Quantum information scrambling through a high-complexity operator mapping

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Quantum information scrambling has attracted much attention amid the effort to reconcile the conflict between quantum-mechanical unitarity and the thermalization irreversibility in many-body systems. Here we propose an unconventional mechanism to generate quantum information scrambling through a high-complexity mapping from logical to physical degrees-of-freedom that hides the logical information into nonseparable many-body correlations. Corresponding to this mapping, we develop an algorithm to efficiently sample a Slater-determinant wave function and compute all physical observables in dynamics with a polynomial cost in system size. The system shows information scrambling in the quantum many-body Hilbert space characterized by the spreading of Hamming distance. At late time we find emergence of classical diffusion dynamics in this quantum many-body system. We establish that the operator mapping enabled growth in an out-of-time-order correlator exhibits exponential-scrambling behavior. The quantum information-hiding mapping approach may shed light on the understanding of fundamental connections among computational complexity, information scrambling, and quantum thermalization.

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I. INTRODUCTION

Recent developments in engineering synthetic quantum devices have achieved unprecedented controllability over a wide range of quantum degrees of freedom [1–7]. Theoretically, it has been realized that there is an intricate difference between few- and many-qubit systems—a few-qubit system undergoing unitary evolution is easily reversible by a quantum circuit, whereas the reverse for many qubits is in general very difficult or practically impossible due to quantum thermalization despite unitarity [8–13]. This conflict between unitarity and irreversibility with many qubits undermines our fundamental understanding of quantum thermalization in a closed quantum system. To reconcile the conflict, the eigenstate thermalization hypothesis has been formulated theoretically [8,9] and confirmed numerically [14]. Nevertheless, the microscopic mechanism of the thermalization remains illusive. Amid the fast-growing research interests on synthetic quantum systems for the purposes of quantum information processing and beyond, a thorough understanding of this problem is key for the field to advance forward [15,16].

The problem of many-body unitarity also emerges in modeling quantum effects of black holes [17]. The black hole information and the recent firewall paradox have received enormous research interest in the last few years [18–29]. One puzzle is that the thermal nature of black-hole evaporation

process through Hawking radiation obtained from semiclassical calculations is fundamentally inconsistent with a full quantum mechanical description.

In both contexts of quantum thermalization and blackhole information paradox, researchers are now approaching a consensus that understanding how quantum information scrambles is crucial to resolve the issues in quantum manybody unitarity [30]. In connecting quantum microscopic degrees of freedom to large-scale classical (or semiclassical) physical processes, the interplay of quantum chaos [30], operator growth [29,31], and computational complexity [32-34] is of deep fundamental interest. Recent developments following the idea of the holographic anti-de Sitter/conformal field theory correspondence [35] have explored the quantum chaotic aspect through the 0+1 dimensional Sachdev-Ye-Kitaev (SYK) model [36,37] and its variants [21,26,27,29], whose out-of-time-order correlators (OTOC) are analytically solvable at large-N limit. However, other equally important aspects such as operator growth, computational complexity, and emergence of semiclassical descriptions are not well captured by the previously studied analytically solvable models [29,36,37], deserving novel theoretical ideas for further progress [32,33].

Here we propose an unconventional approach for quantum information scrambling by applying a high complexity mapping. In our proposed mechanism, despite that the quantum state follows integrable unitary evolution, quantum information is gradually lost in a physical setting, morphing into nonseparable many-body correlations because of the high

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TABLE I. The operator-mapping approach. In our designed framework, quantum information is stored in the logical basis but scrambled in the physical basis. In the mapping from the logical to physical basis (both of which are represented by binary bit strings), we replace 0 and 1 by 0 and 01, respectively, then remove 0 at the left end to handle boundary effects. The mapping is bijective, and the Hilbert space dimensions of the two basis are identical.

