We study stabilizer quantum error-correcting codes (QECC) generated under hybrid dynamics of local Clifford unitaries and local Pauli measurements in one dimension. Building upon 1) a general formula relating the error-susceptibility of a subregion to its entanglement properties, and 2) a previously established mapping between entanglement entropies and domain wall free energies of an underlying spin model, we propose a statistical mechanical description of the QECC in terms of “entanglement domain walls”. Free energies of such domain walls generically feature a leading volume law term coming from its “surface energy”, and a sub-volume law correction coming from thermodynamic entropies of its transverse fluctuations. These are most easily accounted for by capillary-wave theory of liquid-gas interfaces, which we use as an illustrative tool. We show that the information-theoretic decoupling criterion corresponds to a geometric decoupling of domain walls, which further leads to the identification of the “contiguous code distance” of the QECC as the crossover length scale at which the energy and entropy of the domain wall are comparable. The contiguous code distance thus diverges with the system size as the subleading entropic term of the free energy, protecting a finite code rate against local undetectable errors. We support these correspondences with numerical evidence, where we find capillary-wave theory describes many qualitative features of the QECC; we also discuss when and why it fails to do so.

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finite code rate, to a phase where the “error rate” is so high that a finite code rate cannot be sustained.

A complementary approach, as firmly established in hybrid random Haar circuits [19, 20], translates the measurement-driven transition into a “conventional” finite-temperature ordering transition by mapping to an underlying statistical mechanical (stat. mech.) model of spins in \((2 + 0)\)-dimensions with short-range interactions, with the temporal dimension of the circuit viewed as the second spatial dimension. Within this mapping, the disorder-averaged entanglement entropy of a subregion corresponds to the free energy cost upon a change of boundary condition in that subregion. In the low-temperature ordered phase, this change of boundary condition requires the presence of sharp domain walls. This geometrical picture raises the possibility of a stat. mech. description of the entanglement structure, and, in turn, of QECCs in hybrid circuits in terms of these “entanglement domain walls”. The aim of this work is to demonstrate such a description.

We focus on error-correcting properties of stabilizer codes, as generated dynamically after running a random Clifford circuit into the steady state, where the circuit depth scales at most polynomially in the system size. We will mostly focus on the case with a maximally-mixed initial state, and with the sharp domain walls below the transition threshold, \( p < p_c \) (i.e. the mixed phase [18]; see Sec. II A), which was shown to have a finite code rate on relevant time scales.

We start in Sec. II by introducing the model using the stabilizer code formalism, and translate the circuit dynamics into their actions on the code space. We then state a theorem in Sec. II D that applies to all stabilizer codes, which equates the number of independent, undetectable (hence uncorrectable) errors supported on a subregion, with the mutual information between the subregion and the environment.

In Sec. III, we review the domain wall picture of free energies as established analytically in Refs. [19, 20], and numerically for Clifford circuits in Refs. [21, 24]. Since the Clifford stat. mech. model is not known at this stage, we choose to model the “entanglement domain walls” as the simplest type, i.e. that of the liquid-gas interface (or Ising domain walls) in the low-temperature phase, as described by what is called “capillary-wave theory” [26–28]. This simplification allows analytic calculations, and, as we shall see, quite generally captures qualitative features of entanglement domain walls. Using the theorem, we translate certain algebraic properties of the QECC to geometric properties of the domain walls. In particular, the code rate is interpreted as the surface tension, and the code distance as the length scale below which the transverse, entropic fluctuations of the domain wall dominates over the surface energy. The “correctability”

of a subregion, as quantified by the “decoupling principle”, translates into a geometric decoupling condition of domain walls.

In Sec. IV, we perform entanglement entropy calculations for a random Clifford circuit model, and demonstrate that capillary-wave theory gives a qualitatively accurate description of the results. However, quantitative deviations from capillary-wave theory are present in our numerics, which presumably reflects the specific nature of the entanglement domain walls within a stat. mech. description for such Clifford circuit dynamics.

In Sec. V, we discuss implications of our result, and mention several possible future directions.

II. MODEL AND SETTING

A. The random Clifford circuit

We consider “hybrid” circuit models [13–15] as shown in Fig. 1, acting on a set of qubits \( Q = \{1, \ldots , L\} \), arranged in a one-dimensional array. The circuit is composed of nearest-neighbor unitary gates, which we restrict to the two-qubit Clifford group,\(^2\) and sporadic single-qubit projective measurements, which we restrict to be of the Pauli operators. We focus on the quantum trajectories of the state density matrix under circuit evolution, namely,

\[
\rho_Q \rightarrow U \rho_Q U^\dagger, \quad \text{under a Clifford unitary gate,} \quad (1)
\]

\[
\rho_Q \rightarrow P \rho_Q P \frac{\text{Tr}[P \rho_Q P]}{\text{Tr}[P]}, \quad \text{under a Pauli measurement.} \quad (2)
\]

---

1 See also their precursor Ref. [25], in which the mapping was derived for random tensor networks.

2 Recall that the Clifford group contains all unitaries that maps every Pauli string operator to another under conjugation.
Here, the projection operator $P$ is given by $P = \frac{1 + g}{2}$, where $g$ is the Pauli operator being measured, and the plus-minus signs are the (possibly random) outcomes of the measurement. In the case when this outcome is indeed random, we choose either outcome randomly with the corresponding probability given by Born’s rule.

For concreteness, we choose to sample the unitaries uniformly from the two-qubit Clifford group, and perform single-qubit measurements of probability $p$ at each time step, independently on each qubit (the "random Clifford circuit" [16]).

We will focus on the maximally-mixed initial state with maximal entropy, $S(\rho_Q) = |Q| \ln 2 = L \ln 2$. This entropy can be equivalently thought of as the entanglement entropy between $Q$ and a "reference system", $R$, where $Q$ and $R$ together holds a pure state $|\Psi_{QR}\rangle$, and $\rho_Q$ is the reduced density matrix after tracing out $R$,

$$\rho_Q = \text{Tr}_R |\Psi_{QR}\rangle \langle \Psi_{QR}| = \frac{1}{2^{|Q|}} |Q\rangle.$$  \hspace{1cm} (3)

With these specifications, the circuit model is unambiguously defined. Within this model, the entropy of $\rho_Q$ is a monotonically decreasing function of time, $T$, the circuit depth. The decrease of entropy is due to measurements (Eq. (2)) that try to read out some information about the state, while the unitaries (Eq. (1)) "scramble" the information, protecting it from being read-out by local measurements. This competition leads to a "purification transition" at $p = p_c \approx 0.16$ [18, 24], where

- When $p < p_c$, the state $\rho_Q$ retains a finite density of entropy at times $T$ at most polynomial in $|Q|$ (or formally $T = O(\text{poly}(|Q|))$), therefore in the "mixed phase". \hspace{1cm} \cite{footnote:1}

- When $p > p_c$, the entropy density drops to zero on those time scales, therefore in the "pure phase". A subset of $Q$ thus has zero density of entanglement entropy, or equivalently, the entanglement entropy has an area-law scaling.

---

3 In this paper we compute the (von Neumann) entropy by taking the natural logarithm,

$$S(\rho) := -\text{Tr} \rho \ln \rho.$$  

This is the convention adopted in Refs. [19, 20, 25], for which the equality between (average) entanglement entropies and free energies can be made (see Eq. (18)). This choice of convention accounts for the extra factor of $\ln 2$ here, as well as those appearing in Eqs. (7, 13, 23) and Appendix A.

4 In the case of a pure initial state [13, 15] where this transition was first found, $p < p_c$ corresponds to a "volume law entangled phase", where one should be thinking in terms of the reduced (mixed) density matrix of a subsystem which holds a finite entropy density as $T \to \infty$.

We will primarily restrict our attention to "intermediate" time scales, with $T = O(\text{poly}(|Q|))$, since for $T$ exponentially large in $|Q|$, the circuit dynamics should fully purify the state, even when $p < p_c$.

As discussed below, the state $\rho_Q$ at any point of the circuit evolution can be thought of as a "stabilizer code", that can in principle be used for quantum error-correcting purposes. When the state is viewed as a QECC, the purification transition acquires a new interpretation: it is a transition from "good" to "bad" QECCs, where the QECC has a nonzero/zero code rate (which is equal to the entropy density, see below) at relevant time scales, respectively, in the two phases.

B. The stabilizer formalism

Here we summarize several basic notions of stabilizer QECC that are necessary for stating and using the theorem appearing towards the end of this section. We refer the reader to Refs. [9, 29] and Appendix A for details.

A "stabilizer group" $S$ is an abelian subgroup of the Pauli group on $Q$ (denoted $\mathcal{P}(Q)$) generated by $m \leq |Q|$ independent and mutually commuting Pauli string operators,

$$S = \left\{ \prod_{j=1}^m (g_j)^{b_j} | b_j \in \{0, 1\}, g_j \in \mathcal{P}(Q), [g_j, g_{j'}] = 0 \right\} = \langle g \rangle,$$  \hspace{1cm} (4)

where $G = \{g_1, \ldots, g_m\}$ is called a "generating set" of $S$. The group $S$ is abelian, where each element has order 2, and can therefore be viewed as an $m$-dimensional vector space on $\mathbb{F}_2$, the isomorphism being given explicitly above in terms of the $b$-vector.

We list a few more properties that follow from the notion of a stabilizer group [9]:

1. A stabilizer group defines a "code space", that is, the subspace $\mathcal{H}_Q(S)$ of the Hilbert space $\mathcal{H}_Q$ on which all elements of $S$ acts trivially. We have

$$\dim \mathcal{H}_Q(S) = 2^k \equiv 2^{|Q| - |\mathcal{G}|},$$  \hspace{1cm} (5)

where $k := |Q| - |\mathcal{G}| = L - m$ is known as the number of "logical qubits" encoded.

2. The stabilizer group also defines its "code state", namely the maximally-mixed state on the code space. Its density matrix is proportional to the projection operator onto the code space, and is explicitly given by [30],

$$\rho_Q(S) = \frac{1}{2^{|Q|}} \sum_{g \in \mathcal{S}} g.$$  \hspace{1cm} (6)
As an example, the maximally-mixed state (i.e. the initial state of the circuit model in Fig. 1) is such a code state, for which the stabilizer group is empty. Consequently, as we will show below, the state at any point of the random Clifford circuit evolution remains a code state, and therefore is a “stabilizer QECC” in a strict sense.

