General tensor network decoding of 2D Pauli codes

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In this work we develop a general tensor network decoder for 2D codes. Specifically we propose a decoder which approximates maximally likelihood decoding for 2D stabiliser and subsystem codes subject to Pauli noise. For a code consisting of \( n \) qubits our decoder has a runtime of \( O(n \log n + n^{1+\chi}) \), where \( \chi \) is an approximation parameter. We numerically demonstrate the power of this decoder by studying four classes of codes under three noise models, namely regular surface codes, irregular surface codes, subsystem surface codes and colour codes, under bit-flip, phase-flip and depolarising noise. We show that the thresholds yielded by our decoder are state-of-the-art, and numerically consistent with optimal thresholds where available, suggesting that the tensor network decoder well approximates optimal decoding in all these cases.

Novel to our decoder is an efficient and effective approximate contraction scheme for arbitrary 2D tensor networks, which may be of independent interest. We have also released an implementation of this algorithm as a stand-alone Julia package: \texttt{SweepContractor.jl} [1].

A significant barrier to the practical adoption of quantum technologies is the inherent sensitivity of quantum systems to noise. To mitigate this, quantum error correction [2, 3] techniques were developed which allow for robust and fault-tolerant quantum information processing in spite of such noise [4–6]. These techniques utilise quantum codes, which redundantly embed quantum information in larger systems, so as to increase their robustness to noise.

A widely considered variant of error correction is that of active error correction [7, 8], in which a quantum code is augmented with a decoder which helps keep errors within the code down to a sustainable level. Specifically, the decoder is a classical algorithm which takes as input the outcomes of parity-check measurements upon the code, and outputs a candidate correction operator.

There are two properties which are important in decoder design: speed and accuracy. The balance of these two properties required depends on the use case. For decoders designed for use in practical implementations of quantum error correction for example, speed would be of particular importance, as it is known that error correction overheads are a significant contributor to the resource costs of many fault-tolerant quantum information processing algorithms [9, 10]. In this paper we will study tensor network decoding, which focuses primarily on accuracy. By attempting to approximate optimal decoder these decoders allow for the fundamental performance of codes to be probed independent of any decoder heuristics, aiding in the design of quantum codes [11]. Moreover, as we shall see, our decoder utilises tunable approximations, allowing for the speed and accuracy of our decoder to be traded-off as required.

An important family of quantum codes which are often considered as a candidate platform for scalable quantum computing are two-dimensional local codes. This is an important family which includes topological codes [12, 13] such as the toric/surface code [14] and colour code [15, 16]. The two-dimensional nature of these codes allows for them to be laid out in an array, and for the measurement of parity check operators to be performed with only local quantum operations.

Conveniently, there exists an efficient heuristic decoder for the surface code known as the minimum weight perfect matching (MWPM) decoder [17–19]. While the analogous approach does not directly yield an efficient decoder for the colour code [20], there do exist similar decoders which can be adapted to the colour code [20–22]. In all of these cases these decoders are known to be sub-optimal for bit-flip and phase-flip errors, and perform rather poorly for other error models such as depolarising noise when compared to the optimal decoder [23].

Many decoders have since been developed with speed and/or accuracy advantages over MWPM-type decoders, for both surface and for the colour codes, and for other families of 2D codes [22, 24–51]. The issues with these decoders are either that they utilise principles which are highly specialised to specific codes in question, or lack accuracy in such a way that they fall short of the optimal threshold, or—more often than not—both.

In this paper we study tensor network-based decoding. A tensor network (TN) approach to decoding was first introduced in Ref. [52], considered for the surface code in particular in Ref. [53], and utilised for studying thresholds in both the surface and colour codes in Refs. [36, 53–56]. In Ref. [57] it was shown that a similar TN can be constructed for any Pauli code subject to Pauli noise. Importantly, this TN reduces the problem of optimal decoding to the problem of TN contraction. Thus, by combining the TN with a scheme for arbitrarily precise approximate contraction, we are left with an efficient and approximately optimal general-purpose decoder.

Previous 2D TN decoders leverage a contraction scheme introduced in the condensed matter literature in Refs. [58, 59], and first utilised for decoding in Ref. [53]. This scheme explicitly relies on the TN possessing a regular structure, as occurs for codes based on lattice con-
structions. Our main technical contribution in this work is to extend this algorithm to arbitrary 2D codes, utilising algorithmic design concepts from computational geometry.