	Logical basis (f, τ)	Physical basis (c, σ)
Information	Preserved	Scrambled (hidden)
Basis mapping	, 0011001⟩,	, 001010001⟩,
Hilbert space dimension	$\begin{pmatrix} L_{ au} \ N \end{pmatrix}$	$\begin{pmatrix} L_{\tau} = L + 1 - N \\ N \end{pmatrix}$
Hamiltonian	$H = \sum_{k} (f_k^{\dagger} f_{k+1} + \text{H.c.})$	$H = P \sum_{i} (c_{i}^{\dagger} c_{j+1} + \text{H.c.}) P$
Operator growth	$f_k^\dagger f_{k'}$ remains quadratic	$c_j^{\dagger}c_{j'}$ becomes scrambled

complexity of the mapping between integrals of motion and physical observables in the system. This mechanism is demonstrated by a specific carefully designed operator mapping. We show that the system exhibits information scrambling in the exponentially sized Hilbert space. This dynamical process is quantified by a Hamming distance [38]. Corresponding to the mapping, we develop an algorithm and show the system is efficiently tractable on a classical computer—the time-cost scales polynomially in the system size. At late-time quantum dynamics, we find a classical diffusion description emerges out of the microscopic quantum degrees of freedom. The mapping complexity in our proposed mechanism can be in principle generalized to more generic cases, which we expect to open up a wide window to investigate the fundamental interplay of computational complexity, quantum thermalization, and information scrambling.

II. QUANTUM INFORMATION SCRAMBLING THROUGH A HIGH-COMPLEXITY OPERATOR MAPPING

For a generic quantum system the time evolution of a physical observable O is described by the expectation value of the Heisenberg operator $O(t) = e^{iHt}Oe^{-i\bar{H}t}$, with H the system Hamiltonian. Considering a quantum state $|\Psi\rangle$ with information encoded by a set of few-body observables as $O_{\alpha}|\Psi\rangle = \lambda_{\alpha}|\Psi\rangle$ —for example these operators could be local Pauli operators or stabilizers—its quantum evolution is deterministically given by $O_{\alpha}(-t)|\Psi(t)\rangle = \lambda_{\alpha}|\Psi(t)\rangle$. Through this dynamical process the information becomes hidden into $O_{\alpha}(-t)$, which in general involve highly nonlocal manybody operators [29,31,39–41]. Tracking the mapping $\{O_{\alpha}\} \mapsto$ $\{O_{\alpha}(-t)\}\$ for a generic Hamiltonian is exponentially complex, making the extraction of quantum information after long-time evolution practically impossible, i.e., quantum information scrambles under the time evolution. The scrambling process can thus be treated as quantum information initiated in few-body degrees of freedom hidden into nonseparable many-body correlations through a high-complexity mapping. Mathematically, we can define a logical basis that stores the quantum information according to the eigenbasis of $O_{\alpha}(-t)$, and study the information scrambling in the physical degrees of freedom. The information scrambling is thus basis dependent and as a result the intrinsic physics solely relies on the mapping between logical and physical basis (see an illustration in Table I). It is worth emphasizing here that this aspect is not captured by the well-studied SYK model [36,37] or its variants despite they are analytically solvable.

It is evident that the capability of the mapping to build in nonseparable many-body correlations is required to connect the information-preserving logical basis to the scrambling physical basis, but it is unclear whether the mapping having an exponential complexity is also necessary or not. Finding a relatively lower-complexity mapping from few- to many-body operators that still causes information scrambling is not only of fundamental interest but would also assist in constructing exact models for nontrivial quantum many-body dynamics. In constructing such a mapping, we require that (i) the mapping has to be complicated enough to enable scrambling, (ii) the mapping is numerically tractable with low cost, or more specifically having polynomial computational complexity, and (iii) the locality of the quantum evolution must be respected by the mapping—the dynamics in both logical and physical basis should be local. We remark here that the Jordan-Wigner [42] or the Clifford gate unitary mapping [24] does not satisfy these requirements. In the following we devise a suitable mapping based on spin-1/2 qubits. We emphasize here that the proposed theoretical ideas are rather generalizable than restricted to the particular mapping.