Since we will mostly be concerned with codes states as in Eq. (6), we will usually write \( \rho_Q \) as a shorthand notation for \( \rho_Q(\mathcal{S}) \), where its dependence on \( \mathcal{S} \) is implicit.

3. A code state \( \rho_Q \) as in Eq. (6) has a flat spectrum, and all its Rényi entropies are equal to \([30]\)

\[
(\ln 2)^{-1} S(\rho_Q) = |Q| - |\mathcal{G}| = k = \log_2 \dim \mathcal{H}_Q(\mathcal{S}).
\]

That is, the entropy of the code state is equal to \((\ln 2)\) times the number of logical qubits. It follows that the “code rate”, defined to be the ratio of the number of logical and physical qubits, \( \frac{k}{\dim \mathcal{H}_Q(\mathcal{S})} \), is equal to the entropy density of \( \rho_Q \) up to a factor of \( \ln 2 \).

4. We recall that a “logical operator” is an element in \( \mathcal{P}(Q) \) that commutes with all elements in \( \mathcal{S} \), that is, an element of the centralizer \( \mathcal{C}(\mathcal{S}) \).\(^{5}\) A logical operator is “trivial” if it is itself an element of \( \mathcal{S} \), and “nontrivial” otherwise. Consequently, a nontrivial logical operator acts within the code space, but nontrivially, and therefore is a so-called “undetectable and uncorrectable error” of the code.

With this trivial/nontrivial distinction, it is clear that a logical operator is defined up to gauge freedom, that is, up to arbitrary multiplications of elements in \( \mathcal{S} \) (which do not change its action on the code space). Thus logical operators are most easily thought of as “equivalence classes”, or formally, cosets of \( \mathcal{S} \) in \( \mathcal{C}(\mathcal{S}) \). We define the “logical group” \( \mathcal{L} \) as the following quotient group, \( \mathcal{L} := \mathcal{C}(\mathcal{S})/\mathcal{S} \), with \( |\mathcal{L}| = 2^k \). We note that \( \mathcal{L} \) can be generated by representative “logical Pauli X- and Z-operators”, conventionally denoted as \( \{X_{1...k}, Z_{1...k}\} \).

C. The circuit evolution in the stabilizer formalism

We briefly describe the circuit dynamics in the stabilizer formalism, with the help of notions introduced above. We show that the state at any point of the circuit evolution, as governed by Eqs. (1, 2), remains a code state as in Eq. (6).

1. Firstly, we notice that the initial maximally-mixed state is a code state with \( \mathcal{S} = \emptyset \).

2. Under a Clifford unitary gate \( U \) as in Eq. (1)

\[
\rho_Q(\mathcal{S}) \rightarrow U \rho_Q(\mathcal{S}) U^\dagger = \rho_Q(U \mathcal{S} U^\dagger),
\]

where \( \mathcal{S}' = U \mathcal{S} U^\dagger \) is obtained from \( \mathcal{S} \) by conjugating each element of \( \mathcal{S} \) by \( U \). Thus \( \mathcal{S}' \) is also an abelian subgroup of \( \mathcal{P}(Q) \). Moreover, we have \( \mathcal{S}(\rho_Q(\mathcal{S})) = \mathcal{S}(\rho_Q(\mathcal{S}')) \).

3. Under a Pauli measurement of \( g \in \mathcal{P}(Q) \) as in Eq. (2)\(^{6}\), one can easily verify that \([31]\)

(i) When \( g \) anticommutes with some elements of \( \mathcal{S} \) (hence a detectable error), it is always possible to choose \( \mathcal{G} \) such that it has \( \text{exactly one} \) element that anticommutes with \( g \). The updated stabilizer group \( \mathcal{S}' \) is generated by \( \mathcal{G}' \), where

\[
\mathcal{G}' = \{g_j | g_j \in \mathcal{G}, [g_j, g] = 0\} \cup \{g\}. \tag{9}
\]

(ii) When \( g \) commutes with all elements in \( \mathcal{S} \) and is itself within \( \mathcal{S} \) (a trivial logical operator/trivial error),

\[
\mathcal{G}' = \mathcal{G}. \tag{10}
\]

(iii) When \( g \) commutes with all elements in \( \mathcal{S} \) and is itself \( \text{not} \) within \( \mathcal{S} \) (a nontrivial logical operator/undetectable error),

\[
\mathcal{G}' = \mathcal{G} \cup \{g\}. \tag{11}
\]

Thus, if the initial state of the circuit is a “code state” as in Eq. (6), then at any point of the circuit evolution the state is a code state, which admits an efficient representation in terms of \( \mathcal{G} \), and consequently efficient simulation of the circuit dynamics, a result known as the Gottesman-Knill theorem \([31, 32]\).

Moreover, one sees from above that the entropy of the state decreases by \( \ln 2 \) (i.e. the state gets “purified”

\(^{5}\) Throughout the paper, by \( \mathcal{C}(\mathcal{S}) \) we really mean the abelianized centralizer,

\[
\mathcal{C}(\mathcal{S}) = \{g \in \mathcal{P}(Q) | [g, \mathcal{S}] = 0\} / \{\pm 1, \pm i\}.
\]

That is, we “forget about” the (uninteresting) coefficients / commutation relations of the logical operators, and focus on their operator contents. This way, \( \mathcal{C}(\mathcal{S}) \) can be viewed as vector spaces on \( \mathbb{F}_2 \), and group homomorphisms (e.g. those in Appendix A) can be viewed as linear maps between vector spaces. On the other hand, we do care about commutation relations of stabilizers (elements of \( \mathcal{S} \)). We always require \( \mathcal{S} \) to be abelian and hence identical to its abelianization. Thus, \( \mathcal{S} \) is a subgroup of the abelianized centralizer \( \mathcal{C}(\mathcal{S}) \), and the logical group \( \mathcal{L} \) is defined by their quotient.

\(^{6}\) The results here holds generally for all Pauli operators \( g \), although we are mostly interested in single site Pauli operators that are relevant in the context of the circuit model in Fig. 1 and in Eq. (2).
by one unit) if a nontrivial logical operator (or equivalently an “undetectable error”) $g$ is measured (compare Eq. (7)), but remains unchanged otherwise [23, 33]. This observation provides a first clue to a possible connection between the error-correcting properties of the state (when viewed as a QECC) and the purification dynamics.

D. Code distance and the theorem

An important metric of a QECC is its “code distance”, $d$, defined to be the minimal weight of all nontrivial logical operators. In our circuit model that has locality, it is natural to define a similar quantity, the “contiguous code distance” [18, 33], $d_{\text{cont}}$, as the minimal length of a contiguous segment of qubits that supports a nontrivial logical operator. By definition, $d \leq d_{\text{cont}}$.

We say that a logical operator $g \in C(S)$ is “localizable” on a set $A$ of qubits, if there exists $g' \in S$ such that $gg'$ acts trivially on $\overline{A}$, where $\overline{A} := Q - A$ is the complement of $A$ in $Q$. It can be verified that all logical operators localizable on a given set $A$ form a subgroup of $C(S)$. It can also be verified that this subgroup of operators localizable on $A$ contains $S$ as a subgroup, upon taking $g' = g \in S$ above. We take the quotient between these two, and denote the corresponding quotient group as $L_A$, which is a subgroup of $L$ (see Appendix A for a detailed characterization of $L_A$). We have $|L_A| = 2^{\ell_A}$, where $\ell_A$ is an integer, and has the meaning of “the maximal number of independent and inequivalent logical operators (undetectable errors)” on $A$. The quantity $\ell_A$ thus measures how susceptible the QECC is to undetectable errors on $A$.

By definition, any subset (resp. segment) $A$ of qubits with weight (resp. length) smaller than $d$ (resp. $d_{\text{cont}}$) supports no logical operators (therefore $\ell_A = 0$), or equivalently, no “undetectable errors”. An error occurring on $A$ must therefore be either “detectable” (that brings states outside the code space) or “trivial” (that leaves states within the code space unchanged). When a detectable error located on $A$ occurs, an error-correcting unitary supported on $A$ that reverses the effect of the error can be found, given its error syndrome [9, 29].

Following the standard nomenclature, we may say that the circuit defines a $[Q, k, d_{\text{cont}}]$-code over the course of its time evolution, where both $k$ and $d_{\text{cont}}$ are functions of time. A central purpose of this paper is to characterize the code dynamics, and develop an intuitive picture of its error-correcting capabilities as quantified by $k$ and $d_{\text{cont}}$. This is partly achieved by the following relation between $\ell_A$ and the entanglement structure of the state:

Theorem 1. Let $\rho_Q$ be a code state (defined in Eq. (6)) to be the maximally-mixed state on the code space, and $|\Psi_{QR}\rangle$ be an arbitrary purification of $\rho_Q$,

$$\rho_Q = \text{Tr}_R |\Psi_{QR}\rangle \langle \Psi_{QR}|.$$  

Then for any subset $A$ of $Q$, $A \subset Q$, we have

$$\ell_A = (\ln 2)^{-1} I_{A,R},$$

where the RHS is the mutual information between $A$ and $R$.

The proof of the theorem is given in Appendix A. Several comments are in order:

- The quantity $\ell_A$ was introduced and explored in Refs. [34, 35] (see also Ref. [36]), although not explicitly cast in the form of a mutual information. From the theorem it follows directly that $\ell_A + \ell_\mathcal{X} = 2k$, the “cleaning lemma” [36, 37].

- Clearly, from its definition, $\ell_A \leq \ell_{AB}$ since $A \subset AB$, which implies $I_{A,R} \leq I_{AB,R}$ or, equivalently,

$$S(\rho_A) + S(\rho_{ABR}) \leq S(\rho_{AB}) + S(\rho_{AR}),$$

the strong subadditivity inequality.

- We have not specified the pure state $|\Psi_{QR}\rangle$. However, since both sides of Eq. (13) can be defined from $\rho_Q$ alone (see the last line of Eq. (14)), any purification of $\rho_Q$ would work equally well.

