS is in the left *and right* canonical form, so that

$$
\sum_{s} A_{s}^{(j)} A_{s}^{(j)\dagger} = \mathbb{1}_{2^{t-1}}.
$$
 (54)

To see this, we first give the graphical representation of the left-hand side of Eq. (54),

(55)

(56)

where we used unitarity to eliminate the top and bottom gates. Using the motif Eq. [\(53\)](#page-7-0) we can telescope the circuit until

$$
a_{3}
$$
\n
$$
a_{2}
$$
\n
$$
a_{1}
$$
\n
$$
a_{2}
$$
\n
$$
a_{1}
$$
\n
$$
b_{1, s_{2}}
$$
\n
$$
a_{1} \downarrow b_{1, s_{2}}
$$
\n
$$
a_{1} \downarrow b_{1, s_{2}}
$$
\n
$$
a_{1} \downarrow b_{1}
$$
\n
$$
a_{2}
$$
\n
$$
a_{2}
$$
\n
$$
a_{3}
$$

Finally, we use the explicit form of Eq. (51) at the self-dual point to evaluate

$$
\begin{bmatrix}\n a_1 \\
 b_1 \\
 c \\
 c \\
 c \\
 c \\
 f_1 \\
 g'_1\n\end{bmatrix} = \sum_b (U_{12})_{a_1b,cd} (U_{12}^*)_{a'_1b,cd} = \delta_{a_1a'_1},
$$
\n(57)

which corresponds to an initial state equal to a product state in the Z_i basis. This verifies the condition Eq. [\(54\)](#page-7-0).

An MPS in both left and right canonical form describes a *bistochastic* quantum channel: one that preserves the identity. As a result, the ancilla density matrix $R^j = 2^{1-t} \mathbb{1}_{2^{t-1}}$ for all *j*. Evaluating the Rényi entropies yields our result, Eq. [\(47\)](#page-7-0).

It seems likely that a similar analysis can be performed for general Clifford circuits, which also have degenerate entanglement spectra [\[8\]](#page-13-0). However, establishing this in general requires one to carve out several special cases (such as circuits that generate no entanglement at all from a number of initial states [\[30,31\]](#page-14-0)) and we will not pursue this here.

V. NUMERICAL RESULTS: RANDOM UNITARY CIRCUITS

In this section we present numerical results for the evolution of various entanglement measures, and their spatial fluctuations, for random unitary circuits. The coarse features of the evolution of the entanglement spectrum were already discussed in Ref. [\[21\]](#page-14-0). In particular, the bandwidth of the entanglement spectrum broadens linearly in time; as noted in that work, this broadening is a natural consequence of the wide separation between the entanglement and light-cone speeds. For circuits of depth \geqslant 10 this broadening implies that an appreciable fraction of the spectrum of the reduced density matrix is zero to within numerical precision. Our focus here is on the *large* eigenvalues of the reduced density matrix (which dominate Rényi entropies S_n , $n \geq 1$). Because the vast majority of the eigenvalues are near zero, this "lowentanglement-energy" tail can be described accurately by lowrank approximations as in Sec. $IIIC$, allowing us to go to circuits of depth $t = 14$ with modest computational effort.

A. Benchmarking the low-rank approximation

For depths $t \leq 10$ we can compare the entanglement spectra computed by low-rank approximation with the exact ones (Fig. [3\)](#page-9-0). Although the rank of the reduced density matrix in this case is 256, we find that working with the top 20 states allows us to match the low-energy behavior of the entanglement spectrum.

For the largest depths we have considered, exact time evolution is not feasible; however, there is an exact result for the average purity of a semi-infinite system [\[10\]](#page-14-0), viz. $\bar{\gamma} = (4/5)^{t-1}$. (Note that the exponent *t* − 1 here is one less than in [\[10\]](#page-14-0) because our partition of the system lies between the two gates in the top layer of the circuit, which therefore leaves the entanglement unaffected.) For the numerical results presented here we increase the rank of the approximation until the mean computed purity matches this exact result to within statistical error (which is about 1%). For the largest depth we have systematically considered $(t = 14)$ we need to keep \approx 120 states to match the mean purity. This is only about 1% of the spectrum; thus the low-rank approximation is much more efficient than direct propagation of the channel would be.

FIG. 3. Entanglement density of states (i.e., histogram of entanglement energies), at depth $t = 8$, computed both exactly and by low-rank approximation of the quantum channel (Sec. [III C\)](#page-5-0). For rank \geq 20 the low-energy behavior of the entanglement spectrum is well captured.