To show the efficacy of our decoder, we also give numerical evidence that our method can efficiently yield high thresholds for several important exemplary classes of 2D codes. Specifically we study four classes of codes under three noise models, namely regular surface codes, irregular surface codes, subsystem surface codes and colour codes, under bit-flip, phase-flip and depolarising noise. For each code in which either the optimal threshold is known, or there exists an analytic upper bound on the optimal threshold, we show that our decoder saturates these bounds. This suggests that not only is our decoder efficient and effective, but in fact well-approximates the optimal decoder in practice.

I. MAXIMUM LIKELIHOOD DECODING AS TENSOR NETWORK CONTRACTION

A decoder is a classical algorithm which takes as input the error syndrome and attempts to estimate the error which occurred. A natural starting point for such a decoder would simply be an algorithm which outputs the single most likely error which is consistent with this syndrome. While such a decoder is sufficient for classical codes, their quantum counterparts can possess a property called degeneracy, in which multiple logically equivalent errors can possess the same syndrome [60]. For this reason ideal decoding of quantum codes, known as maximum likelihood decoding, involves not just calculating the single most likely error, but the most likely of several classes of errors. For this reason ideal decoding is significantly more difficult for quantum codes than classical, raising the complexity from NP-complete to #P-complete [61].

Building upon earlier work [52, 53], Ref. [57] gave a general construction that reduced the problem of maximum likelihood decoding to TN contraction for arbitrary Pauli codes subject to Pauli noise. Using this construction, an approximate contraction algorithm can be utilised to approximate the maximum likelihood decoder [36, 53–57]. We direct interested readers to Ref. [57] for more details of the construction, and provide an illustrative example in Figure 1.

II. APPROXIMATE CONTRACTION OF 2D TENSOR NETWORKS

In this section we outline our approximate contraction algorithm for 2D TNs. Our algorithm builds upon techniques developed in Refs. [58, 59] for TNs on regular lattices, and utilised in the context of TN decoders in [36, 53–56]. Critically, unlike this previous work, our contraction algorithm works for arbitrary 2D TNs.

As we will not be requiring our TNs to have a regular lattice structure, we first need to be careful to define what it means for a TN to be two-dimensional in the absence of this structure. We note that our definition of a 2D TN are properties not just of the TN itself, but also of its geometric embedding.

Definition 1 (2D tensor network). A 2D tensor network is a tensor network with finite bond dimensions which can be embedded in 2D with finite density, finite range. Specifically, it is a tensor network consisting of $n$ tensors with $O(1)$ bond dimensions, where every tensor can be embedded within an $O(\sqrt{n}) \times O(\sqrt{n})$ such that the density of tensors at any point is $\Omega(1)$, and the length of any bond is $O(1)$.

Before moving on to our algorithm, we note two convenient consequences of Definition 1. The finite range and density conditions mean that a given tensor can only be connected to a bounded number of other tensors, i.e. the underlying graph has a bounded degree. Additionally, by replacing each pair of overlapping bonds with a swap tensor, our TN can always be made planar, at the cost of at most $O(n)$ additional swap tensors. As such, we will henceforth assume that our TN is bounded degree and planar.
A. Sweep line contraction algorithm

To allow our algorithm to function on not-necessarily-regular graphs we utilise the paradigm of sweep line algorithms [63] from the field of computational geometry [64].

The idea of a sweep line algorithm is to consider our data to be laid-out on the plane, and to imagine a line sweeping along this plane, with data being processed as the sweep line passes over it. In our case the idea is to maintain an approximation for the portion of the TN across which we have already swept. This approximation takes the form of a one-dimensional class of TNs, known as a matrix product state (MPS) [65, 66]. A sketch of our sweep line algorithm is given in Figure 2.

Maintenance of the MPS approximation is performed in two steps, which we refer to as CONTRACT and COMPRESS. Firstly, as the sweep line passes over each tensor, we perform CONTRACT, which contracts this tensor into the MPS and reimposes the MPS form. If our algorithm only performed the CONTRACT procedure it would in fact constitute an exact contraction procedure, but this would cause the memory size of our MPS—and therefore the runtime of our algorithm—to grow exponentially [67]. To remedy this, we augment our CONTRACT procedure by routinely running a COMPRESS procedure, which performs an efficient but lossy compression of the MPS. By combining these two procedures we obtain an efficient approximate contraction procedure.