- For concreteness, let us choose $|R| = k$, the minimal number of qubits required, and consider the following “encoded state” as a purification of $\rho_Q$,

$$|\Psi_{QR}\rangle = \frac{1}{\sqrt{2^k}} \sum_x |\pi_Q\rangle |x_R\rangle,$$

where $\{\pi_Q\}$ is an orthonormal basis of the code space, and $\{|x_R\rangle\}$ is an orthonormal basis of $R$.

This pure state can be obtained by starting from $k$ Bell pairs, collecting one qubit from each pair, and encoding this collection of $k$ qubits in the QECC (on $Q$) while labelling the other $k$ qubits as $R$.

The implication of the theorem when $A$ is a contiguous segment and $|A| < d_{\text{cont}}$ is important, and perhaps familiar from general considerations of QECCs. The LHS of Eq. (13) is zero, following the definition of $d_{\text{cont}}$. The RHS is therefore also zero, as it must be [38]: Since all errors on $A$ can be detected (and hence corrected), no observables on $A$ can reveal any information about the encoded state, and there should be no correlations between

---

7 Recall that the weight of a Pauli string operator is the number of qubits on which its content is not the identity operator.
leading to a vanishing mutual information between $A$ and $R$.

On a practical level, the theorem provides a concrete relationship between error correcting capabilities of the QECC and its entanglement structure. For example, one can readily “read off” the code distance of the QECC, assuming a complete knowledge of the entanglement structure.

### III. DOMAIN WALL PICTURE OF ENTANGLEMENT ENTROPIES

#### A. Mapping to a spin model

The upshot of the mapping introduced in Refs. [19, 20] for the hybrid random Haar circuit can be very roughly summarized as follows (compare Fig. 2(a)), where we omit technical details. Recall that the hybrid random Haar circuit considered in Refs. [19, 20] is structurally identical to the circuit in Fig. 1, except with each unitary gate sampled from the Haar measure on $U(4)$, and the sporadic projective measurements replaced by generalized weak measurements of the same strength on each qubit at each time step. This strength plays a role similar to the frequency of sporadic projective measurements in Fig. 1.

1. In the bulk of the circuit, there is one Potts-like spin degree-of-freedom associated with each unitary gate, taking values in $\{a, b, c, \ldots\}$. The bulk spins form a square lattice (see Fig. 2(a), and compare with Fig. 1).

2. At the $t = T$ (final time) boundary of the circuit, there is a spin associated with each physical qubit in $Q$. All spins in $Q$ are fixed to have the same value (say $a$), and therefore corresponds to a “fixed” boundary condition (b.c.) of the spin model.

3. At the $t = 0$ (initial time) boundary of the circuit, there is also a spin associated with each physical qubit in $Q$, and certain short-range entangled initial states on $Q$ corresponds to simple b.c. of the spin model. In particular, a pure product initial state corresponds to a “free” b.c., where each spin can independently take all allowed values. On the other hand, the maximally-mixed initial state corresponds to the same “fixed” b.c. ($a$) as at the $t = T$ boundary.

4. When the spatial b.c. is periodic, the circuit geometry is cylindrical, and there are no other boundaries of the circuit. When the spatial b.c. is open, the circuit geometry is rectangular, and the boundary conditions on the left and right sides of the rectangle are also “free”.

---

\[ S(\rho_{A=[x_1,x_2]}) = - \ln \frac{Z_{\text{circuit}[A]}}{Z_{\text{circuit}}} \]

**FIG. 2.** (a) Illustration of the underlying spin model in Refs. [19, 20] for the hybrid random Haar circuit. This figure is adapted from Ref. [20]. Each bulk unitary maps to a bulk spin (green square), that is “free”. Qubits at the final time $t = T$ (solid dots) correspond to “fixed” spins all pointing in either the $a$ or $b$ direction. A Boltzmann weight is associated with each downward-pointing triangle (shaded), and is a function of spins on its vertices. (b) Representation of the entanglement entropy of the segment $A$ as the difference of two free energies (see main text). In this figure, we have chosen to “zoom in” on a small part near the upper edge of the circuit; the upper edge corresponds to physical qubits $Q$ at the final time $t = T$ of the circuit evolution. The other boundaries are far away, and need not be specified. The illustrations on the RHS represent typical configurations in the low-temperature “ferromagnetic” phase, possibly after a sufficient number of coarse-graining steps. In the denominator ($Z_{\text{circuit}}$), the upper edge is colored blue, corresponding to the fixed boundary condition $a$; thus, the bulk spins tend to also order along $a$. In the numerator ($Z_{\text{circuit}[A]}$) the segment $A$ is colored yellow, and the spins are aligned to have a different value $b$. This will induce the alignment of proximate bulk spins along the same direction $b$. A domain wall is then present where the two domains meet.
5. A Boltzmann weight is defined on each downward-pointing triangle of the lattice (see Fig. 2(a)). The “circuit partition function”, \( Z_{\text{circuit}} \), is obtained by “contracting” all free spin indices of the Boltzmann weights, with the latter viewed as tensors. By construction, when talking about the circuit partition function, one needs to specify the initial state and the spatial boundary conditions.

6. To compute the entanglement entropy of a segment \( A = [x_1, x_2] \) in the final state (at \( t = T \)), one needs to compute another partition function. This partition function is defined by the same boundary condition as \( Z_{\text{circuit}} \), except with all spins in the segment \( A \) at the upper edge of the circuit “aligned” to another different value, say \( b \). We call this new partition function \( Z_{\text{circuit}}[A] \).

As shown in [19, 20, 25], the (ensemble averaged) entanglement entropy follows,

\[
S(\rho_A) = -\ln \frac{Z_{\text{circuit}}[A]}{Z_{\text{circuit}}},
\]

taking the form of a “free energy cost” due to the change of b.c. (see Fig. 2(b)).

7. The purification/entanglement transition corresponds to an ordering transition of the spin model, where the measurement strength \( p \) plays a role similar to temperature. Within the low-temperature ordered phase of the spin model, a well-defined domain wall with finite surface tension must be present to account for the b.c. change (see Fig. 2(b)). The free energy cost, mostly coming from the domain wall, will be extensive, leading to volume law entanglement entropies. We will, for brevity, call it “the entanglement domain wall”. The surface tension decreases with increasing \( p \), and eventually vanishes at the critical point.

As demonstrated in Refs. [19, 20] for random Haar circuits, this mapping requires a replica limit of the spin model (the limit where the number of available values of the Potts spins goes to 1), and enables certain predictions for critical properties of the model [20] for \( n \geq 1 \) Rényi entropies. However, the more general viewpoint of entanglement entropies (namely as free energies of domain walls) has proven useful in understanding the phase transition in other contexts: for the zeroth Rényi entropy [13] (where the entanglement entropy is equal to a “geometrical minimal cut” of the underlying lattice); and for critical properties of the random Clifford circuit [21, 24].

We will henceforth assume this general domain wall picture holds for the random Clifford circuit in the mixed phase. Since a derivation of the underlying stat. mech. model (if it exists) is unavailable at present, the precise nature of the domain walls is unknown. Nevertheless, as we shall see, the domain wall picture alone, with the additional assumption that the domain walls are of the simplest type (“Ising like”; see Fig. 2), captures much of the qualitative aspects of the entanglement entropies in the Clifford circuit. We will devote the rest of this section to capillary-wave theory of Ising domain walls and its implications, and the next section to numerical checks of capillary-wave theory for the Clifford circuit.

B. Capillary-wave theory of Ising domain walls

Capillary-wave theory [26–28] was originally proposed for describing domain walls in the low-temperature ordered phase of the Ising model. For the example in Fig. 2, a sharp domain wall must be present to be consistent with the assigned boundary conditions. For this geometry, one can further argue that it is sufficient to consider configurations with a single domain wall, which also admits the following parametrization as a “height function”，

\[
y : [x_1, x_2] \rightarrow [-T, 0],
\]

\[
x \mapsto y(x),
\]

where \( y(x_1) = y(x_2) = 0 \). With this parameterization, we are neglecting all “overhangs” and “bubbles” that might be present in the relevant configurations; these have a finite typical size in the low-temperature phase, and will eventually disappear under coarse-graining. This reasoning leads to the following approximation for the entanglement entropy,

\[
S(\rho_{A=[x_1,x_2]}) = -\ln \frac{Z_{\text{circuit}}[A]}{Z_{\text{circuit}}}
\]

\[
\approx -\ln \int \mathcal{D}[y(x)] \exp \left[ -\beta \sigma \int_{x_1}^{x_2} dx \sqrt{1 + (\partial_x y)^2} \right],
\]

where \( \beta \) is the “inverse temperature”, and \( \sigma \) the “surface tension”. This resulting “capillary-wave theory” partition function is the canonical ensemble of all domain walls (i.e. height functions \( y(x) \) defined in Eq. (19)), where the energy of each domain wall is the product of the surface tension and its surface area.

After expanding the square root and dropping higher-order irrelevant terms, Eq. (20) becomes a Gaussian theory, and can be readily evaluated. With details in Ap-
upon changing the b.c. of the upper edge (i.e. qubits in $Q$) explicitly that a “zoomed-in” view. In this way, we are assuming implicitly that $|A| \ll L$, and also that the circuit depth is large compared to the vertical extent of the domain wall, $T \gg \sqrt{L}$. It is within this regime that the approximation of $S(\rho_A)$ with $F_{CW}(A)$ in Eq. (21) is established, and is valid regardless of the other boundary conditions (as the domain wall is sufficiently far away from the other boundaries).

The subleading “entropic” correction of $S(\rho_A)$ was found to be characteristic of the mixed phase $0 < p < p_c$ of hybrid circuits \cite{16, 18}, and is now shown to be present generically whenever the fluctuating domain wall picture is valid, though its analytic form ($\frac{3}{2} \ln |A|$) here is special to capillary-wave theory. Its importance will be made clear below; notice, however, that this term is absent at $p = 0$ (random unitary circuits without measurements), or equivalently, at zero temperature of the spin model, where the domain wall cannot fluctuate (see Appendix B).