B. Shape of the entanglement spectrum

In this section we discuss the shape of the entanglement spectrum and its relation to the evolution of the Rényi entropies for RUCs. First, let us recall that the eigenstate thermalization hypothesis predicts that the reduced density matrix of a subsystem should take the form \mathcal{N} exp($-\beta H$), where *H* is the Hamiltonian of the subsystem and β is the inverse temperature. Therefore the entanglement Hamiltonian $-\ln \rho \propto \beta H + \ln \mathcal{N}$, i.e., it is a stretched and shifted version of the physical Hamiltonian. The entanglement spectrum therefore has the same shape as the physical spectrum: for a large subsystem *LA* with a local Hamiltonian, it will be essentially Gaussian in the bulk, with a bandwidth that increases as $\sqrt{L_A}$, although the extreme value statistics (corresponding to the shape of the spectrum near its ground state) are model dependent. When $\beta \sqrt{L_A}$ is large, the entanglement spectrum
dependent. When $\beta \sqrt{L_A}$ is large, the entanglement spectrum will have a large bandwidth, and therefore (because of the Jacobian) the reduced density matrix will have a density of eigenvalues $\rho(\lambda)$ distribution of the form $\rho(\lambda) \sim 1/\lambda$. Infinite temperature is a singular limit, as $\beta = 0$ so the entanglement spectrum is degenerate. In practice, a typical, randomly picked state deviates from infinite temperature by an amount ∼1/ [√]*LA*, so the entanglement "bandwidth" is *LA* independent (up to possible logarithmic dependencies that we are not concerned with here). For Floquet systems or RUCs with no conservation laws, these arguments suggest that the entanglement spectrum of a small subsystem at very late times is degenerate up to finite-size effects. Once finite-size effects are included we expect a Marchenko-Pastur distribution [\[32\]](#page-14-0).

For the case of interest to us—large subsystems at short times—the numerical evidence [\[21\]](#page-14-0) suggests that the spectrum of the reduced density matrix has the density $\rho(\lambda)$ ∼ $1/\lambda$ over many decades, at any time $1 \ll t \ll l_A$; this is qualitatively unlike the (compact) Wishart distribution that obtains at very late times [\[32\]](#page-14-0). The bulk of the spectrum of the reduced density matrix consists of eigenvalues below machine precision whenever $t \ge 8$ (Fig. 4). Here, we are concerned with the "low-energy" or "high-Schmidt-coefficient" edge,

FIG. 4. (a) Cumulative distribution function of the entanglement eigenvalues under random unitary dynamics, for various circuit depths, extracted from exact evolution of the reduced density matrix under the quantum channel corresponding to the unitary circuit. At the largest depths, an appreciable fraction of the entanglement spectrum consists of eigenvalues below machine precision. (b) A more detailed view of the entanglement spectrum near its "lowenergy" edge at various depths *t* under random unitary dynamics. Recall that the rank of the density matrix increases exponentially with the depth. All points except $t = 14$ are averaged over 3000 realizations; $t = 14$ is averaged over 300 realizations. Inset: slope of the linear growth of the entanglement density of states with energy.

which governs the behavior of the Rényi entropies $S^{(n)}$, $n \ge 1$, which we can follow out to later times $t \approx 14$ (Fig. 4). The evolution at early times is nontrivial, but appears to settle down into a well-defined limiting behavior for $t \gtrsim 7$: there is a threshold in the entanglement density of states, followed by a linear increase with entanglement energy that persists out to the energies we can reliably access. The coefficient of this linear growth is approximately $2^{t/2}$ for the larger accessible *t*. This is exponentially slower than the growth of the total number of states, so the fraction of states in the tails thins out exponentially in time.

The Rényi index dependence of $S^{(n)}$ for large *n* follows from this behavior of the limit shape, if we further assume that the entanglement spectrum is self-averaging. A simple model for the spectral density $\rho(\epsilon)$ of the entanglement energies that is consistent with the large deviation form Eq. (3) is

$$
\rho(\epsilon) = \exp\left[\alpha t \Theta(E - v_{\infty}t)\right],\tag{58}
$$

FIG. 5. Rényi-index dependence of the entropy S_{α} . The fits are to the form $S_n = S_\infty(1 - 1/n)$, which works for large *n*, with $S_\infty \simeq t/6$. Deviations from this form grow with time.

in which case

$$
v_n = \frac{\alpha - n v_\infty}{1 - n}.
$$
\n(59)

A large *n* result thus implies $v_n/v_\infty \simeq 1 - 1/n$, which is consistent with our numerical observations (Fig. 5).

