The principle of majorization: application to random quantum circuits

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We test the principle of majorization [J. I. Latorre and M. A. Martín-Delgado, Phys. Rev. A 66, 022305 (2002)] in random circuits. Three classes of circuits were considered: (i) universal, (ii) classically simulatable, and (iii) neither universal nor classically simulatable. The studied families are: {CNOT, H, T}, {CNOT, H, NOT}, {CNOT, H, S} (Clifford), matchgates, and IQP (instantaneous quantum polynomial-time). We verified that all the families of circuits satisfy on average the principle of decreasing majorization. In most cases the asymptotic state (number of gates → ∞) behaves like a random vector. However, clear differences appear in the fluctuations of the Lorenz curves associated to asymptotic states. The fluctuations of the Lorenz curves discriminate between universal and non-universal classes of random quantum circuits, and they also detect the complexity of some non-universal but not classically efficiently simulatable quantum random circuits. We conclude that majorization can be used as a indicator of complexity of quantum dynamics, as an alternative to, e.g., entanglement spectrum and out-of-time-order correlators (OTOCs).

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I. INTRODUCTION

Majorization is a mathematical concept that allows one to decide whether a probability distribution is more disordered/spread than another. Offering an answer to such a fundamental question, the notion of majorization has found applications in many fields of social and natural sciences [1]. The first known use of the majorization notion dates back to the beginning of the twentieth century, and it is due to the economist Max Otto Lorenz who used it as a measure for wealth concentration [2].

Within physics and information theory, the concept of majorization precedes that of entropy. If a probability distribution associated to a random variable \(X\) majorizes the probability distribution of a random variable \(Y\) (see definition in Sec. [1]), then the (Shannon) entropy of \(Y\) is bigger than that of \(X\). However, the reverse statement is not necessarily true: if the entropy of \(Y\) is bigger than that of \(X\), it is not necessarily the case that \(X\) majorizes \(Y\). The majorization relation is thus a finer criteria to compare the spread of probability distributions than the entropy comparison [3]. With that in mind, Ruch and collaborators proposed a stronger version of the second law of thermodynamics, known as the principle of increasing mixing character [3-5]: the time development of a statistical ensemble of isolated systems proceeds in such a way that the probability distributions at earlier times majorize those at later times.

As expected, this intuition remains true in the quantum arena, with the majorization concept also playing an important role in quantum and nano thermodynamics (see, e.g., [6-10]). Within quantum information theory, Nielsen & Vidal’s theorem [11] asserts that majorization determines the possible transformations between bipartite entangled states by means of local operations and classical communication. One recent example related to quantum computation: in the context of boson sampling [12]. Chin and Huh explained quantitatively that the complexity of the computation relates to the majorization ordering of the input and output particle-distribution vectors [13].

While generic dynamics proceeds in the direction of growing disorder, Orús et al noticed that in many quantum algorithms the opposite is true: in the course of the computation the state of the system is step-by-step majorized, with the final result being maximally ordered [14, 15]. Later on, by carrying a systematic analysis of a variety of quantum algorithms, these authors concluded that fast and efficient algorithms (quantum Fourier transform, Grover’s algorithm, the algorithm for the hidden affine function problem, and others) obey a majorization principle (MP), i.e., satisfy step-by-step majorization. Supporting this conjecture, they gave examples of some quantum algorithms not showing any computational speed-up which violate the MP [16]. Further evidence supporting the MP came from adiabatic algorithms [17]. Recently step-by-step majorization for the Fourier Transform was observed experimentally in photonic circuits [18]. A detailed analysis of the Grover algorithm can be found in [19].

In recent years random quantum circuits have grown in importance. Besides various applications in quantum information and communication [20-22], random quantum circuits are becoming the test-bed for the so called quantum advantage. Even within the noisy intermediate scale quantum (NISQ) era, when quantum computers are composed of around hundred qubits and are still not amenable to error correction, sampling from random quantum circuits was proved a hard task for classical simulations [23, 24], and as such it

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a clear demonstration of quantum advantage. Using random quantum circuits of 53 qubits, such quantum advantage was experimentally reported in Ref. [25].