\section*{C. The maximally-mixed initial state}

We have seen in Sec. II C that the hybrid circuit dynamics with the maximally-mixed initial state can be formulated as the dynamics of the corresponding QECC. The entropy of the entire system $Q$, $S(\rho_Q)$, is monotonically decreasing, corresponding to a monotonically decreasing “code rate”. On the other hand, according to the prescriptions summarized at the beginning of this section (for mapping to a spin model), the corresponding circuit partition function $Z_{\text{circuit}}$ is defined by the fixed b.c. $a$ on both the upper and lower edges of the circuit; whereas the entropy $S(\rho_Q)$ is the change in free energy upon changing the b.c. of the upper edge (i.e. qubits in $Q$) to a different, fixed one, $b$. The b.c. relevant to this calculation is illustrated in Fig. 3. Since the maximally-mixed initial state admits a natural purification in terms of $|Q| = L$ Bell pairs where $Q$ consists of one qubit from each pair, the upper and lower edges can be naturally viewed as $Q$ (the system, that is acted upon by the circuit), and $R$ (the “reference”, consisting of the other half of the Bell pairs, that is left un-evolved by the circuit), respectively \cite{24}. We will henceforth adopt this labelling, for we find it intuitive to have a concrete reference $R$ at the far end of the circuit that $Q$ is trying to disentangle itself from, even if this choice of $R$ is not unique.

The dominant contribution to $S(\rho_Q)$ comes from a single domain wall separating the upper and lower edges, going around the “waist” of the circuit (again compare Fig. 3):

- With open spatial b.c., the domain wall endpoints are “free”, and can independently take any vertical coordinate $y(x = 0) \in [-T, 0]$ and $y(x = L) \in [-T, 0]$.

- With periodic spatial b.c., the domain wall is periodic, but otherwise “free” to take any position along the vertical direction, leading to $y(x = 0) = y(x = L) \in [-T, 0]$.

The free energies can then be calculated within capillary-wave theory (Appendix B),

\begin{equation}
S(\rho_Q) = -\ln \frac{Z_{\text{circuit}}[Q]}{Z_{\text{circuit}}} = F_{CW}(Q) = \begin{cases} \beta \sigma L - \ln T, & \text{open b.c.} \\ \beta \sigma L - \ln \frac{T}{\sqrt{L}}, & \text{periodic b.c.} \end{cases} \quad \text{when } T \gg \sqrt{L}. \tag{22}
\end{equation}

The $-\ln T$ term comes from the “center of mass entropy” of the “waist domain wall”, whose form is consistent with an exponentially long purification time within the mixed phase \cite{18} (see Sec. III E). The $\ln \sqrt{L}$ difference between open and periodic b.c. is attributed to the additional endpoint entropy with open b.c., as mentioned above.
We see also that the quantity $\beta \sigma$ can be identified as (\ln 2 times) the code rate,
\[
\lim_{|Q| \to \infty} \frac{k \ln 2}{|Q|} = \lim_{L \to \infty} \frac{S(\rho_Q)}{L} = \beta \sigma,
\] (23)
for $T = \text{poly}(L)$.

\section*{D. Decoupling of domain walls}

We are now ready to investigate the entropy of a contiguous subregion $A$ of $Q$ with arbitrary length. Notice that the previous result in Eq. (21) was obtained for $|A| \ll |Q| = L$, and that Eq. (22) accounts for the limiting case $|A| = |Q| = L$. These two regimes must then be interpolated by some intermediate behavior. For convenience, below, we will instead study the entropy $S(\rho_{\overline{A}})$, defined on the complement of $A$.

Consider first the limit with small $A$, $|A| \ll |Q|$. In this regime, the partition function $Z_{\text{circuit}}(\overline{A})$ (defined by $a$ on $AR$ and $b$ on $\overline{A}$) receives two possibly comparable contributions (see Fig. 4 with periodic b.c.):

1. A single domain wall separating $\overline{A}$ from $AR$ as before. There are then two domains, with spins aligned along $a$ and $b$, respectively (see Fig. 4(a)). The corresponding partition function is approximated within capillary wave theory as
\[
Z_{\text{circuit}}^{(1)}(\overline{A}) \approx Z_{\text{circuit}} e^{-FCW(\overline{A})},
\] (24)

2. Two “decoupled” domain walls, one separating $A$ from $\overline{A}$, and the other, a “waist domain wall”, separating $Q = A\overline{A}$ from $R$. There are now three domains, as shown in Fig. 4(b), and to go from $A$ to $R$ two domain walls must be crossed. The corresponding partition function is approximated within capillary wave theory as
\[
Z_{\text{circuit}}^{(2)}(\overline{A}) \approx Z_{\text{circuit}} e^{-FCW(A) - FCW(Q)}.
\] (25)

After summing these contributions, we have, according to Eq. (18),
\[
S(\rho_{\overline{A}}) \approx - \ln \left[ e^{-FCW(\overline{A})} + e^{-FCW(A) - FCW(Q)} \right].
\] (26)

The first contribution $F_{CW}^{(1)} = FCW(\overline{A})$ is always energetically more favorable than $F_{CW}^{(2)} = FCW(A) + FCW(Q)$, but is not necessarily entropically so. The competition is only present due to fluctuations of the domain walls.

To illustrate this, we evaluate Eq. (26) with periodic b.c. (using Eqs. (21, 22)), where $FCW$ is simply a function of the size of the region, and plot the result in Fig. 4(c). Notice the striking non-monotonic behavior in $S(\rho_{\overline{A}})$, which has a width labelled as $|A|^*$. The non-monotonicity comes from a competition between the two contributions, which we can readily understand for large $|Q| = L$,
\[
S(\rho_{\overline{A}}) \approx - \ln \left[ e^{-FCW(\overline{A})} + e^{-FCW(A) - FCW(Q)} \right] = \min \left\{ FCW(\overline{A}), FCW(A) + FCW(Q) \right\}
\approx \begin{cases} FCW(\overline{A}), & 0 \leq |\overline{A}| < L - |A|^* \\ FCW(A) + FCW(Q), & L - |A|^* < |\overline{A}| \leq L \end{cases}.
\] (27)

Here $|A|^*$ is the length scale when the entropic and energetic terms are comparable, and may be defined as follows,
\[
F_{CW}(L - |A|^*) = FCW(|A|^*) + FCW(L) \Rightarrow |A|^* = \frac{1}{2\beta \sigma} \left( \frac{3}{2} \ln L + \ln \frac{T}{\sqrt{L}} \right),
\] (28)
to leading order for large $T$ and $L$. The length scale $|A|^*$ is thus inversely proportional to the code rate $\beta \sigma$, and grows with both $L$ and $T$ logarithmically. For any circuit depth $T = O(\text{poly}(L))$, $|A|^*$ is proportional to $\ln L$.

In the regime with $L - |A|^* < |A| < L$ (i.e. $0 \leq |A| < |A|^*$), we recognize that the free energies $F_{CW}(A)$ and $F_{CW}(Q)$ in Eq. (27) represent the corresponding entanglement entropies $S(\rho_A)$ and $S(\rho_Q)$ according to Eqs. (21, 22). The last line in Eq. (27) can then be rewritten as,

$$0 \leq \frac{|A|}{|A|^*} \leq 1 \Rightarrow S(\rho_T) \approx S(\rho_A) + S(\rho_Q)$$

$$\Rightarrow S(\rho_{AR}) \approx S(\rho_A) + S(\rho_R)$$

$$\Rightarrow I_{A,R} \approx 0.$$ 

(29)

We thereby conclude that if $\frac{|A|}{|A|^*} < 1$, the subsystems $A$ and $R$ decouple. This decoupling corresponds to the regime where the configuration in Fig. 4(b) dominates, i.e. when the domain wall decouples, with two domain walls separating $A$ and $R$.

In Fig. 4(c) we have also plotted (half) the mutual information between $A$ and $R$, $I_{A,R}$, as computed from capillary-wave theory using Eqs. (14, 21, 26). Notice the (near) vanishing of $I_{A,R}$ for $0 \leq |A| < |A|^*$, consistent with Eq. (29).

A more detailed calculation shows that,

$$I_{A,R} \approx \ln \left[ 1 + e^{F_{CW}(A) + F_{CW}(Q) - F_{CW}(A)} \right]$$

$$\approx \ln \left[ 1 + e^{-2\beta \sigma(|A|^* - |A|)} \right]$$

$$\approx \begin{cases} 
\frac{2\beta \sigma(|A|^* - |A|)}{\ln \tanh \frac{1}{2}}, & |A| < |A|^*; \\
0, & |A| > |A|^*. 
\end{cases}$$ 

(30)

Here we have used Eq. (28), and only kept the leading linear terms in $F_{CW}$. Since $|A|^*$ diverges in the thermodynamic limit (see Eq. (28)), for $\frac{|A|}{|A|^*} \in [0, 1)$ the mutual information $I_{A,R}$ vanishes exactly. On the other hand, $I_{A,R}$ is strictly positive if $\frac{|A|}{|A|^*} > 1$.

Upon combining with Theorem 1 in Sec. IID, we conclude that $\ell_A = 0$ if and only if $|A| < |A|^*$. We can then make the important identification between the code distance $d_{\text{cont}}$ and $|A|^*$,

$$d_{\text{cont}} = |A|^*.$$ 

(31)

With this equality, we may deduce from Fig. 4(c) (see the figure caption) that

$$k = (\ln 2)^{-1}S(\rho_Q) \leq L - 2|A|^* = |Q| - 2d_{\text{cont}}.$$ 

(32)

This is essentially the quantum Singleton bound [4], with $d \rightarrow d_{\text{cont}} \gg 1$.

To summarize, capillary-wave theory predicts that the dynamically generated QECC has code distance that diverges with system size on relevant time scales, while also keeping a finite code rate. Qubit segments with length smaller than $|A|^* = d_{\text{cont}}$ are protected from undetectable errors by thermodynamic fluctuations of the entanglement domain walls.