Consider a one-dimensional spin chain $\sigma_j^{x,y,z}$ with the site index $j \in [1, L]$. We assume that the total spin S_z component is conserved, and shall work in the Hilbert space sector having $\sum_{j=1}^{L} \sigma_j^z/2 = S_z$. This S_z conservation holds in the quantum dynamics to be specified below. We introduce a highly nonlocal projective mapping as

$$\tau_1^z = \sigma_1^z P,
\tau_{k\neq 1}^z = \sum_{\vec{s}} \sigma_{k+\sum_{j=1}^{k-1} (s_j+1/2)}^z Q_{\vec{s}}^k P,$$
(1)

where the two projectors are $P=\prod_{j=1}^{L-1}[1-(\sigma_j^z+1)(\sigma_{j+1}^z+1)/4],\ Q_{\vec{s}}^k=\prod_{j=1}^{k-1}[s_j\tau_j^z+1/2],\ \text{with } \vec{s}_k=\{s_1,s_2,\ldots,s_{k-1}\},\ s_j=\pm 1/2,\ k\in[1,L_\tau]\ (L_\tau\equiv L/2+1-S_z).$ Correspondingly, we have $\tau_k^+\tau_k^-=\sum_{\vec{s}_k}\sigma_{k+\sum_{j=1}^{k-1}(s_j+1/2)}^kQ_{\vec{s}}^k$ $\sum_{\vec{s}_{k'}}\sigma_{k'+\sum_{j=1}^{k'-1}s_{j}+1/2}^-Q_{\vec{s}'}^{k'}P,\ \text{with }\sigma^\pm=(\sigma^x\pm i\sigma_y)/2.$ These τ operators satisfy spin-1/2 algebra. Although the mapping appears a bit cryptic, it becomes more explicit when looking at the transformation of the eigenbasis of τ_z and σ_z . Taking an eigenstate of τ_z , say $|0011001\rangle$, we first map each 0 to 0, and each 1 to 01, and then remove the leftmost 0. The

transformed state is an eigenstate of σ_z , with the example state mapped to $|001010001\rangle$ (see Table I). This mapping was previously used to study the ground-state Luttinger liquid [43,44] and also the excited-state many-body localization [45], but an operator form of the mapping was not known, and an efficient algorithm to calculate physical observables under the mapping was technically lacking—the calculation in the previous literature was consequently restricted to small system sizes [44].

We shall consider a quantum state with information encoded as $\tau_k^z |\Psi\rangle = (2m_k^0-1)|\Psi\rangle$ that undergoes dynamical evolution according to the Hamiltonian

$$H = \frac{1}{2} \sum_{k=1}^{L_{\tau}} \left[\tau_k^x \tau_{k+1}^x + \tau_k^y \tau_{k+1}^y \right]. \tag{2}$$

An open boundary condition is adopted in this work. We note here that the lattice constant is set to be the length unit and that the tunneling is the energy unit. In the σ basis, the system is still local, and we have

$$H = \frac{1}{2} \sum_{i=1}^{L} P \left[\sigma_{j}^{x} \sigma_{j+1}^{x} + \sigma_{j}^{y} \sigma_{j+1}^{y} \right] P$$
 (3)

and $\sigma_j^z|\Psi\rangle=(2n_j^0-1)|\Psi\rangle$. The eigenvalues m_k^0 and n_j^0 are related by the transformation rule specified above.

The dynamics in this system is more transparent in the fermion picture through a Jordan-Wigner transformation, where we introduce f_k and c_j corresponding to τ and σ degrees of freedom, respectively. In the fermion picture, total S_z conservation implies that the total particle number N (= $S_z + L/2$) is conserved. Corresponding to Eq. (2), f_k operators follow free fermion dynamics with a Hamiltonian $\sum_k (f_k^{\dagger} f_{k+1} + \text{H.c.})$, and c_j follow dynamics of a spinless hard-core fermion model with infinite nearest neighbor interaction, whose ground state is shown to be an interacting Luttinger liquid [43,44]. To proceed, we introduce the natural orbital bases $C_j(t)$ associated with c operators by diagonalizing the time-dependent correlation function by

$$\langle \Psi(t)|c_j^{\dagger}c_{j'}|\Psi(t)\rangle = \sum_i \lambda_i(t)U_{j,i}^c(t)U_{j',i}^{c*}(t) \tag{4}$$

and $C_i = \sum_j U_{j,c}^c c_j$. The natural orbital bases associated with f operators are defined in the same way as F_k , which is given by $F_k(t) = e^{-iHt} f_k e^{iHt}$. During the dynamical evolution, the quantum information is thus stored as $F_k^{\dagger} F_k | \Psi(t) \rangle = m_k^0 | \Psi \rangle$, whereas in the σ bases (or equivalently the c bases) the dynamical $\sigma_j^z(-t)$ operator holding the information has a nonseparable many-body character. Under this mathematical construction, the degrees of freedom τ and σ correspond to logical and physical basis, respectively. Properties of the mapping between the two are summarized in Table I.

