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Tensor-network simulations of the surface code under realistic noise

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The surface code is a many-body quantum system, and simulating it in generic conditions is computationally hard. While the surface code is believed to have a high threshold, the numerical simulations used to establish this threshold are based on simplified noise models. We present a tensor-network algorithm for simulating error correction with the surface code under arbitrary local noise. We use this algorithm to study the threshold and the sub-threshold behaviour of the amplitude-damping and systematic rotation channels. We also compare these results to those obtained by making standard approximations to the noise models.

The working principle behind quantum error correction is to ‘fight entanglement with entanglement’, i.e., protect the data against local interaction with the environment by encoding it into delocalized degrees of freedom of a many-body system. Thus, characterizing a fault-tolerant scheme is ultimately a problem of quantum many-body physics.

While simulating quantum many-body systems is generically hard, particular systems have additional structure that can be taken advantage of. For example, free-fermion Hamiltonians have algebraic properties that makes them exactly solvable. An analogy in stabilizer quantum error correction is Pauli noise, where errors are Pauli operators drawn from some fixed distribution. Due to their algebraic structure, Pauli noise models can be simulated efficiently using the stabilizer formalism [1]. Beyond Pauli noise, noise composed of Clifford gates and projections onto Pauli eigenstates can also be simulated efficiently using the same methods [2].

While such efficiently simulable noise models can be useful to benchmark fault-tolerant schemes, they do not represent most models of practical interest. For instance, qubits that are built out of non-degenerate energy eigenstates are often subject to relaxation, a.k.a. amplitude damping. Miscalibrations often result in systematic errors corresponding to small unitary rotations [3]. Given that these processes do not have efficient descriptions within the stabilizer formalism, understanding how a given fault-tolerant scheme will respond to them is a difficult and important problem.

The simplest approach to such many-body problems is brute-force simulation, where an arbitrary state in Hilbert space is represented as an exponentially large vector of coefficients. Using such methods, small surface codes (up to distance 3) have been simulated under non-Clifford noise [4]. In another study, brute-force simulation of the seven-qubit Steane code was performed without concatenation [5]. Simulation of such low distance codes allow comparison of noise at the logical level to the noise on the physical level, however it is difficult to infer quantities of interest such as thresholds or overheads from such small simulations. Another approach, akin to the use of tight-binding approximations in solid-state physics, is to approximate these noise processes with efficiently simulable ones [2, 6]. However, the accuracy of

these approximations can be very poor as we will show below.

In this work, we import quantum many-body methods developed in the context of solid-state physics to study quantum error correction with realistic, non-Clifford noise models. Our construction hinges on the fact that the surface code is a projected-entangled-pair state (PEPS) with low bond dimension [7]. As a basic demonstration, we use our method to simulate the surface code under two non-Clifford local noise models: amplitude damping and systematic rotation. We assume that syndrome measurements are performed perfectly. We obtain thresholds for these noise models and also study error correction in the region of practical interest, where the noise strength is low relative to the threshold. These results are compared with those obtained using standard Pauli approximations, and significant discrepancies are observed. We have performed exact simulations on codes of up to 153 data qubits, while in contrast, previous studies using brute-force simulations were limited to 13 data qubits and 12 syndrome qubits [4].

Tensor networks — For our purposes, a tensor A_{i_1, i_2, \dots, i_n} is an n -index array of complex numbers, where i_k runs from 0 to $D_k - 1$, where D_k is called the bond dimension of i_k . We represent a tensor graphically as a box, and each index of the tensor as an edge emanating from that box. When two tensors are linked by an edge, the connected indices are summed over, for instance

$$\sum_{\alpha} A_{\alpha, i_1, i_2, \dots} B_{\alpha, j_1, j_2, \dots} = \begin{array}{c} \begin{array}{|c|} \hline i_1 \\ \hline \end{array} \begin{array}{|c|} \hline i_2 \\ \hline \end{array} \begin{array}{|c|} \hline \vdots \\ \hline \end{array} \begin{array}{|c|} \hline \alpha \\ \hline \end{array} \begin{array}{|c|} \hline j_1 \\ \hline \end{array} \begin{array}{|c|} \hline j_2 \\ \hline \end{array} \begin{array}{|c|} \hline \vdots \\ \hline \end{array} \\ \hline \end{array} \quad .$$

It will often be convenient to group multiple indices together, using the notation $A_{i_1, i_2, \dots, i_n} = \mathbf{A}_i$ where a bold subscript is taken to mean that the index is composed of multiple indices.