Although the late-time entanglement spectrum is not numerically accessible, our results allow us to comment on a few possible qualitative scenarios of the entanglement spectrum. First, it is clear numerically that the probability density of states is exponentially small near the low-energy edge of the entanglement spectrum. This turns out to be necessary for the Rényi entropies to have distinct velocities. [If one considers, e.g., a box-shaped entanglement density of states (DOS), it is simple to show that all Rényi entropies with *n* > 0 must have the same velocity, regardless of the aspect ratio of the box. Similar results hold for Marchenko-Pastur and other possible compact shapes.] Second, one might suppose the entanglement spectrum has a Gaussian shape. Matching exact results for S_0 , the normalization of ρ , and S_2 requires the Gaussian to have a linearly growing mean and variance. Numerically, the entanglement bandwidth grows linearly in time rather than as a square root, possibly because of level repulsion. Developing a theory of how the entanglement spectrum evolves is an important question for future work: at present we do not have even a phenomenological Brownian-motion model of this growth.

C. Statistics and spatial correlations of entanglement

Out to the latest times we have considered, the sampleaveraged Rényi entropies have Gaussian distributions. (Thus, quantities such as the purity are log-normally distributed.) Whether these distributions become anisotropic at much larger system sizes is unclear; however, we have not seen any sign of incipient skewness out to the times we can simulate (Fig. 6).

We now turn to fluctuations of the entanglement across spatial cuts. The prediction of Ref. [\[8\]](#page-13-0), based on a mapping to the KPZ equation, is that the entanglement fluctuations are spatially correlated, with a correlation length $\xi(t) \sim t^{2/3}$, and

FIG. 6. Histograms of S_2 and S_{∞} for $t = 12$; lines are fits to a Gaussian.

that the width of the entanglement distribution scales as $t^{1/3}$ (i.e., the entanglement "roughens"). The method used here works with an infinite system at a fixed depth, and enables one to address these spatial correlations. We find that the spatial correlations of entanglement do get longer-ranged in time, as their power spectrum clearly narrows in *k* space (Fig. 7). The

FIG. 7. Top: power spectrum of the Fourier transform of minentropy across spatial cuts, normalized to one for all circuits. Note the narrowing of the Fourier transform with increasing *t*. Data are for a single system of length 5000. Bottom: estimate of the correlation length ξ extracted from the width of the Fourier peak.

FIG. 8. Average purity of a random unitary circuit with $q = 2$ computed by applying the quantum channel (red circles) and by the trajectory method (green bars) for $L = 1000$ steps. Good agreement is found with the result $\bar{\gamma} = (4/5)^{t-1}$ from Ref. [\[10\]](#page-14-0) (dashed blue line).

Fourier transform has a characteristic width, from which we can extract a correlation length that clearly grows sublinearly with *t*. However, the correlation length remains short out to the latest times we can access, so we do not have the dynamic range to extract meaningful exponents.

The KPZ picture also predicts that the entanglement "roughens" with time, i.e., its standard deviation grows. This is consistent with what we see, although, again, the roughening is too weak to extract meaningful exponents.

D. Trajectory approach for purity

Finally, we demonstrate the trajectory approach described in Sec. [III D.](#page-5-0) To evaluate the purity using Eq. [\(41\)](#page-6-0) we evolve a pair of trajectories $s_{1:L}$ and $t_{1:L}$ with the transition probabilities given in Eq. [\(39\)](#page-6-0). For a random unitary circuit with a purity that fluctuates with spatial position, evaluating the unaveraged purity at a point would involve averaging over many trajectories with the same set of gates. As a proof of principle we instead focus on the ensemble averaged purity, and average the fidelity $|\langle \psi_{t_{1:l}} | \psi_{s_{1:l}} \rangle|^2$ for $l = 1, \ldots L$ with a trajectory of $L = 1000$. In this case we can compare with the known exact result $\bar{\gamma} = (4/5)^{t-1}$ for random unitary circuits [\[10\]](#page-14-0). Since we are now evolving vectors in the ancilla space rather than density matrices we can simulate deeper circuits. Figure 8 shows the average purity for depths up to 18, comparing the trajectory method with the exact result, as well as with the density-matrix approach for depths up to 12. Good agreement is found in all cases.