1. CONTRACT

Firstly we find all of the MPS tensors connected to the next tensor from the network, and contract these into a single big tensor (Figure 3a). We note that, because the networks we are contracting are planar, the MPS tensors involved in this step are necessarily contiguous within the MPS. This, together with the bounded degree of the graph underlying the TN, and the bounded bond dimensions, guarantee that this step can be performed in constant time. Secondly we need to return our tensors back to the form of an MPS. To do this, we need to decompose our large multi-legged tensor into a standard single-legged MPS tensor. To do this we can utilise any matrix decomposition, but given that we are not concerned with memory costs in CONTRACT, we can simply perform this step using reshaped identity matrices (Figure 3b). Once decomposed, we can reinsert our tensors into the MPS, concluding CONTRACT (Figure 3c).

2. COMPRESS

A key feature of MPSs is that they admit an efficient procedure to compress them, known as bond truncation, allowing us to reduce the memory footprint of our MPS. Conveniently, by adjusting the bond dimensions involved we can tune this approximation at will, allowing us to trade-off the approximation error and the time/memory costs at will. See Refs. [53, 65] for details on the bond truncation procedure.

The difference from the previously considered algorithms for regular 2D TNs is when the compression step is performed. In the algorithm considered in Refs. [36, 53–56], the COMPRESS step was executed after each column of the network was contracted in, reducing the bond dimension down to $\chi$. For potentially irregular networks however there doesn’t necessarily exist a clean notion of a ‘column’ anymore. Instead we introduce a maximum bond dimension cut-off $\chi’ > \chi$: whenever the maximum bond dimension of the MPS rises above $\chi'$, we compress the entire MPS down to a bond dimension no larger than $\chi$.

B. Time and space costs

For simplicity, suppose our two bond dimensions are of the same order, $\chi' = O(\chi)$. By construction the COMPRESS step never allows our bond dimension to grow beyond $\chi'$. The length of the MPS corresponds to the number of bonds in our TN which are cut by the sweep line. Returning to Definition 1, the finite density and range constraints mean the sweep line can only cut $O(\sqrt{n})$ bonds in the worst case, and so we can conclude that our MPS has a memory cost of $O(\sqrt{n}^2)$. The total time cost of all the CONTRACT and COMPRESS procedures together are $O(n\chi^2)$ and $O(n\chi^3)$ respectively [53]. Lastly we need to recall that our procedure required us to sort our tensors by their $y$-coordinate (see Figure 2), which takes $O(n \log n)$ time to complete. Putting everything together we conclude:

\begin{align*}
\text{Time:} & \quad O(n \log n + n\chi^3) \\
\text{Space:} & \quad O(n + \sqrt{n}\chi^2)
\end{align*}

We note that if the tensors are input in an already sorted fashion, we can remove the linearithmic time cost, returning us to the $O(n\chi^3)$ run-time seen for regular 2D TN contraction [53]. To give a sense of practical runtimes, all of the numerics contained within the paper took approximately 80 CPU-years of cluster time, or approximately 35,000 CPU-hours per threshold.

We have also released a stand-alone implementation of this algorithm, in the form of a Julia package called SweepContractor.jl [1].

III. CODES CONSIDERED

To demonstrate the power of TN decoding for 2D codes, we consider applying it to four important families of 2D codes. In this section we will describe the codes considered within each family. For each code, we will determine the thresholds under three Pauli noise models: bit-flip noise, phase-flip noise, and depolarising noise. For
a more detailed description of properties of the the codes we are studying, we direct interested readers to Ref. [7].

A. Regular surface codes

The first family of codes we consider are the regular surface codes. The surface codes are a family of codes which generalise the famous toric code [14]. A surface code can be defined for any planar graph, with $Z$-type stabilisers which reside on the faces and $X$-type stabilisers which reside on the vertices.