We will define an N -particle projected entangled pair state (PEPS) $|\psi\rangle$ to be a quantum state whose tensor of coefficients $\psi_{i_1, i_2, \dots, i_N} = \langle i_1, \dots, i_N | \psi \rangle$ is the contraction of a network of N tensors $\mathcal{T} = \{A_{i_1, \alpha_1}^{(1)}, A_{i_2, \alpha_2}^{(2)}, \dots, A_{i_N, \alpha_N}^{(N)}\}$, each of which has one physical index labelled i and some number of virtual indices labelled α . We assume that the physical particles have

dimension d , that each virtual index has bond dimension D , and the number of virtual indices per tensor in \mathcal{T} is less than a constant n . This implies that the tensor network description of $|\psi\rangle$ is memory efficient: while the tensor $\psi_{i_1, i_2 \dots i_N}$ has d^N entries, each tensor in \mathcal{T} has at most dD^n entries and thus the whole set \mathcal{T} can be specified with only NdD^n complex numbers.

The surface code tensor network — We consider the optimized layout of the surface code introduced in [8] where qubits are placed on the vertices of a $W \times L$ rectangular lattice and x -check operators $A_f = \prod_{i \in f} X_i$ and z -check operators $B_f = \prod_{i \in f} Z_i$ are defined on alternating faces of the lattice in a checkerboard pattern. This layout is illustrated in Supplemental Material III. The logical qubit state $|0\rangle_L$ is defined to be the simultaneous $+1$ eigenspace of all check operators and of the logical operator \bar{Z} , where \bar{Z} is a string of Z 's along the left boundary of the code. Likewise, the state $|1\rangle_L$ is fixed by all check operators but is a -1 eigenstate of \bar{Z} , it can be obtained as $\bar{X}|0\rangle_L$, where \bar{X} is a product of X operators along the bottom boundary of the code. Given that the product state $|0\rangle^{\otimes N}$ has non-zero overlap with $|0\rangle_L$, and that $|0\rangle^{\otimes N}$ is a $+1$ eigenstate of every B_f and \bar{Z} , we have

$$|0\rangle_L \propto \prod_f \frac{1}{2} (I + A_f) |0\rangle^{\otimes N}, \quad (1)$$

where $\frac{1}{2}(I + A_f)$ is the projector onto the $+1$ eigenspace of A_f , and the product is taken over all x -checks.

Let $W_{i, i', \alpha}(C)$, be a tensor with two physical indices i, i' and an arbitrary number of virtual indices $\alpha = \alpha_1, \alpha_2, \dots$, that depends on a 2×2 matrix C . Each index of $W(C)$ has bond dimension two, and the only non-zero entries of $W(C)$ are

$$W_{i, i', \mathbf{0}}(C) = \delta_{i, i'} \quad \text{and} \quad W_{i, i', \mathbf{1}}(C) = C_{i, i'}, \quad (2)$$

where $\mathbf{0}(\mathbf{1})$ means that all virtual indices are set to 0 (1), and the symbol $\delta_{i, i'}$ denotes the Kronecker delta. For convenience, we define $Q^\pm = W(\pm X)$ and $R^\pm = W(\pm Z)$.

Consider the projector $\frac{1}{2}(I + A_f)$ acting on particles 1, 2, 3 and 4 ordered clockwise around a face. It can easily be verified that the tensor $\langle i_1, i_2, i_3, i_4 | I + A_f | i'_1, i'_2, i'_3, i'_4 \rangle$ can be expressed as a contraction of four tensors

$$\sum_{\alpha_1, \alpha_2, \alpha_3} Q_{i_1, i'_1, \alpha_1}^+ Q_{i_2, i'_2, \alpha_1, \alpha_2}^+ Q_{i_3, i'_3, \alpha_2, \alpha_3}^+ Q_{i_4, i'_4, \alpha_3}^+. \quad (3)$$

We remark that the tensor description for the projection onto the -1 eigenspace of A_f is identical to Eq. (3) but with any one of the Q^+ tensors replaced with Q^- . Projections onto eigenspaces of B_f operators can be defined analogously by replacing all Q 's with R 's. The tensor network corresponding to the product of projectors in Eq. (1), is obtained by overlapping the tensor projector in Eq. (3) over the whole lattice.