III. METHOD AND COMPLEXITY FOR COMPUTING PHYSICAL OBSERVABLES

Despite the many-body nature of the mapping from the logical to physical qubits in Eq. (1), we find that the computational complexity to extract the physical observables is polynomial in the system size. For the dynamical quantum

state $|\Psi(t)\rangle = [F_k^\dagger(t)]^{m_k^0}|0\rangle$, the time-dependent wave function $\Psi_{\vec{m}}$ in the basis of $|\vec{m}\rangle = \prod_k [f_k^\dagger(0)]^{m_k}|0\rangle$ is a Slater determinant. It is then straightforward to calculate the few-body observables in terms of the logical qubits via Wick theorem. However, the Wick theorem does not carry over to observables composed of physical qubits. A direct calculation of the expectation value of such a physical observable O, $\langle O \rangle = \sum_{\vec{m},\vec{m}'} O_{\vec{m}\vec{m}'} \Psi_{\vec{m}}^*(t) \Psi_{\vec{m}'}(t)$, requires evaluating an exponentially large (in the system size) number of Slater determinants, and is thus intractable for large systems.

To overcome the exponential difficulty, we develop an algorithm to explicitly sample the Slater determinants efficiently with the computational complexity reduced to polynomial. This algorithm is inspired by a recent work on boson sampling which proposes a sampling method for permanents [46]. Considering a Slater-determinant state represented by a matrix U where each column represents a single-particle wave function. The algorithm contains three steps. The first step is to generate a random permutation of the integer sequence $[1, \ldots, N]$, which has complexity O(N). The generated sequence is denoted as a vector \vec{v} . The second step is to iteratively sample x_k (k is from 1 to N in the iteration) according to a conditional probability distribution

$$P(x_k|x_1,\ldots,x_{k-1};v_1,\ldots,v_k) \propto \frac{1}{k!} |\text{Det}[U_{x_1...x_k,v_1...v_k}]|^2.$$

Then we choose an all zero vector \vec{m} and then set $m_{x_k} = 1$. The vector \vec{m} satisfies the required probability distribution $|\Psi_{\vec{m}}|^2$

$$\frac{1}{N!} \sum_{\vec{v}} \left[\prod_{k} P(x_k | x_1, \dots, x_{k-1}; v_1, \dots, v_k) \right] = |\Psi_{\vec{m}}|^2, \quad (5)$$

which follows from

$$\sum_{x_k} |\text{Det } U_{x_1,\dots,x_k;v_1,\dots,v_k}|^2 = \sum_{v \in [v_1,\dots,v_k]} |\text{Det } U_{x_1,\dots,x_{k-1};[v_1,\dots,v_k] \setminus v]}|^2,$$

with $[v_1, \ldots, v_k] \setminus v$ keeping the elements in $[v_1, \ldots, v_k]$ except v. The sampling algorithm substantially improves the calculation efficiency compared to the previously used approach in studying ground state Luttinger liquids [44], and allows us to treat much larger system sizes.

The major cost to calculate the physical observables arises from sampling the determinants. In the algorithm provided above, the complexity for M_s samples scales as $O(M_sL_\tau N^{z+1})$, with N^z the computational complexity to calculate the determinant of a $N \times N$ matrix (z = 2.373 with best known algorithm). Since the sampling error scales as $\epsilon \sim 1/\sqrt{M_s}$, the computational complexity to calculate one physical observable is $O(\delta^{-2}L_\tau N^{z+1})$, with δ the required error threshold.

The overall computation complexity to calculate one physical observable is reduced to $O(L_{\tau}N^{3.373}/\delta^2)$ with δ the desired error threshold. The dynamics for the interacting physical system defined by the mapping [Eq. (1)] is thus *almost* exactly solvable, in the sense that the computational complexity does not scale exponentially.