Following on from Ref. [68], which analysed the thresholds of the regular surface codes using the sub-optimal [17, 57] minimum weight perfect matching decoder [17, 69], we will study the surface code on seven different lattices. The surface code defined on a lattice and its dual differ by a Hadamard transformation, i.e. the exchange of bit-flip and phase-flip errors. As such, we arrange the lattices considered into four dual pairs:

- Square (self dual)
- Triangular/Hexagonal
- Kagome/Rhombille
- Truncated Hexagonal/Asanoha

B. Irregular surface codes

Above we stated that the surface code can be defined on any planar graph, which needn’t necessarily be a regular lattice. The next family of codes we consider are surface codes defined on irregular and randomly constructed graphs. As we shall see in Section IV, despite their irregular structure these codes have performance comparable with their regular counterparts. One interesting property of irregular surface codes is that their resilience to errors varies qubit-to-qubit, making them a candidate for error correction in the presence of non-uniform noise [70]. Here we will study two pairs of random graph constructions:

- Rand. Triangulation/Rand. Trivalent
- Rand. Quadrangulation/Rand. Tetravalent

For such random codes, it is not clear a priori how one should define the threshold. Numerically, we compute the threshold by taking an average over such code constructions. We discuss the consequences of this definition in Appendix C.

C. Subsystem surface code

The surface codes defined above are stabiliser codes, meaning that all the check operators that define the code commute, and none are redundant. Another important construction of Pauli codes used in quantum error correction are the subsystem codes [71]. These codes are constructed with lower weight checks, requiring more measurements, with checks no longer commuting. In Ref. [72] a construction is given of a subsystem code whose encoded logical states are equivalent to the surface code, which is known as the subsystem surface code.

D. Colour code

Lastly we consider a family of topological codes which do not lie within the larger family of surface codes, the colour code. Unlike the surface code, the colour codes are self-dual, with both $X$- and $Z$-type generators lying on faces. As such, the colour code is equally robust to bit- and phase-flip errors. Unlike the surface code which can be defined for any planar graph, the colour code has
IV. RESULTS

We describe in detail the threshold procedure used in Appendix A. The main threshold results are given in Table 1.

A. Saturated upper bounds

There exists a correspondence between quantum codes and classical statistical mechanical models [17, 57] that allows for estimates of the optimal thresholds, which are an upper bound for the threshold that any specific decoder can exhibit. In Table 1, alongside our numerically determined TN decoder thresholds, we have also provided the available upper bounds from studying such statistical models from Ref. [23, 73–75]. Importantly, for every code where such an upper bound exists, our deviation from the optimal threshold is comparable to our numerical precision. This provides strong evidence that our TN decoder is not only an efficient decoder, but also well-approximates optimal decoding. By increasing the code sizes and bond dimensions used, we believe that the accuracy of this method can likely be further increased, albeit at the cost of additional time and memory.

B. Hashing bound

The thresholds of surface codes on lattices other than the standard square lattice were first considered in Ref. [68], in which they were analysed under the sub-optimal minimum weight perfect matching decoder [17]. It was observed that, based on the connectivity of the underlying graph, there is a trade-off between the bit-flip and phase-flip thresholds. Furthermore, it was conjectured that this trade-off was captured by the zero-rate hashing bound [76, 77]. Specifically, it was conjectured for such codes that

\[ h(\tau_X) + h(\tau_Z) = 1, \]

where \( \tau_X, \tau_Z \) are the bit-flip and phase-flip thresholds, and \( h \) is the binary entropy. We note that this is in fact equivalent under the statistical mechanical mapping [57] to a conjecture relating the multi-critical points of random bond Ising models present in the spin glass literature [74, 78–80].

For the TN decoder, we find even closer agreement with the hashing bound, providing even more stringent evidence of this conjecture. A plot of these thresholds against the hashing bound is given in Figure 4, and the observed entropies are given in Table 2. While our thresholds do not fully saturate the hashing bound, we once again believe this gap may be closed further by considering larger code sizes and bond dimensions.

V. CONCLUSION

In this paper we have shown how to extend TN decoding to give a general-purpose decoder for arbitrary 2D Pauli codes subject to Pauli noise, and numerically demonstrated its power on several classes of 2D codes.