To determine properties of quantum random circuits that help to characterize their “complexity” is thus an important task. Recently, in a series of articles [26–28], it was established an apparent connection between the entanglement spectrum statistics of the output of a random circuit and its complexity – as defined by the universality of the gate set determining ensemble. More concretely, let $G$ be a set of quantum gates, and take an ensemble of $N$ qubit quantum circuits formed by uniformly sampling from $G$. Initiating the circuit with a random product state, the authors observed that the level spacing statistics for the entanglement spectrum (the Schmidt values of the balanced partition) at the end of the circuit follows the random matrix theory (RMT) prediction if $G$ forms a universal set of gates, and a Poissonian distribution in the opposite case [27]. The authors go one step further and suggest a connection between the impossibility of returning from a stationary entangled state to a separable state (via a Metropolis like algorithm) and the complexity of the random circuit: universal set of gates would lead to an effectively irreversible entanglement dynamics (when the sequence of gates leading to the stationary state is forgotten).

Another characteristic of complex quantum dynamics is that local information spreads quickly over the whole system. Widely used indicators of this process are the out-of-time-ordered correlators (OTOCs): Exponentially fast changes of OTOCs have been taken as a sign of “quantum chaos”, i.e., quantum complexity [29–31].

It is the aim of the present work to analyze the majorization principle in different ensembles of random quantum circuits. Here we show that not only the majorization principle holds (on average) for random circuits, but the fluctuations of cumulant vectors (see below) do seem to correctly identify complex computation. In addition, we show that non-universal but classically non-simulatable quantum circuits might also lead to a RMT entanglement spectrum. Our calculations demonstrate that this intermediate class of circuits can be identified by the majorization criteria.

II. MAJORIZATION & RANDOM CIRCUITS

In the following we define majorization, succinctly describe the various types of quantum random circuits used, and touch upon some related questions.

A. Majorization

Let $x$ and $y$ be real vectors of length $N$. It is said that $x$ is majorized by $y$ (or $y$ majorizes $x$), denoted by $x \prec y$, if, for all $k < N$,

$$
\sum_{i=1}^{k} x_i^{↓} \leq \sum_{i=1}^{k} y_i^{↓},
$$

and

$$
\sum_{i=1}^{N} x_i^{↓} = \sum_{i=1}^{N} y_i^{↓},
$$

where $x^{↓}$ means that the components of $x$ have been arranged in nonincreasing order. The partial sums in the equations above will be called cumulants. The $k$-th cumulant of $x$ will be denoted $F_x(k)$. Here we will always use majorization for comparing probability vectors, i.e., real vectors of non-negative components and normalized to unity: $x_i \geq 0, \forall 1 \leq i \leq N$ and $F(N) = 1$.

A very useful way of visualizing the majorization (partial) ordering $x \prec y$ is by plotting the Lorenz curves $F_x(k)$ and $F_y(k)$ vs $k/N$ [2]. Then, one has that $x \prec y$ iff the Lorenz curve for $y$ is above the curve for $x$ for all values of $k/N$. Some examples of Lorenz curves can be seen in Fig. [1]. Note that there exist vectors $x$ and $y$ for which neither $x$ majorizes $y$, nor $y$ majorizes $x$.

B. The random circuits

We consider unitary quantum circuits with $n$ qubit lines ($n = 8$ in our numerical calculations). The total system dimension, and the dimension of the probability vectors, is thus $N = 2^n$. The system evolution through the quantum circuit is given in unit time steps. Fixed a gate set $G$, at each time step a gate from $G$ is chosen and it is applied to a selected set of qubits. In a quantum random circuit the choice of gate to be applied at a given time, and the selection of qubits to which this gate is going to be applied, are both probabilistic – as defined by preset measures. For each step we evaluate the state in the computational basis, the associated probabilities, and the cumulants $F(k)$, $k = 1, \ldots , N$. This is repeated for a number $N_s$ of time steps.

We always use completely factorized initial states, i.e., $|\psi\rangle = |\psi_1\rangle \otimes \ldots \otimes |\psi_n\rangle$. Each $|\psi_i\rangle$ may be random or a fixed state. In the first case, we chose each factor state independently and uniformly distributed on the Bloch sphere (Haar measure for vectors). Diagonal-gate circuits (IQP) use always $|0\rangle^{\otimes n}$ as input. In some cases we tested both kinds of initial states for a given type of circuit.

Starting from a separable state, the successive application of gates eventually leads to highly entangled states showing statistical properties typical of random vectors. This is clearly true for the universal sets, e.g., $G_3$ (see below). Likewise, most families of circuits considered here show a similar behavior. In particular, the Lorenz curves associated to asymptotic (large number of gates) states are very close to those corresponding to random complex vectors. We denote the ensemble of $n$-qubit states, with the induced Haar measure, as Haar-$n$.