We mention here that the matrix elements of the dynamical many-body operator $\langle \vec{m}|e^{-iHt}c_j^{\dagger}c_je^{iHt}|\vec{m}'\rangle$ holding the information in the physical basis can be calculated at a time

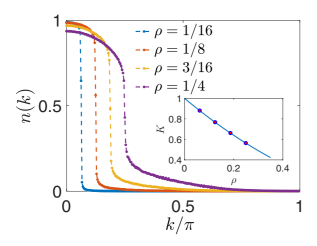


FIG. 1. Momentum distribution of the ground state hard-core spinless fermions [Eq. (3)]. The momentum distribution $n(\mathbf{k})$ of the ground state is shown here, and it exhibits Luttinger liquid behavior. As we decrease the particle density, interaction effects become weaker, and the momentum distribution is approaching Fermi-surface type. The interaction effects in this model can also be quantified by the deviation of the the ground state Luttinger parameter K from the fermion case which has K=1 (see the inset). The number of lattice sites used here is L=512.

cost which scales polynomially, much more efficient than the exponential scaling in conventional settings.

The algorithm developed here is also a significant technical *advance* in solving hard-core fermion models, which allows for calculation of much larger system sizes than previously accessible. In Fig. 1 we show the results for the ground state of Hamiltonian in Eq. (3) which can be compared to the results in Ref. [44]. A straightforward variant of our algorithm could solve all finite-range interacting hard-core fermion models efficiently.

We note here that a different method for sampling the Slater determinant is also developed in literature [47,48], which has been termed determinantal point process. This method also has a polynomial computational cost as ours. How the two different methods compare with each other in detail is left for a future study.

IV. HAMMING DISTANCE AS A MEASURE FOR INFORMATION SCRAMBLING

To quantify information scrambling from few- to many-body degrees of freedom, we generalize a previously introduced Hamming distance [38] to natural orbital occupation basis. In this basis, the Hilbert space is spanned by the time-dependent states $|\vec{n};t\rangle = [C_1^{\dagger}]^{n_1}[C_2^{\dagger}]^{n_2}\cdots|0\rangle$. The Hamming distance that characterizes the distance between the dynamical state and the initial state in the Hilbert space is

$$\mathcal{D}(t) = \sum_{\vec{n}} (\vec{n} - \vec{n_0})^2 |\langle \vec{n}; t | \Psi(t) \rangle|^2, \tag{6}$$

where \vec{n}_0 represents the initial occupation of natural orbitals. This Hamming distance is also a measure of *operator growth* [29]. The natural orbital basis is defined by sorting the eigenvalues λ_j in descending order, so that $\lambda_{j \leq N} = 1$, and $\lambda_{j > N} = 0$

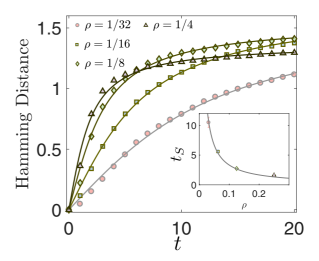


FIG. 2. Quantum information scrambling in Hilbert space measured by Hamming distance. In this figure, the shown Hamming distance is normalized by N. The system size is fixed to be 256 in this plot. We average over 10 random initial states and sampling 15 000 states in Hilbert space to minimize the sampling error. The numerical results shown by symbol points fit to a function of $\arctan(t/t_S)$. The fitted form is shown by the lines in the plot. The characteristic timescale t_S exhibits a $1/\rho$ dependence as shown in the inset.

for the initial state, and $\lambda_j \geqslant \lambda_{j+1}$ in general. For systems that do not exhibit information scrambling such as free fermions or many-body localized systems, $\mathcal{D}(t)$ remains to be an intensive quantity in dynamics, i.e., $\mathcal{D}(t)/L \to 0$, whereas $\mathcal{D}(t)/L$ becomes finite at long time when the information spreads over the Hilbert space.

A direct calculation of the Hamming distance is still exponentially difficult. We thus rewrite it in terms of observables as $\mathcal{D}(t) = 2N[1-\chi(t)]$, with $\chi(t) = \frac{1}{N} \sum_{l} n_{0,l} \lambda_{l}$. Now the Hamming distance quantifying information spreading in the physical bases can be efficiently calculated at a polynomial time cost.