The state $|0\rangle_L$ is then obtained by applying this projector to the state $|0\rangle^{\otimes N}$, which, in the tensor network

picture, corresponds to fixing the second index to zero, thus effectively removing it. A square-lattice tensor network $\mathcal{T} = \{A_{i_1, \alpha_1}^{(1)}, A_{i_2, \alpha_2}^{(2)}, \dots, A_{i_N, \alpha_N}^{(N)}\}$ for the state $|0\rangle_L$ is thus formed from contractions of Q^+ tensors.

While this tensor network describes the logical $|0\rangle_L$ state, we will need to be able to represent other states in order to fully characterise the transformation of the encoded information during error correction. Specifically, we want to compute the logical channel \mathcal{E}_L that is applied to the logical qubit during a round of error correction. By the Choi-Jamiołkowski isomorphism, this channel can be inferred directly from the resulting output when a Bell state of the form $|\Psi^+\rangle = |0\rangle_L |0\rangle_a + |1\rangle_L |1\rangle_a$ is input, where the first qubit is encoded in a surface code and is subject to error correction, while the second qubit is unencoded and assumed to be noise free. We can obtain a tensor network description for $|\Psi^+\rangle$ by a simple modification to the tensor network describing $|0\rangle_L$. Consider the tensor

$$\overline{\text{CNOT}} := \sum_{\alpha_1, \alpha_2, \dots, \alpha_L} Q_{i_1, i'_1, \alpha_1}^+ Q_{i_2, i'_2, \alpha_1, \alpha_2}^+ \dots Q_{i_L, i'_L, \alpha_L, \alpha_a}^+, \quad (4)$$

which can be thought of as an L qubit operator (expressed in tensor form) with a single uncontracted virtual index α_a , which we call the ancilla index. If α_a is set to 0 then $\overline{\text{CNOT}}$ is simply the identity on its physical indices, while if $\alpha_a = 1$, $\overline{\text{CNOT}}$ is an L -fold tensor product of X . Therefore if $\overline{\text{CNOT}}$ is applied to the bottom row of the tensor network for $|0\rangle_L$, the resulting tensor network state $\psi_{i_1, i_2, \dots, i_N, \alpha_a} = \langle i_1, i_2, \dots, i_N, \alpha_a | \psi \rangle$ (which includes the ancilla index as a physical particle) will be the desired Bell state $|0\rangle_L |0\rangle_a + (\bar{X}|0\rangle_L) |1\rangle_a = |\Psi^+\rangle$.

The above tensor network definitions are for pure states. However as we will be considering non-unitary noise, we will want a tensor network description for density matrices. Given a tensor network $\{A_{i_1, \alpha_1}^{(1)}, A_{i_2, \alpha_2}^{(2)}, \dots, A_{i_N, \alpha_N}^{(N)}\}$ for a pure state $|\psi\rangle$, we obtain the tensor network $\{B_{i_1, \alpha_1}^{(1)}, B_{i_2, \alpha_2}^{(2)}, \dots, B_{i_N, \alpha_N}^{(N)}\}$ for the density matrix $\langle i_1, \dots, i_N | \psi \rangle \langle \psi | i'_1, \dots, i'_N \rangle$ by defining $B_{i, \alpha}^{(k)} = A_{i, \alpha'}^{(k)} A_{i', \alpha''}^{(k)*}$, where $i = i, i'$ and $\alpha = \alpha', \alpha''$ represent combined sets of physical and virtual indices respectively. The resulting tensor network is illustrated in Fig. 1a).