When speaking of classical simulatability of quantum circuits one can distinguish between two main notions: strong (all output probabilities are calculated) or weak (only a sample of the output probability is required). It is said that a simulation by classical means is efficient if it runs in polynomial time.
in the input size. The efficiency of a classical simulation may depend on the input type (arbitrary product state or computational basis state), and on how many output lines are measured (single-line or multiline measurement). Other ingredients like intermediate measurements, post-selection, etc., will not be considered. We have given a very compact definition of (efficient) classical simulatability. Thorough definitions may be found in the references: \[32, 36\] (Clifford circuits), \[37, 39\] (IPQ/diagonal circuits), \[40, 42\] (matchgates).

We employed seven classes of circuits: \(G1, G2, G3, MG, D2, D3, Dn\). They are described in the following three sections.

1. Circuits constructed from a few generators

We considered the three sets used in Ref. \[27\], i.e., \(G1=\{\text{CNOT, H, NOT}\}\), \(G2=\{\text{CNOT, H, S}\}\), and \(G3=\{\text{CNOT, H, T}\}\), where CNOT is the controlled-NOT gate, H stands for Hadamard, and S and T are \(\pi/4\) and \(\pi/8\) phase gates, respectively. The set \(G3\) is universal and thus approximates the full unitary group \(U(N)\) to arbitrary precision. Both sets \(G1\) and \(G2\) contain only Clifford gates, thus are nonuniversal and classically simulatable (in the setting of the Gottesman-Knill theorem \[32, 35\]). The circuits constructed from \(G2\) generate the Clifford group \[32\]. The gates in \(G1\) generate a subgroup of Clifford, then the simulatability of \(G2\) implies the simulatability of \(G1\) (in the same settings).

The probability for a gate to be selected at given time is always \(1/3\) for the four families above. We also choose with equal probability the qubits or pairs of qubits to which a selected gate is applied.

2. Matchgate circuits (MG)

Matchgates are two-qubit gates formed from two one-qubit gates A and B with the same determinant: A acts on the even parity subspace (spanned by \(|00\rangle\) and \(|11\rangle\)) and B acts on the odd parity subspace (spanned by \(|01\rangle\) and \(|10\rangle\)). A and B are randomly chosen according the Haar measure in the unitary group \(U(2)\). All pairs of qubits are equiprobable. Circuits of matchgates acting on nearest-neighbor lines only are classically simulatable; however, if the nearest-neighbor restriction is lift, the resulting circuits are universal for quantum computation \[41, 42\].

3. Diagonal-gate circuits (\(D2, D3, Dn\))

These circuits are made up from gates which are diagonal in the computational (\(Z\)) basis. The initial state is set to \(|0\rangle^n\) and Hadamard gates are placed at the beginning and ending of each line \[37\]. This class of circuits is also called IQP (instantaneous quantum polynomial-time): As diagonal gates commute, they can be applied simultaneously and thus there is not a natural time ordering of gates for a given circuit.

Diagonal circuits cannot perform universal computation, however, in general, they are not classically simulatable \[37, 43\].

We shall deal with a particular subclass of diagonal circuits: the \(r\)-qubit phase-random circuits \[44\]. In this case, each gate acts upon \(r\) qubits and has the form

\[
W_r = \text{diag}\{e^{i\phi_1}, e^{i\phi_2}, \ldots, e^{i\phi_{2^r}}\},
\]

with the \(\phi\)'s independent random uniform in \([0, 2\pi]\). The gates are applied on all combinations of \(r\) (out of \(n\)) qubits, the ordering being random. Here we use only three values for \(r\), namely \(r = 2, 3, n\). For \(r = 2\) the circuits contain \(n(n-1)/2\) gates, while for \(r = n\) the circuits consist of only one random gate. So, our diagonal circuits have a well defined number of gates. On the contrary, \(G1, G2, G3\) and \(MG\) may have an arbitrary length.

In the case \(r = 2\) we also consider the situation of gates acting on nearest-neighbor qubits. Assuming a ring topology, these circuits contain only \(n\) gates, and they are classically simulatable \[49\].

C. Abbreviations

As we are considering various types of circuits, connectivity, and initial states, for the sake of compactness, we shall employ abbreviations to designate the different possibilities. To a triplet \{circuit, connectivity, initial state\} we shall associate the word \(A-B-C\). \(A\) stands for the type of circuit: \(G1, G2, G3, MG, D2, D3, Dn\); \(B\) may equal \(m\) (nearest neighbor), \(mn\) (random neighbors), or \(all\) (all combinations of \(r\) qubits –only for diagonal circuits); and \(C\) may be \(rs\) (random initial state) or \(0\) (\(|0\rangle^n\) as input).