The results for Hamming distance are shown in Fig. 2. The information scrambling time t_S is determined by fitting the numerical results to $\mathcal{D}(t) = \mathcal{D}_{\infty} \arctan(t/t_S)$. We find that at long time limit \mathcal{D}_{∞} is an extensive quantity, i.e., \mathcal{D}_{∞}/N is nonzero, and it approaches $2N(1-\rho)$ as we increases total N, with the fermion density $\rho = N/L$. This means the quantum information encoded in the logical τ basis scrambles to the whole many-body Hilbert space in the σ basis. The scrambling time t_S , here referring to the time for the one-dimensional system to fully scramble, is found to have a $1/\rho$ dependence, which is qualitatively as expected because the effective interactions between physical fermions increase upon increasing the particle number density.

V. CORRELATION RELAXATION DYNAMICS AND EMERGENCE OF CLASSICAL DIFFUSION AT LATE TIME

Apart from Hamming distance, the quantum memory loss in the physical basis can also be seen in the relaxation dynamics of correlations. Figure 3 shows the evolution of the natural orbital occupation numbers, which are obtained through the eigenvalues of the correlation matrix $\langle \Psi(t)|c_i^{\dagger}c_j|\Psi(t)\rangle$. In

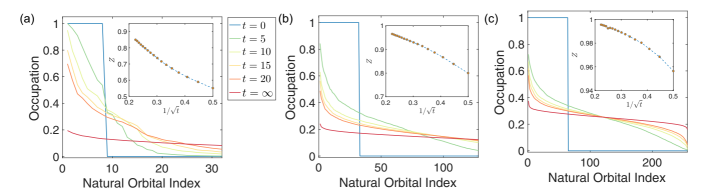


FIG. 3. Correlation relaxation dynamics of the physical degrees of freedom. (a)–(c) The occupation numbers of natural orbital basis (see main text), with different N=8,32,64, respectively. The total system size is fixed at L=256. The $t=\infty$ results correspond to $t=10^6$ in the calculation. Despite the logical degrees of freedom f_k are noninteracting, the physical ones show effective interaction effects in that natural orbital occupation exhibits relaxation behavior because otherwise they would not. The inset shows Z(t) (see main text), which fits to a power-law form of $1-Z(t)=\sqrt{t_r/t}+O(1/L)$.

the dilute limit of $\rho \to 0$, the interaction effects are vanishing, and the system is essentially formed by noninteracting fermions that do not relax. Away from that limit, we find efficient relaxation in the correlation, and the relaxation rate becomes stronger as we increase the particle number N. To quantify the relaxation dynamics, we introduce $Z(t) = \frac{1}{N} \sum_{l} \sqrt{\lambda_{l}(t) \lambda_{l}(t \to \infty)}$ that approaches 1 in the long-time limit.

We find that for large systems the long-time behavior in Z(t) is captured by $1-Z(t)=\sqrt{t_r/t}+O(1/L)$, which is attributed to classical diffusive dynamics. At a time t much longer than microscopic timescales, the system is formed by domains of fermions in a vacuum background. The typical domain size is expected to be \sqrt{Dt} , with D the diffusion constant. The late-time behavior determined by the domain-size growth would lead to the $1/\sqrt{t}$ relaxation. This behavior also implies that a classical description could emerge from quantum many-body dynamics after the quantum information scrambles in the system.

The classical description consistent with numerical results is that at late time the particle density behaves as a classical superposition of localized domains, whose domain size grows according to the classical diffusion physics, i.e., $l_d = \sqrt{Dt}$, with D the diffusion constant. With the randomly distributed domains, the central limit theorem implies that the fermion density $\langle c_j^\dagger c_j \rangle$ satisfies a normal distribution with a mean N/L and variance $\mathrm{Var} = N/L(l_d^{-1} - L^{-1})$. The average of $\sqrt{\langle c_j^\dagger c_j \rangle}$ is then given as

$$\overline{\sqrt{\langle c_j^{\dagger} c_j \rangle}} = \sqrt{N/L} \left[1 - \frac{1}{8} \frac{\text{Var}}{(N/L)^2} \right] + O(1/L), \quad (7)$$

from which we get $1 - Z(t) \sim \sqrt{t}$, after substitution of l_d by \sqrt{Dt} .