Simulation of error correction — The evolution of the tensor network during the error-correction simulation is illustrated in Fig. 1. We start with the density matrix for the half encoded Bell state described above. The noisy state is obtained by applying the desired CPTP map \mathcal{E} to every qubit in the code. This corresponds to the local tensor update

$$B_{i, i', \alpha} \leftarrow \sum_{j, j'} \mathcal{E}_{ijj' i'} B_{j, j', \alpha}, \quad (5)$$

where $\mathcal{E}_{ijj' i'} := \langle i | \mathcal{E}(|j\rangle \langle j'|) | i' \rangle$, as depicted in 1b).

During error correction, every check will be measured, yielding a set of measurement outcomes $s = (m_1, m_2, \dots)$

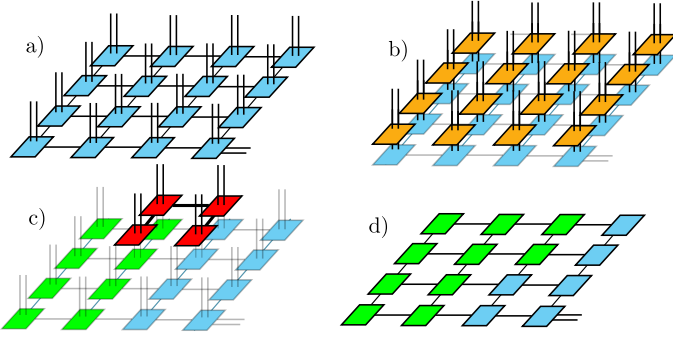


FIG. 1: Simulation of error correction with the surface code. a) The surface-code density operator as a tensor network. The two additional indices on the bottom right tensor are ancilla indices. b) Applying noise local noise to the state. c) Applying a check projector P_k (red) to the state. Green tensors have been involved in previous check measurements, while blue tensors have not. d) The resulting tensor network representing $\text{Tr}(P_{k+1}\rho_k)$ obtained by capping off pairs of physical indices.

corresponding to a syndrome s with probability $p_s = \text{Tr}(\Pi_s \rho)$, where Π_s is the projection onto the syndrome subspace corresponding to s and ρ is the noisy code state. We will assume that measurements are performed perfectly. In order to sample from the distribution p_s we imagine measuring checks sequentially. We can compute the conditional probability $p(m_k|m_1, m_2, \dots, m_{k-1})$ for any k as follows. Let ρ_k be the state of the system after k check measurements and let P_{k+1} be the projection onto the $+1$ eigenspace of the $k+1$ th check. The probabilities of obtaining an outcome of 1 and -1 are given respectively by $q = \text{Tr}(P_{k+1}\rho_k)$ and $1 - q$. To obtain a tensor network for $P_{k+1}\rho_k$, we apply the tensor corresponding to the check projector to the tensor network of ρ_k as illustrated in Fig. 1c). For example, a 4-qubit x -check can be decomposed as a contraction of four Q^+ tensors, as in Eq. (3). To apply the check to ρ_k we contract these Q^+ tensors with the measured physical indices, specifically

$$B_{i,i',\alpha} \leftarrow \sum_j Q_{i,j,\alpha'}^+ B_{j,i',\alpha''}, \quad (6)$$

where the updated virtual indices α contain the original virtual indices of B as well as those of the appended Q^+ . Then, to evaluate q , we take the trace of every tensor ($B_\alpha \leftarrow \sum_i B_{i,i,\alpha}$). The resulting tensor network has no physical indices and is illustrated in Fig. 1d). We then contract all virtual indices in the network.

This last step corresponds to contracting a square-lattice tensor network. Generically, this problem is $\#P$ -complete and therefore no efficient algorithm is believed to exist [9]. However, many efficient algorithms have been developed to obtain approximate solutions to this problem [10–15]. In this work we have used both an exact, albeit inefficient, contraction algorithm as well as an efficient, approximate contraction algorithm for finite-sized PEPS with open boundary conditions [10].

To contract the tensor network exactly, we merge all tensors of the left column into a single tensor, then contract this sequentially with the remaining tensors from left to right. The algorithm is inefficient because the amount of memory required to store the tensor associated to a column is exponential in the lattice width. We note, however, that we have used a number of optimizations to significantly improve the efficiency of the algorithm. These are described in Supplemental Material III. With exact contraction, the complexity of sampling a single syndrome is $\mathcal{O}(W4^W)$, and thus the complexity of sampling all syndromes (and of the entire algorithm) is $\mathcal{O}(LW^24^W)$. For efficient, approximate contraction, we use a well known algorithm described in [16], which represents the contraction of the network as the repeated multiplication of a matrix-product state by matrix-product operators. This algorithm depends on an accuracy parameter χ and contraction of the lattice takes time $\mathcal{O}(LW\chi^3)$.