For instance, \(MG-m-rs\), denotes a circuit composed from matchgates acting on arbitrary neighbors, with random initial state. Analogously, \(D3-all-0\) denotes a diagonal 3-qubit random phase circuit.

Before proceeding with the numerical calculations, we note that, for our purposes, the various families of circuits to be analyzed can be split into three categories: (i) universal, (ii) classically simulatable, and (iii) neither universal nor classically simulatable. We shall try to correlate the different features in our calculations with these three classes.

III. NUMERICAL CALCULATIONS

We start by giving an example of Lorenz curves for a particular realization of a circuit \(G1\) and a random initial state. In Fig. \[1a\] we plot a set of Lorenz curves for different times. We see that, as time grows, there is an overall tendency to decreasing majorization. For example, if \(x_t\) denotes the probability vector at time \(t\), then we have \(x_0 \succ x_{25} \succ x_t\) for \(t = 50, 75, 100, \ldots\). However, there are cases of increasing majorization (curve swapping, e.g., \(x_{125} \prec x_{150}\) as well as cases of no definite majorization order (curve crossing, e.g., \(x_{75}\) and \(x_{100}\)). We have observed analogous behavior for other
realizations of $G_1$, and also for circuits belonging to other families. Not surprisingly, crossings and swappings vanish by averaging over realizations and/or initial conditions, and then majorization order appears. This can be clearly seen in Fig. 1 [panels (b)-(d)], where we exhibit Lorenz curves for selected times, but now averaged over 500 circuits/initial states. Results for families $G_1$-rn-rs, $D_3$-all-0, and $MG$-rn-rs are qualitatively similar, all showing decreasing majorization as the number of gates is increased.

Though the rate of change of the Lorenz curves depends of the type of circuit, the asymptotic curves are almost identical. From here on, we focus on the asymptotic Lorenz curves – asymptotic meaning large number of gates for circuits $G_1$, $G_2$, $G_3$, MG or the maximum number of gates for $D_2$, $D_3$, $D_n$.

Figure 2 depicts averages of the asymptotic Lorenz curves for all classes of circuits considered. Most curves coincide (within visual resolution) with that corresponding to random vectors of 8 qubits ($Haar$-8).

We have verified numerically that the Lorenz curves associated with random vectors tend to a limiting curve, $Haar$-$\infty$,
Here we observe a larger spread of the Lorenz curves, with a partial removal of the coalescence of Fig. 3. G1 and G2 (both nonuniversal) have separated off the main group, exhibiting larger fluctuations. The smallest, universal fluctuations correspond to the circuits G3-rn-rs, MG-rn-rs, DN-all-containing larger fluctuations. The smallest, universal fluctuations correspond to random vectors decrease as the number of qubits grows (cf. Fig. 3). This raises the questions: What is the associated scaling law? Do the fluctuations of the various families obey different laws? Answering these questions, though a source of potentially valuable information, lays outside of the scope of the present paper.

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A. Entanglement spectra

Following [27] we considered a 50-50 bipartition and calculated the spectra of the asymptotic reduced matrices. Let us denote the eigenvalues $\lambda_1, \lambda_2, \ldots, \lambda_{N/2}$ in decreasing order. Use the gaps $\epsilon_i = \lambda_{i+1} - \lambda_i$ to form the quotients $r_i = \epsilon_{i+1}/\epsilon_i$. Then construct the histogram $P'(r)$ by sampling within each class of circuit. If the distribution of gaps follows a Wigner-Dyson law (characteristic of Gaussian random matrix ensembles [46]), the theoretical prediction for $P'(r)$, in the limit of large matrices, is [47]

$$P_{WD}(r) = \frac{(r + r^2)^\beta}{Z (1 + r + r^2)^{1+3\beta/2}},$$

where $Z = \frac{8}{2\pi}$ for the Gaussian Orthogonal Ensemble (GOE) with $\beta = 1$, and $Z = \frac{4}{\sqrt{3}} \frac{\pi}{16}$ for the Gaussian Unitary Ensemble (GUE) with $\beta = 2$.