E. Crossover to late times

In this subsection, we deviate from our main focus on polynomial time scales $T = O(\text{poly}(L))$, and briefly discuss how the domain wall picture can account for the late time crossover behavior when $T$ is exponential in $L$. On these time scales, the entropy of the code state $S(\rho_Q)$ is expected to decay to zero [18], i.e. the state is completely purified.

Previously, when computing $S(\rho_Q)$ from Fig. 3 obtaining the result in Eq. (22), we only took into account configurations with a single waist domain wall - valid since the energy term $\beta \sigma L$ is always dominant over the entropy term $-\ln T$ when $T = O(\text{poly}(L))$, and single-domain wall configurations have the lowest energy. This simplification eventually breaks down when $T \gg \exp[\beta \sigma L]$, and we have to consider the possibility of multiple waist domain walls. In particular, $Z_{\text{circuit}}$ will now receive contributions from all configuration with an even number of waist domain walls, and $Z_{\text{circuit}}(Q)$ from those with an odd number. Here we are again assuming the Ising nature of these domain walls.

Moreover, the vertical (i.e. time direction) extent of each domain wall scales as $\ell_A |A|^*$, consistent with Eq. (29).

$$I_{A,R} \approx \ln \left[ 1 + e^{F_{CW}(A) + F_{CW}(Q) - F_{CW}(A)} \right]$$

$$\approx \ln \left[ 1 + e^{-2\beta \sigma(|A|^* - |A|)} \right]$$

$$\approx \begin{cases} 
\frac{2\beta \sigma(|A|^* - |A|)}{\ln \tanh \frac{1}{2}}, & |A| < |A|^*; \\
0, & |A| > |A|^*. 
\end{cases}$$ 

(30)

Notice that

$$\lim_{T \rightarrow \infty} S(\rho_Q) = 0,$$ 

(34)

as expected for a pure state.

The same reasoning leads to a similar modification of $S(\rho_T)$ in Eq. (26),

$$S(\rho_T) \approx -\ln \left[ e^{-F_{CW}(A)} + e^{-F_{CW}(Q)} \right] \tanh \left( e^{-F_{CW}(Q)} \right),$$ 

(35)
which implies,
\[
\lim_{T \to \infty} S(\rho^A) = -\ln \left[ e^{-F_{CW}(A)} + e^{-F_{CW}(A^*)} \right] \\
\approx \min \{ F_{CW}(A), F_{CW}(A^*) \},
\]
again as expected for the pure state $\rho_Q$ that is dynamically generated on exponentially long times.

Inclusion of multiple domain walls also introduces some crossover time dependence in $d_{cont}$, accounting for its eventual linear scaling in $L$ when $T \gg \exp[\beta\sigma L]$. Indeed, the decoupling conditions Eqs. (29, 30) retain their forms in this limit, and $d_{cont}$ can still be identified with $|A|^*$, which approaches $L/2$ in the long time limit.

IV. NUMERICAL RESULTS

In this section, we compare our capillary-wave theory results with numerical computations in the random Clifford circuit (Fig. 1) for the observables explored in the previous section. Overall, we find qualitative agreement between the two, but as we shall see, a complete quantitative agreement is lacking. We interpret the former as support for the general entanglement domain wall picture, and the latter as an indication of a more complex nature of these domain walls for the random Clifford circuit.

A. Code distance for Clifford QECC’s

The most striking qualitative prediction of the domain wall picture from Sec. III is the phenomenon of decoupled domain walls as illustrated in Fig. 4. To explore this for random Clifford circuits, we compute $S(\rho^A)$ and $I_{A,R}$ with varying $|A|$, taking a maximally-mixed initial state and averaging over the random ensemble of circuits, as well as over a time window $6L < T < 8L$. Within this time window, we have $T \gg L^{1/2}$, so that Eqs. (21, 22) should apply.

Our numerical results are shown in Fig. 5(a) for $L = 1024$ and $p = 0.08 \approx 0.5p_c$. Strikingly, we observe the same non-monotonicity in $S(\rho^A)$, decreasing with $|A|$ in the range $L - |A|^* < |A| \leq L$, in accordance with the capillary-wave theory results in Fig. 4. Moreover, within this range, $I_{A,R}$ is very small, showing a plateau with height $\approx 0$. The Clifford numerical results for both $S(\rho^A)$ and $I_{A,R}$ are thus fully consistent with the domain wall decoupling results in Fig. 4. Evidently, the domain wall picture holds for random Clifford circuits, being qualitatively consistent with capillary-wave theory.

Our particular choice of $p = 0.08$ was unimportant. Indeed, for the Clifford circuit we find consistency with Fig. 4 for a wide range of $p$ with $0 < p < p_c$ (not shown). This is as expected, since the domain wall picture should be valid at any “temperature” $p$ below the “critical temperature” $p_c$ of the spin model, i.e. throughout the “ordered phase”.

We also explore finite size effects on $I_{A,R}$ (see Fig. 5(b)). For a fixed $|A| < |A|^*$, we find that $I_{A,R}$ decreases with increasing system size $L$, consistent with Eq. (30). We therefore expect that in the thermodynamic limit, $I_{A,R} = 0$ if and only if $|A| < |A|^*$; the identification between $|A|^*$ and $d_{cont}$ can then be made. In a finite system, we define $|A|^*$ as the size of $A$ for which $I_{A,R} \approx \epsilon$, where $\epsilon$ is a small number independent of $L$. 

![Fig. 5](attachment:image.png)
Scaling of algebraic definition of distance also grows with $L$ consistently with capillary-wave theory. The code distance $d$ and the time window $6$ in agreement with a direct computation (from the algebraic definition of $d$) in Ref. [18]. The power-law dependence on the system size, $d_pbc(\rho_Q) - d_{obc}(\rho_Q) = \zeta \ln L$, with $\zeta \approx 0.56$. This difference has a weak time dependence, but not displayed here. (b) The halfcut mutual information $I_{A,\overline{A}}$ for $|A| = |\overline{A}| = L/2$ as a function of $L$, with periodic b.c., where we find $I_{A,\overline{A}} \propto L^{0.36}$ with $\gamma_2 \approx 0.36$.

With this identification, we may now examine how $d_{cont} = |A|^\gamma$ depends on the system size (as obtained from Fig. 5(a,b)). As shown in Fig. 5(c), we find that the code distance $d_{cont}$ increases with increasing $p$, qualitatively consistent with capillary-wave theory. The code distance also grows with $L$, but as a power-law function, $d_{cont} \propto L^{\gamma_1}$. The exponent is estimated to be $\gamma_1 \approx 0.38$, in agreement with a direct computation (from the algebraic definition of $d_{cont}$) in Ref. [18]. The power-law scaling of $d_{cont}$ quantitatively differs from capillary-wave theory in Eq. (28), where a logarithmic scaling was found.

B. Clifford dynamics versus (generalized) capillary-wave theory

We next numerically compute a few more quantities that we can compare with capillary-wave theory, as shown in Fig. 6. Once again, these results were obtained for the random Clifford circuit with a maximally-mixed initial state, upon averaging over both circuit realizations and the time window $6L < T < 8L$.

In Fig. 6(a), we plot the difference between $S(\rho_Q)$ with periodic and open boundary conditions, $\Delta S(\rho_Q) := S_{pbc}(\rho_Q) - S_{obc}(\rho_Q)$. Capillary-wave theory (Eqs. (21, 22)) predicts cancellations of the time dependence as well as of the “surface energy” term, leaving only the extra endpoint entropy term, $(1/2) \ln L$. This logarithmic scaling is indeed observed numerically, with the coefficient of $\ln L$ given by $\approx 0.56$, close in value to that of capillary-wave theory. We have also confirmed a very weak $T$-dependence of $\Delta S(\rho_Q)$, but the data is not displayed here.

In Fig. 6(b), we plot the “halfcut mutual information” [18], $I_{A,\overline{A}}$ with $|A| = |\overline{A}| = L/2$ versus $L$ with periodic b.c. Upon varying $L$, we find $I_{A,\overline{A}} \propto L^{\gamma_2}$ with $\gamma_2 \approx 0.36$, and the overall amplitude having a weak dependence on $p$. As for the code distance in Fig. 5(c) which grows with a similar power $\gamma_1 \approx \gamma_2$, this power-law scaling is quantitatively different from capillary-wave theory. The latter predicts a logarithmic scaling, $I_{A,\overline{A}} = (7/2) \ln L$ for $T \propto L$.

To account for the power-laws in Fig. 5(c) and Fig. 6(b), we introduce a phenomenological description, which we call “generalized capillary-wave” (GCW), with the following (minimal) modifications of the free energies for “pinned” and “waist” domain walls (Eqs. (21, 22)), respectively

$$F_{GCW}(A) = \beta \sigma |A| + \chi |A|^\gamma, \text{ when } T \gg L^\delta \gg |A|^\lambda,$$  \hspace{1cm} (37)

$$F_{GCW}(Q) = \left\{ \begin{array}{ll} \beta \sigma L - \ln T, & \text{open b.c.} \quad \text{when } T \gg L^\delta. \\ \beta \sigma L - \ln \frac{T}{\lambda}, & \text{periodic b.c.} \end{array} \right.$$

Here $0 \leq \gamma < 1$ is the exponent characterizing domain wall free energies in GCW, and $0 < \zeta < 1$ is the exponent of vertical extent of the domain walls.\(^{10}\) The constant $\chi$ is expected to be independent of $|A|, L, T, \beta \sigma$. Capillary-wave theory thus has $\zeta_{CW} = 0$ and $\gamma_{CW} = 1/2$; compare Eqs. (21, 22). This generalization of capillary-wave theory remains qualitatively consistent with Fig. 5(a,b) and Fig. 6(a), where we found $\zeta \approx 0.56$.

Eqs. (37, 38), together with the definition of $|A|^\gamma$ in Eq. (28),\(^{11}\) lead to the following scaling behaviors for the code distance and half-cut mutual information,

$$d_{cont} = |A|^\gamma \approx \frac{\chi}{2\beta \sigma} L^\lambda, \hspace{1cm} (39)$$

$$I_{A,\overline{A}} \propto L^\lambda. \hspace{1cm} (40)$$

These are both consistent with our Clifford numerics in Fig. 5(c) and Fig. 6(b), provided we take $\gamma = \gamma_1 = \gamma_2$.