When all check outcomes have been obtained, we have sampled a syndrome s from the distribution p_s , and the tensor network encodes a noisy state with appropriate check projectors applied. The state is then returned to the code space via a Pauli operator, which we choose to be a product of X operators connecting each flipped B_f check to the left boundary and Z connecting each flipped A_f check to the top boundary.

At this point, a classical algorithm, called the decoder, is used to select one of four possible corrections $\bar{I}, \bar{X}, \bar{Y}, \bar{Z}$ to be applied to the state. The decoder uses information about the syndrome and the noise model to select the correction that minimises the overall logical error. For efficiency this is usually only done approximately [17–26]. In this work we perform decoding by choosing the correction that minimizes the distance between the computed logical channel and the identity. The entire procedure (application of noise, check measurements, returning to the code space, correction) yields a CPTP map \mathcal{E}_L acting on the encoded qubit. The calculation of \mathcal{E}_L as well as the decoding algorithm are performed using essentially the same method as the syndrome sampling and are described in detail in Supplementary Material I.

Numerical results — Two non-Pauli noise models are considered: systematic rotation about the z -axis $\mathcal{E}_{SR}(\rho) = e^{-i\theta Z} \rho e^{i\theta Z}$ where $\theta \in [0, \pi)$ is the rotation angle, and amplitude damping $\mathcal{E}_{AD}(\rho) = \sum_i K_i \rho K_i^\dagger$, which has two Kraus operators

$$K_0 = |0\rangle\langle 0| + \sqrt{1-\gamma}|1\rangle\langle 1|, \quad K_1 = \sqrt{\gamma}|0\rangle\langle 1|, \quad (7)$$

where $\gamma \in [0, 1]$ is the damping parameter. As a test, we also performed simulations with the depolarising channel $\mathcal{E}_{DP}(\rho) = (1-\epsilon)\rho + \frac{\epsilon}{3}X\rho X + \frac{\epsilon}{3}Y\rho Y + \frac{\epsilon}{3}Z\rho Z$, which is a well studied Pauli channel.

We have also considered two different Pauli approximations of these channels. The Pauli twirl approximation (PTA) to an arbitrary channel expressed in the Pauli basis $\mathcal{E}(\rho) = \sum_{i,j} \chi_{i,j} P_i \rho P_j$ is the channel $\mathcal{E}^{\text{PTA}} =$

	Exact	PTA	HPA
DP (ϵ)	$18.5 \pm 1.5\%$	$18.5 \pm 1.5\%$	$18.5 \pm 1.5\%$
AD (γ)	$39 \pm 2\%$	$39 \pm 2\%$	$21 \pm 1\%$
SR (θ/π)	> 0.15	0.17	0.055

TABLE I: Computed thresholds for three noise models and their Pauli approximations. PTA and HPA to SR were computed using the exact value of the threshold under bit flip noise. Pauli approximations of DP are identical to the exact channel, so only the exact channel was simulated.

$\sum_i \chi_{i,i} P_i \rho P_i$, obtained by removing the off-diagonal elements of $\chi_{i,j}$ [27, 28]. It can produce noise models that are much better behaved, and thus provide poor insight into the performance of the real channel. For this reason, an honest Pauli approximation (HPA) was introduced [6], which seeks the channel \mathcal{E}^{HPA} which is as close as possible to the original channel yet produces a noisier output on every possible input. This approximation is thus expected to provide a pessimistic bound on the performance of a fault tolerant scheme under some noise process \mathcal{E} .

As a first application, we have used the exact simulation algorithm to estimate thresholds, which are presented in table I. Details on how thresholds were determined are provided in Supplemental Material II. The largest lattice sizes simulated were 9×9 for depolarising, 11×11 for systematic rotation and 9×17 for amplitude damping. For amplitude damping we found that a non-square lattice, where the \bar{X} logical operator runs along the long dimension of the code, performed significantly better than a square one.