On the other side, if the spectrum is uncorrelated, then the corresponding statistics if Poissonian:

$$P_{\text{Poisson}}(r) = \frac{1}{(1 + r)^2}.$$  

Given that the circuits considered here do no exhibit time-reversal symmetry, their spectral statistics should follow the predictions of GUE – provided they exhibit complex dynamics. In fact, it was shown in [27] that $P'(r)$ for the universal family G3 agrees with GUE, i.e., Eq. (4) with $\beta = 2$. On the other side, the families G1 and G2 showed Poisson-like fluctuations.

Our matrices are not large enough to use the asymptotic expressions above. So, in order to account for finite dimensions effects, we resorted to numerical calculations. In the case of GUE statistics, instead of employing Eq. [4], we used the spectra of reduced density matrices obtained via partial trace of Haar random vectors of $n$ qubits. The results obtained from these spectra will be labeled Haar-n. In the other side, we generated Poissonian spectra simply by choosing $N/2$ levels $\in (0, 1)$ independently and random-uniformly.

We extend the spectral calculations of $P'(r)$ in [27] (restricted to G1, G2 and G3) to matchgates and diagonal circuits. In Fig. 4 we show histograms representing $P'(r)$ for
all the circuits. We observe that they can essentially be divided into two groups: (a) Poisson-like, i.e., decreasing probability distributions (G1-rn-rs, G2-rn-0, D2-all-0, D2-nn-0), and (b) GUE-like, i.e., peaked histograms. The latter category includes the circuits G3-rn-rs, D3-all-0, DN-all-0, MG-rn-rs, which show excellent agreement with the GUE prediction (Haar-8), and MG-rn-0 which coincides with Haar-6 (this was to be expected because MG-rn-0 is universal but has parity symmetry [48]).

The spectra corresponding to the families G1-rn-0 and G2-rn-0 consist of several multiply degenerated levels. Thus, the associated $P(r)$ are singular (nor Poisson- neither GUE-like) and were not plotted.

In the previous sections we have limited ourselves to presenting the results of our calculations, reserving the discussion thereof for the next, concluding section.

IV. DISCUSSION & FINAL REMARKS

The purpose of this paper was to characterize the complexity of quantum random circuits using majorization criteria. The authors of [27] had showed that the entanglement spectrum could be used to distinguish universal from non-universal families of circuits, using as examples the families G1, G2, G3. Here we extended the analysis of [27] in two directions. First, we considered additional families of circuits, i.e., those constructed either from matchgates or diagonal gates.

Second, we inquired about the complexity (according to majorization criteria or entanglement spectrum) of the families that, in spite of being non-universal, cannot be efficiently simulated in a classical computer. We found that some families of diagonal circuits (non-universal) must be classified as complex according to the above mentioned criteria.

After verifying that all circuit families satisfy the principle of decreasing majorization (on average), we focused on the Lorenz curves of the output states. Our results, displayed in Figs. 2, 3, 4, are summarized in Table I. The last two columns of Table I based on the strict agreement (or not) between our data and the predictions for random outputs. If we descend to the semi-quantitative level, the complexity indicators may become in conflict. For instance, MG-nn-rs [OUT(1)-STRONG simulatable] has indeed a GUE-like spectrum but intermediate-size cumulant fluctuations. On the other side, D2-all-0 [OUT(1)-WEAK simulatable] has Poisson-like spectrum, however its fluctuations are close to universal. The case of MG-nn-0 [OUT(MANY)-STRONG] is clearer: mildly GUE-like spectrum and large fluctuations. In the previous cases both indicators cooperate to characterize the respective circuit families as not complex.

Some unexpected behaviors, e.g., MG-nn-rs having a spectrum close to GUE, or D2-all-0 having Poisson-like spectrum and small cumulant fluctuations, call for further studies. In particular one should analyze how the cumulant fluctuations decrease as the system size increases. In this respect, it would be very useful to have analytical results for the fluctuations of Lorenz curves in the case of complex random vectors, which seems feasible [53].

We verified that the fluctuations of the Lorenz curves qualify as an indicator of complexity, producing essentially the same classification as the entanglement spectrum. Both measures, therefore, not only discriminate between universal and non-universal classes of random quantum circuits, but they also detect the complexity of some non-universal but not classically efficiently simulatable quantum random circuits. It should be noted, however, that the fluctuations of the Lorenz curves are more easily obtained than the entanglement spectrum. The former requires simple evaluation/measurement of the probabilities of the computational basis, while the latter requires full tomography of half the qubits and further diagonalization of the obtained reduced density matrix.