Our decoder requires that the code in question is 2D local, and that it encodes a constant number of logical qubits. This suggests several natural potential extensions, such as:

- 2D codes in the presence of correlated noise
- 2D codes in the presence of noisy measurements
- Higher dimensional or hyperbolic codes
- Low-density parity check codes

By using the results Ref. [57] our decoder can also be applied to correlated noise. For all other extensions however the resulting TN is no longer 2D, meaning that we need to adapt our contraction scheme to different structures of TN. We leave this for future work.
Table 1. The threshold results given by our TN decoder (see Appendix A for details). All listed uncertainties denote a standard deviation. The upper bounds come from statistical mechanical mappings considered in Refs. [23, 73–75].

### Bit-flip

| Surface code (reg.) | Observed | Upper bound |
|---------------------|----------|-------------|
| Square              | 10.917(5)% | 10.9187% [73] |
| Tri./Hex.           | 16.341(7)% | 16.4015% [73, 74] |
| Kag./Rho.           | 9.875(5)%  | ?           |
| T.H./Asa.           | 4.297(7)%  | ?           |
| Surface code (irr.) |          |             |
| Rand. Tri.          | 17.128(15)% | ?           |
| Rand. Quad.         | 12.195(12)% | ?           |

### Phase-flip

| Subsystem SC | Observed | Upper bound |
|--------------|----------|-------------|
| Square       | 6.705(13)% | 6.7407% [73] |

### Depolarising

| Surace code (irr.) | Observed | Upper bound |
|--------------------|----------|-------------|
| Rand. T ri.        | 17.436(15)% | ?           |
| Rand. Quad.        | 12.195(12)% | ?           |

### Subsystem SC

| Colour Code | Observed | Upper bound |
|-------------|----------|-------------|
| Hexagonal   | 10.910(5)% | 10.9(2)% [75] |

Table 2. Threshold entropy for surface codes under independent bit-and-phase-flip noise. The surface code is conjectured to saturate the Hashing Bound [68, 74, 78–80], corresponding to unit entropy (see Section IV B).

|                     | Entropy   |
|---------------------|-----------|
| **Regular**         |           |
| Square              | 0.9948(3) |
| Tri./Hex.           | 0.9989(3) |
| Kag./Rho.           | 0.9918(3) |
| T.H./Asa.           | 0.9915(6) |
| **Irregular**       |           |
| Rand. Tri.          | 0.9974(7) |
| Rand. Quad.         | 0.9948(7) |

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Appendix A: Threshold procedure

In this appendix we will explicitly describe the procedure we use to approximate the threshold of our decoders. Our method is based on the critical scaling approach used widely in the literature [17, 69].

For each code distance $d$ and error probability $p$, we sample $n$ error configurations. For the randomised constructions laid out in Section III B we also sample a new code for each error considered (see Appendix C for a discussion of the consequences of this). For each configuration we contract the appropriate TNs, and determine whether the decoding failed or succeeded, and counting the occurrences of each outcome.

Suppose now that our decoder succeeded in correctly decoding $s$ times, and failed $f$ times. Naively we might estimate the corresponding logical error rate as

$$p_L \approx \frac{f}{s + f} \pm \sqrt{\frac{sf}{(s+f)^3}}. \tag{A1}$$

To estimate the logical error rate, we follow the approach used by Laplace in his resolution of the Sunrise Problem [81]. If we take a uniform prior on our logical error rates and apply Bayes’ Theorem, we find that our posterior distribution over logical error rates, conditioned on whether the decoding failed or succeeded, and counting the occurrences of each outcome.

Taking the mean and standard deviation of this we may estimate our logical error rate to be

$$p_L = \frac{f + 1}{s + f + 2} \pm \sqrt{\frac{(s + 1)(f + 1)}{(s + f + 2)^2(s + f + 3)}}. \tag{A3}$$

which approaches our naïve estimate Equation (A1) for large $n$, as long as our logical error rate is well gapped away from zero. We note that if the logical error rates of our simulation were quite low, then characterising the posterior simply by its mean and standard deviation would be inappropriate due to high skewness, and other characterisations would be necessary [82].

Once we have approximations for our logical error rate for a range of different physical error rates $p$ and code distances $d$, we now use a critical scaling ansatz to estimate our threshold. Specifically we take the logical error rate $p_L$ to only depend on a non-dimensional error rate of the form $d^{1/\nu}(p - \tau)$, where $\nu$ is the critical exponent and $\tau$ the threshold, i.e.

$$p_L = f \left( d^{1/\nu}(p - \tau) \right), \tag{A4}$$

for some function $f$. We present the critical exponents observed in Table 4. As we only consider error rates around the threshold—exactly the regime in which this ansatz is believed to hold—we can take the function $f$ to be a low-order polynomial. We determine the parameters of this model by fitting to our data using a weighted least-squares fit. An example threshold fit is given in Figure 5 for the random-triangulation surface code under bit-flip noise.