\(^{10}\) Notice that with $0 < \zeta < 1$, we still have $T \gg L^\delta$ for $6L < T < 8L$, the time window we took in the numerics. As we saw in Fig. 6(a), the exponent $\zeta \approx 0.56$ falls within this range, and seems to be close in value to $\zeta_{CW} = 1/2$.

\(^{11}\) We note that the form of Eq. (30) is identical for capillary-wave theory and its generalization in Eqs. (37, 38), since in its derivation we only kept the leading linear term, which is common for both cases. Thus the identification between $|A|^\gamma$ and $d_{cont}$ can still be made for GCW.
We emphasize that Eqs. (37, 38) are phenomenological, motivated by both capillary-wave theory and our Clifford numerics (specifically Fig. 5(c) and Fig. 6(b)). At this moment we do not have a theory from which these free energies can be derived.

We note that direct numerical computations of $S(\rho_A)$ (for $|A| \ll L$) and $S(\rho_Q)$ are qualitatively consistent with both capillary wave theory (Eqs. (21, 22)) and its generalization in Eqs. (37, 38), as established in Refs. [16, 18]. In particular,

- For the approximation $S(\rho_A) \approx F_{CW}(A)$ when $|A| \ll L$, the “linear plus log” form of $F_{CW}(A)$ is consistent with the stabilizer length distribution [16];
- For the approximation $S(\rho_Q) \approx F_{CW}(Q)$, the $-\ln T$ dependence on circuit depth is consistent with an exponentially long purification time [18].

On the other hand, a quantitative comparison between capillary-wave theory and GCW is tricky, due to the difficulty in distinguishing a logarithmic function from a small power-law in the presence of a background linear term. Thus, we will not here attempt to compare capillary-wave theory and GCW for the quantities $S(\rho_A)$ (with $|A| \ll L$) and $S(\rho_Q)$.

V. DISCUSSION

A. Summary

In this paper we established a correspondence between QECCs generated by random hybrid Clifford circuit dynamics, and the statistical mechanics of fluctuating “entanglement domain walls”. The number of encoded logical qubits $k$ of the QECC maps to the “surface energy” that is extensive in the number of physical qubits $|Q|$, and the code distance maps to a crossover length scale proportional to the “entropy” of transverse fluctuations, that is subextensive in $|Q|$. Fluctuations of entanglement domain walls are entirely responsible for the diverging code distance, which protects the state against local (undetectable/uncorrectable) errors, a characteristic property of QECCs.

Our results rest upon two well-motivated assumptions, namely the validity of the entanglement domain wall picture, and the “linear plus sublinear” form of their free energies. The former has been analytically established in the context of random Haar circuits [19, 20], and the latter follows from the former within capillary-wave theory. We expect that both assumptions are also valid for Clifford circuits, as supported by the Clifford numerics in Refs. [16, 18], as well as those in Sec. IV.

We emphasize that the qualitative properties of the QECC do not depend crucially on the specific form of the entropic term, which diverges logarithmically with $|Q|$ in capillary-wave theory, and as a small power-law in our Clifford numerics. The latter is possibly described by a certain generalization of capillary-wave theory. In some sense, one can view capillary-wave theory as a “mean-field theory” of the entanglement domain walls.

B. The diverging code distance as a self-consistency condition

The error-correcting nature of the dynamically generated state, as exemplified by the diverging code distance, is consistent with the resilience of this finite entropy-density (and code rate) state to repeated local measurements. Indeed, a measurement decreases the entropy only if the measured operator is a non-trivial logical operator (see Sec. II C), and with a diverging code distance the probability of each local measurement in the circuit (Fig. 1) being a logical operator (denoted $p_{\text{logical}}$) vanishes in the thermodynamic limit (equivalently, each qubit in $Q$ decouples from the reference state $R$ with probability one). We can estimate $p_{\text{logical}}$ by setting $|A| = 1$ in Eq. (30), giving $p_{\text{logical}} \propto \ell_A I_{A,R} \approx \exp[-2\beta\sigma_{\text{cont}}]$, leading to $p_{\text{logical}} \propto L^{-2}$ within capillary-wave theory for $T \propto L$, and $p_{\text{logical}} \propto \exp[-\chi L^7]$ within a generalized capillary-wave description. In either case, this leads to a vanishing rate of purification in the mixed phase when $O(L)$ measurements are made in each time step [18], and subsequently to the stability of the finite code rate.\footnote{When $T$ is allowed to be independent of and longer than $L$, capillary-wave theory gives, $p_{\text{logical}} \propto (LT)^{-1} \Rightarrow \frac{dS(\rho_Q)}{dT} \approx -(pL)p_{\text{logical}} \propto T^{-1}$, consistent with Eq. (22) and an exponentially long purification time (see Sec. III E).}

Our discussion above is not an explanation of the stability of the mixed phase, but a requirement of self-consistency, since the diverging code distance is itself computed from the steady state within the mixed phase. The domain wall picture itself also requires the assumption of an ordered phase.

Moreover, a quantum Hamming bound [7, 43] on $p_c$, as in Ref. [23], cannot be inferred from our discussion. Besides the code distance being subextensive rather than extensive, here we are viewing the one-qubit measurements within one circuit time step as a sequence of one-qubit errors, rather than a single $p(|Q|)$-qubit error. With respect to these single qubit errors, the code is highly degenerate, and the Hamming bound does not apply.

Finally, we mention that QECCs can also be dynamically generated for circuits with a pure initial state in the volume law entangled phase when $p < p_c$ [17, 23], if we take the “system” $Q$ to be an extensive subsystem, $R$ to be the complement of $Q$ with $|R| > |Q|$, and consider the decoupling of $A \subset Q$ from $R$ (see Fig. 7). Indeed, in this...
accounting for the power-law dependences in d. This result suggests that disorder could dramatically modify the structure of the domain walls, possibly even being entirely absent. In the regime where A < d_{cont}, the dominant domain wall configuration is the “rainbow diagram” shown in (b), implying that A and R should fully decouple (for |Q| large), with vanishing mutual information. In this regime, an error on A will have no effect on |S(\rho_A)|.

C. The role of disorder

As for random Haar circuits [19, 20] and random tensor networks [25], the identification in Eq. (18) is between free energies in the stat. mech. model and entanglement entropies averaged over an ensemble of circuits. Thus we have been studying the averaged entropies, and comparing them with (generalized) capillary-wave theory. Capillary-wave theory assumes translational symmetry by construction, with no reference to sample-to-sample fluctuations or the role of disorder. We note that similar scaling behaviors have been found in random unitary circuits without measurements [49–52]. Without an analytic theory, we cannot determine if the entanglement domain walls are indeed DPRM-like. In this context, it could be interesting to find “clean” circuit models, for which effects of quenched disorder are absent, so that the subleading “entropic” term only receives contribution from thermal fluctuations, just like simple Ising domain walls. These open issues are left for future work.

D. Outlook

We have made extensive use of the random Clifford circuit and the stabilizer formalism to establish our results. It would be interesting to see if the entanglement domain wall picture is valid in a broader class of models, such as hybrid random Haar circuits [13, 19, 20, 22, 53], or other “generic” models of non-unitary dynamics [54–63], or circuit models with measurements only [33, 64–66].

While a general QECC does not necessarily have “locality”, spatial locality and spatial dimensionality is important in the hybrid circuit dynamics. For this reason, we have been considering the “contiguous code distance”, rather than the conventionally defined code distance in the QECC context. We have also restricted our attention to (1+1)-dimensional circuits, in which the domain walls are one-dimensional objects in a two-dimensional background. It would be interesting to test this picture in higher dimensions [13], in tree-like geometries [67], or even in “all-to-all” models [18, 68] where locality is entirely absent.

The dynamically generated QECCs are found to have a finite code rate and a subextensive code distance. They are therefore not “good codes” in the conventional sense [5], which have a finite code rate and an extensive...
code distance. For example, a good code can be obtained by running a random unitary circuit without measurements into the steady state, starting from a mixed state with a finite entropy density. It would be interesting to see if the domain wall picture sheds any light on these good codes, and more generally on QECCs that are not dynamically generated and/or stabilizer-based [69], e.g. those commonly used in an error correction context (see for example Ref. [70] for a recent survey).

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Appendix A: More on stabilizer codes

1. Entanglement entropies of stabilizer codes

Let the stabilizer group $S$ be an abelian subgroup of $\mathcal{P}(Q)$ as defined in Eq. (4). Let $\rho_Q(S)$ be the corresponding stabilizer code state as in Eq. (6) [30],

$$\rho_Q(S) = 2^{-|Q|} \sum_{g \in S} g.$$  \hfill (A1)

We can directly compute its Rényi entropies [30],

$$(\ln 2)^{-1} S^{(n)}(\rho_Q(S)) = \frac{1}{1-n} \log_2 \text{Tr} [(\rho_Q(S))^n]$$

$$= \frac{1}{1-n} \log_2 \text{Tr} \left[ \left( 2^{-|Q|} |S| \right)^{n-1} \rho_Q(S) \right]$$

$$= |Q| - \log_2 |S|. \hfill (A2)$$

Since this result is independent of the Rényi index $n$, we will suppress it henceforth. As in Sec. II, we take

$$|S| = 2^m \hfill (A3)$$

and define

$$k := |Q| - m = (\ln 2)^{-1} S(\rho_Q(S)). \hfill (A4)$$

Given a bipartition of the system, $A \subset Q$, $\overline{A} = Q - A$, we define the following group homomorphism

$$\text{proj}_{\overline{A}} : \mathcal{P}(Q) \rightarrow \mathcal{P}(\overline{A})$$

$$g_A \otimes g_{\overline{A}} \mapsto g_{\overline{A}} \hfill (A5)$$

We take the following reduced density matrix on $A$,

$$\rho_A(S) = \text{Tr}_{\overline{A}} [\rho_Q(S)]$$

$$= 2^{-|Q|} \sum_{g \in S} \text{Tr}_{\overline{A}} (g)$$

$$= 2^{-|Q|} \sum_{g \in S \cap \ker \text{proj}_{\overline{A}}} g$$

$$= 2^{-|Q|} \sum_{g \in S_A} g, \hfill (A6)$$
where we noticed that $\text{Tr}_\mathcal{F}(g)$ is nonzero only if $\text{proj}_\mathcal{F}(g) = 1$; and defined $\mathcal{S}_A$ to be the following subgroup of $\mathcal{S}$,

$$
\mathcal{S}_A := \mathcal{S} \cap \text{Ker} \text{proj}_\mathcal{F}.
$$

Thus, we have

$$
\mathcal{S}_A \cong \frac{\mathcal{S}}{\text{proj}_\mathcal{F}(\mathcal{S})},
$$

$$
|\mathcal{S}_A| = \frac{|\mathcal{S}|}{|\text{proj}_\mathcal{F}(\mathcal{S})|}.
$$

and from Eqs. (A2, A6),

$$
(\ln 2)^{-1} S(\rho_A(\mathcal{S}))
= |A| - \log_2 |\mathcal{S}_A|
= |A| - \log_2 |\mathcal{S}| + \log_2 |\text{proj}_\mathcal{F}(\mathcal{S})|.
$$