To conclude, we have introduced a criterion based on majorization (fluctuations of Lorenz curves) as an indicator of complexity of quantum computations/dynamics. This indicator is intended to serve as an alternative or complement to the well known entanglement spectrum and OTOCs.

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| Circuit name  | UNIV? | Non classically simulatable? | Classically simulatable? | Ave-H? | Fluc-H? | Spec-RMT? |
|--------------|-------|------------------------------|--------------------------|--------|---------|-----------|
| G3-mm-rs     | Yes   | Always                       | Never                    | Yes    | Yes     | Yes       |
| G2-mm-rs     | No    | OUT(MANY) WEAK [37]          | OUT(1) STRONG [33]      | Yes    | No      | No        |
| G1-mm-rs     | No    | (?))                         | OUT(MANY) STRONG [33]   | Yes    | No      | No        |
| G2-mm-0      | No    | Never                        |                          | Yes    | No      | No        |
| MG-mm-rs     | Yes   | Always                       | Never                    | Yes    | Yes     | Yes       |
| MG-mm-0      | Yes   | Always                       | Never                    | Yes    | Yes     | Yes       |
| MG-mm-rs     | No    | (?))                         | OUT(1) STRONG [41][42]   | No     | No      | No*       |
| MG-mm-0      | No    | Never                        | OUT(MANY) STRONG [40][43][50] | No | No | No |
| DN-all-0     | No    | OUT(MANY) WEAK [37]          | OUT(1) WEAK [37]        | Yes    | Yes     | Yes       |
| D3-all-0     | No    | OUT(MANY) WEAK [37]          | OUT(1) WEAK [37]        | Yes    | Yes     | Yes       |
| D2-all-0     | No    | OUT(MANY) STRONG [39]        | OUT(1) STRONG [39]      | Yes    | Yes     | Yes       |
| D2-mm-0      | No    | OUT(MANY) STRONG [39]        | OUT(1) WEAK [39]        | Yes    | Yes     | Yes       |
|              |       |                               |                          | Yes    | Yes     | Yes       |

TABLE I. Summary of the results. The first column lists the circuits. The second column says if the circuit family is universal or not. The third and fourth columns inform about the classical simulatability (or not) and the settings of the corresponding proofs. The acronyms STRONG and WEAK refer to the tasks of calculating or sampling from the output probability, respectively (see II B). OUT(1) and OUT(MANY) say whether the task is single- or multi-line. Ave-H: average cumulants of the final states coincide with those of random vectors (Haar measure); Fluc-H: same as before, but for the fluctuations of the cumulants; Spec-RMT: reduced density matrix spectra well described by random matrix theory; (?): No results, to best of our knowledge; No*: RMT-like, coincides with Haar-6, but for this non-symmetric case RMT predicts Haar-8; No**: Very close to Haar-6, however, as before, the prediction of RMT is Haar-8.

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[48] The final states of the circuits $MG_{m-0}$ possess parity symmetry. It is easy to show that the corresponding reduced density matrices also are parity-symmetric. Because of this, their entanglement spectra consist of two uncorrelated sequences, each having a definite parity. Thus, we separated the ensemble of spectra into two symmetry classes, and then calculated two sets of histograms. Both set of histograms are consistent with the GUE distribution. Some considerations apply to $MG_{m-0}$. In both cases we considered only the even-symmetry spectra (odd spectra produce statistically identical histograms).

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[51] The family $D2_{nn-0}$ might be almost strongly simulatable. Indeed, Fujii and Morimae [39] devised a simulatability proof for IQP circuits formed by gates of the type $\exp(i\theta Z_j \otimes Z_k)$, with $(j,k)$ denoting nearest-neighbor qubit lines. They name the simulability class of nearest-neighbor IQP “almost strongly simulatable”, between strongly simulatable (in the exact sense) and weakly simulatable. However, strictly speaking, one can not assert that both $D2_{nn-0}$ and nearest-neighbor IQP belong to the same simulability class because $D2_{nn-0}$ contains nearest-neighbor IQP.

[52] Matchgates acting on nearest neighbors, having random product states as inputs, are OUT(MANY)-STRONG simulatable on the line (open boundary conditions) [50]. Can this result be extended to the cycle (periodic boundary conditions, as used in the present paper)? Brod presents some geometric arguments in contrary, however, he admits that these may not be definitive: “Curiously, the circuit [...] for periodic boundary conditions corresponds to a geometry where matchgates are universal [...]”, although it might just use this geometry in a very restricted manner that does not break the simulability” [50].

[53] Tobias Micklitz and Felipe Monteiro, private communication.