For the depolarising channel, we can compare our obtained threshold with exact results. We find that our threshold estimate agrees with the optimal depolarising threshold of $18.9(3)\%$ [29].

The systematic rotation channel was simulated for rotation angles between $0.025\pi \leq \theta \leq 0.3\pi$. A large region showed below threshold behaviour, however no clear transition behaviour could be identified.

The thresholds of Pauli approximations did not always agree with the thresholds of the exact channels. However, the twirl approximation to amplitude damping, yielded the same threshold as the exact channel (to within the accuracy of our data). As expected, the honest Pauli approximations provided pessimistic values of the threshold for non-Pauli channels.

As another application of our algorithm, we have performed exact simulations at fixed noise rates well below threshold. The results are presented in Fig. 2 for amplitude damping with $\gamma = 9\%$, and z -rotation with $\theta = 0.005\pi$. Again, the behaviour of the twirl approximation agrees quite well with that of the exact amplitude damping channel. However the behaviour of both approximations differed considerably from the exact z -rotation. For instance, for a code with $W = 5$, the PTA underestimates the logical error rate by a factor of about

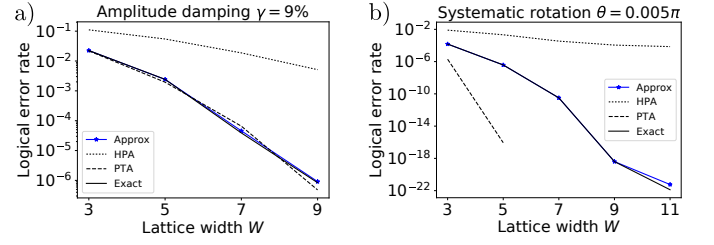


FIG. 2: Empirical logical error rate vs lattice dimension for two non-Pauli noise models. Solid black lines represent exact channels while dashed and dotted lines represent the PTA and the HPA, respectively. The left plot is $\gamma = 9\%$ amplitude damping, while the right plot is $\theta = 0.005\pi$ systematic rotation. For each set of parameters, 1.32×10^5 syndromes were sampled to acquire statistics. For the PTA of SR with $W > 5$, error rates are numerically zero. Error rates obtained for the exact channels using an approximate contraction algorithm with $\chi = 8$ are also plotted in blue. Note that the exact average logical error could significantly differ from the empirical average observed over millions of syndrome measurements if failures are dominated by outliers.

10^{10} , while the HPA over-estimates the logical error rate by a factor of about 10^4 . The observed discrepancies highlight the fact that the performance of the code under an efficiently simulable approximation to a noise model can differ significantly from its performance under the exact noise model and motivates the development of more efficient algorithms, such as the one presented here.

We have also computed error rates using the exact channels but with an approximate contraction algorithm in place of the (inefficient) exact algorithm. Setting $\chi = 8$ in the calculation of the logical channel, we observe remarkably good agreement with exact data. For instance, at high noise rates, we obtain the same threshold (within statistical error) for the amplitude damping channel (see the threshold plot in Supplemental Material II). We also observe very good agreement with exact data for systematic rotation and amplitude damping at low logical error rates, as is displayed in Fig. 2).

Conclusion — We have presented a simple tensor-network based algorithm for simulating the surface code under arbitrary local noise. The algorithm can be made exact within statistical fluctuations, allowing accurate simulation of systems with well over one hundred qubits. We have used exact simulation to estimate thresholds of non-Pauli noise models, and to calculate error rates well below threshold.

In order to demonstrate scalability of the simulation, we have also used an approximate, efficient algorithm to calculate the logical channel in place of the exact algorithm. We found very good agreement with the exact data for non-Pauli noise at both high, and low logical error rates.

While our simulation algorithm has assumed perfect check measurements, in practice, measurement errors are unavoidable. In surface-code error correction, measurement errors are detected by performing multiple rounds of check measurements and observing how the syndrome evolves in time. This procedure could potentially be sim-

ulated using PEPS time-evolution algorithms [30–33], or by contracting 3D tensor networks [34].

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