E. Circuits with more structure

The transfer-matrix method discussed here extends directly from random unitary circuits to any other type of circuit that can be decomposed into a "brickwork" arrangement of two-site gates. We discuss two examples here: circuits with a conservation law and translation-invariant circuits.

FIG. 9. *Entanglement spectra for circuits with a conservation law*. Upper panel: behavior of the entanglement spectrum near its edge for three classes of initial states: random product states, random computational-basis bit strings ("number sharp"), and the nonrandom Néel state. Lower panel: fluctuations of the min-entropy for these three classes of states. All data are for depth $t = 12$, averaged over 3000 samples. Only the initial Néel state approaches a Gaussian distribution.

1. Number-conserving circuits

As an illustrative example we now turn to circuits with a single conservation law, which we choose to be the number of \uparrow spins in the computational basis [\[33,34\]](#page-14-0). For a random circuit, all classes of product states are equivalent; however, circuits with a conservation law yield very different entanglement DOSs depending on the initial state. Figure 9 shows the results for three classes of initial states: random product states, random bit strings in the computational basis, and a uniform Néel state. While the Néel state behaves analogously to the random unitary circuit, at least at these depths, we see that the other two types of product states give rise to very different entanglement spectra, with substantially higher DOSs at low energies. The difference can be attributed to rare states with anomalously large weight on configurations with long strings of aligned spins, which do not entangle under number-conserving dynamics [\[12,13\]](#page-14-0).

2. Translation-invariant circuits

Next, we consider translation-invariant circuits, in which all the gates at a given time step are identical. Gates could be the same at different time steps (giving a Floquet system) or random at every time step (giving a system with perfectly

FIG. 10. Power spectrum of min-entropy fluctuations in the *XXZ* chain; the initial state is a random product state in the computational basis. As in random unitary circuits, the correlation length of the entanglement fluctuations grows in time.

spatially correlated noise). We focus here on the former case. For concreteness we focus on the integrable Trotterization of the *XXZ* model that was recently introduced [\[35–37\]](#page-14-0). This model is parametrized by two parameters (η, λ) , and the dynamics consists of repeated application of the two-site gate,

$$
U(\eta,\lambda) \equiv \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & \frac{\sin\eta}{\sin(\eta+\lambda)} & \frac{\sin\lambda}{\sin(\eta+\lambda)} & 0 \\ 0 & \frac{\sin\lambda}{\sin(\eta+\lambda)} & \frac{\sin\eta}{\sin(\eta+\lambda)} & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}.
$$
 (60)

When η is imaginary and λ is real, this is a Trotterized version of the Ising phase of the *XXZ* chain, with larger η corresponding to larger easy-axis anisotropy.

One could consider the dynamics of entanglement for either random or homogeneous initial states. For random states, we expect (and find) spatial fluctuations of entanglement, which have a growing correlation length as in random

FIG. 11. Evolution of the min-entropy under the quantum channel, after *L* steps, for depth $t = 12$. The data shown are for the Trotterized *XXZ* chain at two values of the anisotropy, and (for comparison) for a translation-invariant circuit with a randomly chosen gate. The integrable *XXZ* chain has nonmonotonic entanglement growth, which is generically absent. In all cases the initial state is the Néel state ↑↓↑↓

FIG. 12. Upper panel: entanglement spectra starting from the Néel state, under Trotterized *XXZ* dynamics, vs anisotropy parameter η , at depth 12. For large η the dynamics freezes, so S_{∞} remains close to zero out to late times. Lower panel: time evolution of S_{∞} for a fixed anisotropy $\eta = 1.5i$.

unitary circuits (Fig. 10). At the accessible times, we are unable to extract any clear qualitative difference between the behavior of the correlation length in these integrable circuits and the random unitary case. For translation-invariant states (specifically the Néel state), the quantum channel converges to a definite steady state after a time interval on the order of the circuit depth (Fig. 11). Comparing the transient behavior between the *XXZ* circuit and a (presumably nonintegrable) circuit consisting of tiling a random two-site gate, we see that the transient behavior of the integrable case is different: the min-entropy overshoots its steady-state value in the integrable case, but not in the random case. Figure 12 shows the lowest 40 entanglement energies at $t = 12$ as a function of the anisotropy parameter η ; as one would expect, increasing the anisotropy slows down the growth of entanglement. This manifests itself as the top eigenvalue in the Schmidt spectrum drifting toward zero, leading to a large gap in the Schmidt spectrum. However, the min-entropy, starting from the Néel state, grows linearly out to the circuit depths we can access, with no signs of curvature. This is consistent with the intuitive picture of ballistic entanglement growth in integrable systems [\[38\]](#page-14-0), since quasiparticles move ballistically although spin dynamics is diffusive [\[37,39–41\]](#page-14-0).