2. Proof of Theorem 1 in Sec. II D

Recall that $\mathcal{C}(\mathcal{S}) \subset \mathcal{P}(Q)$ is the abelianized centralizer of $\mathcal{S}$ in $\mathcal{P}(Q)$. Recall also that the group of logical operators is defined as the quotient group $\mathcal{L} = \mathcal{C}(\mathcal{S})/\mathcal{S}$ (see Sec. II B). The homomorphism $\text{proj}_\mathcal{F}$ naturally induces the following homomorphism between quotient groups,

$$
\widetilde{\text{proj}}_\mathcal{F} : \mathcal{L} \to \frac{\text{proj}_\mathcal{F}(\mathcal{C}(\mathcal{S}))}{\text{proj}_\mathcal{F}(\mathcal{S})},
$$

$$
g \cdot \mathcal{S} \mapsto \text{proj}_\mathcal{F}(g) \cdot \text{proj}_\mathcal{F}(\mathcal{S})
$$

It can be straightforwardly verified that this homomorphism is well defined, and is in fact surjective,

$$
\widetilde{\text{proj}}_\mathcal{F}(\mathcal{L}) = \frac{\text{proj}_\mathcal{F}(\mathcal{C}(\mathcal{S}))}{\text{proj}_\mathcal{F}(\mathcal{S})}.
$$

Recall that the group $\mathcal{L}_A$ is defined in Sec. II B as follows

$$
\mathcal{L}_A := \{ g \in \mathcal{C}(\mathcal{S}) \mid \text{proj}_\mathcal{F}(g) \in \text{proj}_\mathcal{F}(\mathcal{S}) \},
$$

where $\text{proj}_\mathcal{F}$ is understood as from $\mathcal{P}(Q)$ to $\mathcal{P}(\mathcal{A})$, as in Eq. (A5). It follows from the definitions that

$$
\mathcal{L}_A = \text{Ker} \ \widetilde{\text{proj}}_\mathcal{F} \subset \mathcal{L},
$$

thus

$$
|\mathcal{L}_A| = \left| \text{Ker} \ \widetilde{\text{proj}}_\mathcal{F} \right|
= \frac{|\mathcal{L}|}{|\widetilde{\text{proj}}_\mathcal{F}(\mathcal{L})|}
= \frac{|\mathcal{C}(\mathcal{S})| \cdot |\text{proj}_\mathcal{F}(\mathcal{S})|}{|\mathcal{S}| \cdot |\text{proj}_\mathcal{F}(\mathcal{C}(\mathcal{S}))|}.
$$

In the following, we associate these factors with entanglement entropies, using Eq. (A9).

We state without proof that an arbitrary generating set of $\mathcal{S}$ can be extended into one of $\mathcal{C}(\mathcal{S})$ [8, 31]:

$$
\mathcal{G}_S = \{ g_1, \ldots, g_m \},
$$

$$
\mathcal{G}_{\mathcal{C}(\mathcal{S})} = \{ g_1, \ldots, g_m, h_1^x, \ldots, h_k^x, h_1^z, \ldots, h_k^z \}.
$$
Each of $\mathcal{G}_S$ and $\mathcal{G}_{\mathcal{C}(S)}$ is a set of independent operators in $\mathcal{P}(Q)$; thus
\begin{equation}
|S| = 2^{\lvert \mathcal{G}_S \rvert} = 2^m, \tag{A17}
\end{equation}
\begin{equation}
|\mathcal{C}(S)| = 2^{\lvert \mathcal{G}_{\mathcal{C}(S)} \rvert} = 2^{m+2k} = 2^{|Q|+k}. \tag{A18}
\end{equation}
Each of $\{g_1 \ldots m\}$, $\{h^X_1 \ldots k\}$, and $\{h^Z_1 \ldots k\}$ is a set of mutually commuting operators in $\mathcal{P}(Q)$. In addition, the $g$’s commute with the $h^X$’s as well as with the $h^Z$’s; and $h^X_i h^Z_j = (-1)^{\delta_{ij}} h^Z_i h^X_j$. The $h$ operators can be thought of the so-called “representative logical $X$- and $Z$-operators”.

Next, we construct a purification of the state $\rho_Q(S)$. Let $R$ be a system of $k$ qubits, and let $\tilde{S} \subset \mathcal{P}(QR)$ be generated by the following set $\tilde{G}$, obtained from $\mathcal{G}_{\mathcal{C}(S)}$ by “extending” its elements to $QR$,
\begin{equation}
\tilde{G} = \left\{ (g_j)_Q \otimes 1_R \middle| j = 1 \ldots m \right\} \cup \left\{ (h^X_j)_Q \otimes (X_j)_R \middle| j = 1 \ldots k \right\} \cup \left\{ (h^Z_j)_Q \otimes (Z_j)_R \middle| j = 1 \ldots k \right\}, \tag{A19}
\end{equation}
where $(X_j)_R$ is the Pauli $X$-operator on the $j$-th qubit of $R$; and similarly for $(Z_j)_R$. It is clear that $\tilde{G}$ is a set of independent, mutually commuting elements of $\mathcal{P}(QR)$, and thus defines a physical state on $QR$,
\begin{equation}
\rho_{QR}(\tilde{S}) = 2^{-|QR|} \sum_{g \in \tilde{S}} g. \tag{A20}
\end{equation}
Since $\lvert \tilde{G} \rvert = \lvert \mathcal{G}_{\mathcal{C}(S)} \rvert = \lvert Q \rvert + k = \lvert QR \rvert$, we have $\lvert \tilde{S} \rvert = 2^{|QR|}$, and from Eq. (A2)
\begin{equation}
(\ln 2)^{-1} S(\rho_{QR}(\tilde{S})) = 0. \tag{A21}
\end{equation}
Moreover, by construction,
\begin{align*}
\rho_Q(\tilde{S}) &:= \text{Tr}_R \left[ \rho_{QR}(\tilde{S}) \right] \\
&= 2^{-|QR|} \sum_{g \in \tilde{S}} \text{Tr}_R(g) \\
&= 2^{-|Q|} \sum_{g \in \tilde{S}} g \\
&= \rho_Q(S). \tag{A22}
\end{align*}
Therefore, $\rho_{QR}(\tilde{S})$ is a purification of $\rho_Q(S)$ on $QR$, as claimed.

On the other hand, let us compute the reduced density matrix on $R$,
\begin{align*}
\rho_R(\tilde{S}) &= \text{Tr}_Q \left[ \rho_{QR}(\tilde{S}) \right] \\
&= 2^{-|QR|} \sum_{g \in \tilde{S}} \text{Tr}_Q(g) \\
&= 2^{-R|} \mathbb{1}_R, \tag{A23}
\end{align*}
i.e. the maximally-mixed state on $R$, as expected. Thus we have
\begin{equation}
(\ln 2)^{-1} S(\rho_R(\tilde{S})) = |R| = k. \tag{A24}
\end{equation}
It is easy to verify that for $A \subset Q$ and $\overline{A} := Q - A$,
\begin{equation}
\text{proj}_\overline{A}(\mathcal{C}(S)) = \text{proj}_\overline{A}(\tilde{S}), \tag{A25}
\end{equation}
where $\text{proj}_\overline{A}$ on the LHS is understood as from $\mathcal{P}(Q)$ to $\mathcal{P}(\overline{A})$, and that on the RHS from $\mathcal{P}(QR)$ to $\mathcal{P}(\overline{A})$. Thus, using Eq. (A9), but now for $AR \subset QR$, $\overline{A} = Q - A = QR - AR$, and $\tilde{S}$, we have
\begin{align*}
(\ln 2)^{-1} S(\rho_{AR}(\tilde{S})) &= |AR| - \log_2 |\tilde{S}| + \log_2 |\text{proj}_\overline{A}(\tilde{S})| \\
&= |AR| - \log_2 |\tilde{S}| + \log_2 |\text{proj}_\overline{A}(\mathcal{C}(S))|. \tag{A26}
\end{align*}
Combining this equation with Eqs. (A9, A14), we have (compare Eq. (13))

\[
\ell_A = \log_2 |\mathcal{L}_A| = \log_2 |\mathcal{C}(S)| - \log_2 |\mathcal{S}| + \log_2 |\text{proj}_T(S)| - \log_2 |\text{proj}_T(C(S))|
\]

Thus, we have proven the result stated in Sec. II D, by constructing a particular purification of \(\rho_Q(S)\) using a particular generating set of \(\mathcal{C}(S)\). But this choice is really arbitrary, and there is no surprise that it should work. In fact, any purification of \(\rho_Q(S)\) on QR with \(|R| = k\) has a generating set of the form in Eq. (A19), and thus gives a generating set of \(\mathcal{C}(S)\).

### Appendix B: Capillary-wave theory calculations

We compute within capillary-wave theory the free energies of two types of domain walls: those with pinned endpoints, as in Fig. 2(b); and those with free endpoints that wrap around the “waist” of the circuit, as in Fig. 3.