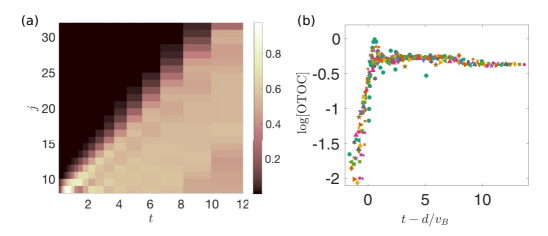


FIG. 4. Out-of-time-order correlator (OTOC) of the physical operators G_{ij} in dynamics. (a) The color plot of OTOC of $G_{i=8,j}$. In dynamics, the correlator exhibits a light-cone behavior with a finite butterfly velocity (v_B) . Once the operator separation range in OTOC is within the light cone, they quickly scramble and locally equilibrate. In this figure the temperature is fixed to be 1 here. (b) The base-10 logarithm of OTOC $G_{i=8,j}(t)$ versus $t-d/v_B$, with d the operator distance d=j-8. Different symbols correspond to different j index. The OTOC early time dynamics is consistent with an exponential growth. In the calculation we choose system size L=32 and N=8.

VI. OPERATOR-MAPPING ENABLED OTOC GROWTH

Despite the polynomial time cost in calculating all equaltime observables, our physical system still exhibits quantum chaotic behavior in the OTOC [49] as in a generic thermalizing system to be explained below. The correlator we use here takes a form of $G_{jj'}(t) = \langle [\sigma_j^z(t), \sigma_{j'}^z(0)]^{\dagger} [\sigma_j^z(t), \sigma_{j'}^z(0)] \rangle_{\beta}$, with $\sigma_j(t)$ the Heisenberg operator, and $\langle \cdots \rangle_{\beta}$ the thermal ensemble average. The sampling method we developed to calculate dynamics in few-body observables would breakdown in calculating OTOC at long time, because the sampling error would explode as $\sqrt{2^{3V_L t}/M_s}$, with M_s being the sampling number, and v_L the Lieb-Robinson bound velocity. We thus evaluate OTOC with exact methods, and afford to simulate the dynamics up to a system size L=32, with gpu techniques.

Figure 4 shows the results for OTOC where the exponential scrambling behavior of quantum information is observed. Effective interaction effects among the physical fermions are revealed as the OTOC spreading behaves as "ball-like" instead of "shell-like," which differentiates interacting and free fermions [18,19]. This further confirms the information scrambling in this system. The early time behavior of OTOC gives the Lyapunov exponent λ_L which exhibits a mild dependence of the temperature, similar to the situation in [50]. Besides, the butterfly velocity is found to be approximately equal to the sound velocity in the conformal field theory at low energy, and its implication to black hole physics is worth further investigation.

Given that the physical system exhibits quantum chaotic behavior and that the computational complexity for equal-time observables is polynomial in the system size, our quantum operator-mapping scheme is expected to motivate further researches on the minimal microscopic computational complexity for chaotic phenomena, which is of fundamental interest in understanding complex many-body systems.

VII. CONCLUSION

We have presented a mechanism for quantum information scrambling by hiding the information behind a highly nonlocal mapping, which relates the physical degrees of freedom to the logical ones holding the quantum information. A concrete operator mapping is provided to support this mechanism. We develop an algorithm to calculate physical observables, the complexity of which is polynomial in the system size. This allows us to simulate quantum dynamics of large systems. By calculating Hamming distance, correlation relaxation, and out-of-time-order correlators in dynamics, we confirm the physical system exhibits quantum information scrambling despite the fact that the mapping enabling scrambling has much lower complexity than exponential. Our discovery offers a better understanding of computation complexity, quantum thermalization, and information scrambling, and the connections among them. Furthermore, our approach can be readily generalized to construct exact solutions for novel interacting quantum dynamics. Building nontrivial but tractable mappings from quantum integrable models may inspire ideas to formulate quantum Kolmogorov-Arnold-Moser theorem [51,52], an important open question relevant to a broad range of physics spanning quantum chaos and quantum many-body localization.

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