An interesting quantitative difference between integrable dynamics and generic chaotic dynamics is that (for intermediate values of η) the entanglement spectrum stays "narrow"

in the integrable case: Schmidt coefficients do not rapidly spread out over many decades the way they do under random unitary dynamics. This is consistent with the quasiparticle picture, which predicts that entanglement should spread with a characteristic quasiparticle velocity that does not depend on the Rényi index. (We note that attempts to make this intuitive picture more quantitative have been technically challenging [\[42\]](#page-14-0).) How this picture extends to the case of strong anisotropy, in which the quasiparticles have a broad distribution of velocities, is an interesting open question.

VI. CONCLUSIONS

In concluding, let us summarize the technical achievements of the quantum channel approach.

(1) Unitary circuits are presented directly as matrix product states in canonical form.

(2) The spectrum of the reduced density matrix is related to that of the ancilla states exposed by the diagonal cut.

(3) The resulting quantum channel allows us to work in the infinite width limit at finite depth.

(4) Analytical results are obtained for the kicked Ising model at the self-dual point using a simple graphical calculus, simpler than the approach of Ref. [\[20\]](#page-14-0).

(5) The numerical evaluation of the channel may be improved by making a low rank approximation for the ancilla density matrix or by unraveling a quantum trajectory over the physical states.

In this work, we benchmarked the quantum channel approach against exact results for the purity in random unitary circuits, and used it to compute the shape and fluctuations of the entanglement spectrum at low "entanglement energies." With relatively little computational effort we were able to get converged results for depths $t = 14$ for the entanglement spectrum and $t = 18$ for the purity. We expect that there is room to optimize the algorithm and perform more resourceintensive computations, allowing us to access somewhat later times than we have in the present work. Our results support and extend earlier work [\[21\]](#page-14-0) showing that the entanglement spectrum has a bandwidth that grows rapidly in time, with a sharp onset. Under random unitary dynamics (with or without a conservation law), most of the Schmidt coefficients that are generated are exponentially small in circuit depth, and can therefore be truncated, allowing for efficient computation of the entanglement spectrum. We were able to compute the spatial fluctuations of entanglement for large systems,

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and finally for translation-invariant systems we obtained converged results for the entanglement spectrum by evolving the associated quantum channel to convergence.

A key distinction between the quantum channel approach and standard methods (such as time-evolving block decimation [\[43\]](#page-14-0)) for evolving a matrix-product state is that the quantum channel approach constructs the entanglement spectrum without explicitly representing the physical wave function at time *t*. Our truncation and sampling schemes are also conceptually different from that in time-evolving block decimation (TEBD) [\[24,43\]](#page-14-0). Thus the quantum channel offers benefits if one wants to compute the entanglement spectrum (since it requires storing only the object of interest); however, it is not clear how one would apply our methods to compute observables in the physical (rather than ancilla) space.

A number of avenues for future work present themselves. The thermodynamic shape of the entanglement spectrum of a semi-infinite circuit is still not well understood beyond its coarsest features. At a technical level, applying our approach to the entanglement of a finite region will involve considering the quantum channel at different "times," instead of the stationary (distribution of the) ancilla density matrix. Finally, our analysis of the kicked Ising model shows that the soluble self-dual point arises simply from the relation Eq. [\(53\)](#page-7-0), and suggests a criterion for searching for more models in the same class (see also Ref. [\[44\]](#page-14-0)). The ancilla-space density matrices for these "dual-unitary" circuits are proportional to the identity and thus have no operator entanglement [\[45\]](#page-14-0) at any time; they can be represented as matrix-product operators of bond dimension 1. For circuits close to these dual-unitary points, therefore, we expect the ancilla-space operator entanglement to grow slowly, so representing the reduced density matrix as a matrix-product operator and using TEBD methods to apply the quantum channel is a promising strategy. For generic unitary circuits, we have checked that the ancilla-space operator entanglement grows linearly in time with a prefactor of order unity, so this TEBD approach is less promising.

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