A full listing of all the parameters used in our simulations are given in Table 3.

Figure 5. Threshold plot for the random triangulation surface code under bit-flip noise. a) The logical failure rate $p_L$ as a function of the physical error rate $p$. Error bars indicate one standard error, and lines correspond to the critical scaling fit through the data. Dashed line and grey region indicate the threshold and standard error thereof, $\tau = 17.128(15)\%$. b) The logical failure rate $p_L$ as a function of the non-dimensional error rate $d^{1/\nu}(p - \tau)$. The black line is the critical scaling fit. The collapse of the data here supports the validity of the critical scaling ansatz in this regime.
Table 3. Simulation parameters used in threshold calculations: $p$ is the range of physical error rates considered (always in increments of 0.1%), $d$ the code distances considered, $\chi$ the bond-dimension used, and $n$ the total number of samples taken at each point. The notation $a : b : c$ is used as a shorthand for the set $\{a, a + b, \ldots, c\}$.

Table 4. Critical scaling exponents observed during threshold fitting. The exponents for the surface codes under bit-flip/phase-flip are consistent with $\nu = 1.5$, as has previously been noted for MWPM [68].

Appendix B: Random graph constructions

In this appendix we present the two random surface code constructions described in Section III. Both constructions are based upon Delaunay Triangulations [83] of random point sets, with special care taken for the boundaries and distances of the resulting codes.

Start by considering a construction consisting of a triangulation of $n$ points randomly placed within a unit square. This code would have two issues. Firstly, this code naturally has only smooth boundaries, and not the combination of smooth and rough boundaries necessary for those codes to support a logical qubit. Secondly, because of the difference between the average degree and average co-degree, the code has a higher $Z$ distance than $X$ distance, by roughly a factor of $1.5$.

The boundary issue is resolved by fixing the locations
of the boundary qubits to be regularly placed, as they would be for the square lattice, giving the code the same clearly defined rough and smooth boundaries. The differences between the two distances is resolved by adjusting the aspect ratio of the code, randomly distributing points of a $3:2$ rectangle instead of a square. We show an example of this construction in Figure 6a.

To construct quadrangulations we can extend our triangulations by adding vertices and edges, combining both the triangulation and its dual. Specifically we construct a new graph which contains vertices for any vertex/edge/face in the original graph, connecting any vertices corresponding to an adjacent edge-vertex pair or edge-face pair in the original graph. Applying the operation to a triangulation yields a quadrangulation. Once again utilising a $3:2$ aspect ratio, as well as fixed boundary qubits, this once again can yield a surface code. We show an example of this construction in Figure 6b.

Appendix C: Self-averaging of random codes

The descriptive power of the threshold lies in its ability to describe the performance of a code family in the limit of large distance: below the threshold the logical error asymptotically vanishes, and above the threshold it approaches a maximal value.

For the randomised code families considered in Section III B, we calculate a threshold for the distribution as a whole, averaging over randomly constructed codes. While this threshold tells us about the average performance of a code sampled from our family, it is no longer clear a priori if it is still descriptive of the typical performance of a code in this family for large code sizes—or indeed if such a thing still exists.

To address this, we need to show that the random code families considered exhibit self-averaging. In Figure 7 we show the logical error rates for individual random code instances, as well as for the average over such codes, for small and large codes. While there can be considerable variation in the observed logical error rates between given code instances for smaller distances, for larger distances the codes perform far more uniformly, a hallmark of self-averaging.
Figure 7. Self-averaging of the logical error rates of surface codes on random triangulations. Each column corresponds to an error model, run on smaller ($d = 8$) and larger ($d = 64$) code examples. Each plots shows the logical error rates found after 10,000 samples of 3 specific fixed random triangulations (coloured), versus the average over 10,000 graphs (black). For clarity we have omitted the error bars for the lines corresponding to individual instances, and the lines for the average. For the smaller codes the logical error rates vary from graph to graph, but the larger codes exhibit significant self-averaging.