#### 1. Domain walls with pinned endpoints

For the case in Fig. 2(b), we have (compare Eq. (20))

\[
F_{\text{CW}}(A) = -\ln \int \mathcal{D}[y(x)] \exp \left[ -\beta\sigma \int_{x_1}^{x_2} dx \sqrt{1 + (\partial_x y)^2} \right],
\]

where the functional integral over \(y(x)\) is over the following class of “height functions”

\[
y : [x_1, x_2] \rightarrow [-T, 0], \quad x \mapsto y(x),
\]

with the additional constraint that the endpoints are “pinned”, \(y(x_1) = y(x_2) = 0\). To regularize the path integral, we will however take \(y(x_1) = y(x_2) = \epsilon\) to be a small constant, which can be understood as the lattice spacing.

We first expand the square root,

\[
\sqrt{1 + (\partial_x y)^2} = 1 + \frac{1}{2} (\partial_x y)^2 + O\left( (\partial_x y)^4 \right),
\]

and neglect quartic and higher order terms in \((\partial_x y)\); these are irrelevant under a renormalization group transformation. Thus we have a Gaussian theory,

\[
F_{\text{CW}}(A) = -\ln \int \mathcal{D}[y(x)] \exp \left[ -\frac{\beta\sigma}{2} \int_{x_1}^{x_2} dx \left( 1 + \frac{1}{2} (\partial_x y)^2 \right) \right]
\]

The second term in this equation is the summation over all admissible configurations of paths/height functions \(y(x)\), and can be viewed as a “random walk” with “diffusion constant” \((\beta\sigma)^{-1}\). It is thus regarded as the “thermal entropy”
of transverse fluctuations of the domain walls. The magnitude of the fluctuation scales with \( x_{12} \) identically to that of a random walker, and can, for example, be quantified by the following quantity,

\[
\sqrt{\langle |y(\bar{x}) - y(x_1)|^2 \rangle} \propto \sqrt{\frac{|A|}{\beta \sigma}}, \tag{B6}
\]

where \( \bar{x} \equiv (x_1 + x_2)/2 \).

In the following we will, for convenience, treat the path integral in Eq. (B5) as a quantum mechanical transition amplitude, from which Eq. (B6) can also be deduced. However, we note that there are other ways to evaluate this integral, e.g. by solving the diffusion equation subject to the constraint \( y(x) \in [-T, 0] \).

We now “quantize” the path integral, with the spatial direction \( x \) viewed as “imaginary time”. We then have an imaginary time path integral of a free quantum particle with mass \( \beta \sigma \), confined within a potential well \( y \in [-T, 0] \),

\[
\exp [-F_{CW}(A) + \beta \sigma |A|] = \int_{y(x) \in [-T, 0], y(x_1) = y(x_2) = \epsilon} \mathcal{D}[y(x)] \exp \left[ -\frac{\beta \sigma}{2} \int_{x_1}^{x_2} dx \left( \partial_x y \right)^2 \right] = \langle y(x_2) | \exp \left[ -\hat{H}_{x_{12}} \right] | y(x_1) \rangle, \tag{B7}
\]

where the Hamiltonian is that of a “particle in box” problem,

\[
\hat{H} = \frac{\beta \sigma}{2M} + V(y), \quad \text{where} \quad V(y) = \begin{cases} 0, & -T \leq y \leq 0; \\ \infty, & \text{otherwise}. \end{cases} \tag{B8}
\]

The eigenstates and their corresponding energies are

\[
\phi_n(y) = \langle y | n \rangle = \sqrt{\frac{2}{T}} \sin \left( \frac{n \pi y}{T} \right), \quad y \in [-T, 0],
\]

\[
E_n = \frac{1}{2\beta \sigma} \left( \frac{n \pi}{T} \right)^2, \quad n = 1, 2, 3, \ldots \tag{B9}
\]

We expand Eq. (B7) in the eigenbasis,

\[
\exp [-F_{CW}(A) + \beta \sigma |A|] = \sum_{n=1}^{\infty} \langle y(x_2) | n \rangle \langle n | y(x_1) \rangle \exp [-E_n x_{12}] = \frac{2}{T} \sum_{n=1}^{\infty} \sin^2 \left( \frac{n \pi \epsilon}{T} \right) \exp \left[ -\frac{1}{2\beta \sigma} \left( \frac{n \pi \sqrt{x_{12}}}{T} \right)^2 \right]. \tag{B10}
\]

When \( \pi \sqrt{x_{12}}/T \ll 1 \), we may approximate the summation with the following integral over \( u = \frac{n \pi \sqrt{x_{12}}}{T} \),

\[
\exp [-F_{CW}(A) + \beta \sigma |A|] \approx 2 \int_{\pi \sqrt{x_{12}}/T}^{\infty} \frac{T}{\pi \sqrt{x_{12}}} \sin^2 \left( \frac{ue}{\sqrt{x_{12}}} \right) \exp \left( -\frac{u^2}{2\beta \sigma} \right),
\]

\[
\approx \frac{2}{\pi \sqrt{x_{12}}} \int_{0}^{\infty} du \sin^2 \left( \frac{ue}{\sqrt{x_{12}}} \right) \exp \left( -\frac{u^2}{2\beta \sigma} \right) \approx \frac{2e^2}{\pi (x_{12})^{3/2}} \int_{0}^{\infty} du u^2 \exp \left( -\frac{u^2}{2\beta \sigma} \right) = \frac{2e^2}{\pi |A|^{3/2}} \int_{0}^{\infty} du u^2 \exp \left( -\frac{u^2}{2\beta \sigma} \right) = \sqrt{\frac{2}{\pi}} e^2 (\beta \sigma)^{3/2} |A|^{-3/2}, \tag{B11}
\]
and thus (compare Eq. (21))

\[ F_{CW}(A) = \beta \sigma |A| + \frac{3}{2} \ln |A| + \text{const., when } |A| \ll T. \]  
(B12)

In arriving at this result, we made the following replacement in the integrand

\[ \sin^2 \left( \frac{ue}{\sqrt{x_{12}}} \right) \exp \left( -\frac{u^2}{2\beta \sigma} \right) \rightarrow \left( \frac{ue}{\sqrt{x_{12}}} \right)^2 \exp \left( -\frac{u^2}{2\beta \sigma} \right); \]  
(B13)

this is valid when

\[ \frac{\epsilon \sqrt{\beta \sigma}}{\sqrt{x_{12}}} \ll 1 \iff \epsilon \ll \sqrt{|A|/\beta \sigma}. \]  
(B14)

Physically, it means that the temperature cannot be too low, so that the transverse fluctuation of the domain wall is large compared to the “lattice spacing”, \( \epsilon \).

2. “Waist” domain walls

As shown in Fig. 3, the “waist” domain wall for open b.c. has two independent free endpoints; whereas for periodic b.c. the two endpoints must coincide, but otherwise free.

In the case of open b.c., let \( y_1 = y(x_1 = 0) \), and \( y_2 = y(x_2 = L) \). The analog to Eq. (B7) reads

\[
\exp [-F_{CW}(Q) + \beta \sigma |Q|] \\
= \int_{y(x) \in [-T,0]} D[y(x)] \exp \left[ -\frac{\beta \sigma}{2} \int_0^L dx \left( \partial_x y \right)^2 \right] \\
= \int_{-T}^0 dy_1 \int_{-T}^0 dy_2 \langle y_2 \exp [-HL] | y_1 \rangle \\
= \int_{-T}^0 dy_1 \int_{-T}^0 dy_2 \sum_{n=1}^{\infty} \langle y_2 | n \rangle \langle n | y_1 \rangle \exp [-E_n L] \\
= \frac{2}{T} \int_{-T}^0 dy_1 \int_{-T}^0 dy_2 \sum_{n=1}^{\infty} \sin \left( \frac{n\pi y_1}{T} \right) \sin \left( \frac{n\pi y_2}{T} \right) \exp \left[ -\frac{1}{2\beta \sigma} \left( \frac{n\pi \sqrt{L}}{T} \right)^2 \right] \\
= \sum_{n \text{ odd}} \frac{8T}{n^2 \pi^2} \exp \left[ -\frac{1}{2\beta \sigma} \left( \frac{n\pi \sqrt{L}}{T} \right)^2 \right] \\
\approx \frac{1}{2} \int_{\pi \sqrt{T/L}}^\infty T du \frac{8L}{\pi \sqrt{T \beta \sigma} u^2} \exp \left[ -\frac{u^2}{2\beta \sigma} \right] \\
= \frac{4\sqrt{L}}{\pi} \int_{\pi \sqrt{T/L}}^\infty du u^{-2} \exp \left[ -\frac{u^2}{2\beta \sigma} \right] \\
= \frac{4\sqrt{L}}{\pi} \left\{ \left. -u^{-1} \exp \left[ -\frac{u^2}{2\beta \sigma} \right] \right|_0^{\infty} - \int_{\pi \sqrt{T/L}}^\infty du (u^{-1}) \left( -\frac{u}{\beta \sigma} \right) \exp \left[ \frac{u^2}{2\beta \sigma} \right] \right\} \\
\approx \frac{4}{\pi^2} T,
\]  
(B15)

where we assumed \( T \gg \sqrt{L} \) throughout.
For periodic b.c., letting \( y = y(x_1 = 0) = y(x_2 = L) \), we have
\[
\exp[-F_{CW}(Q) + \beta\sigma|Q|] = \int_{y(x) \in [-T,0]} \mathcal{D}[y(x)] \exp \left[ -\frac{\beta\sigma}{2} \int_0^L dx (\partial_x y)^2 \right]
\]
\[
= \int_{-T}^0 dy \langle y | \exp \left[ -\hat{H} L \right] | y \rangle
\]
\[
= \int_{-T}^0 dy \sum_{n=1}^\infty \langle y | n \rangle \langle n | y \rangle \exp \left[ -E_n L \right]
\]
\[
= \sum_{n=1}^\infty \exp \left[ -\frac{1}{2\beta\sigma} \left( \frac{n\pi \sqrt{L}}{T} \right)^2 \right]
\]
\[
\approx \int_0^\infty \frac{T du}{\pi \sqrt{L}} \exp \left[ -\frac{u^2}{2\beta\sigma} \right]
\]
\[
\approx \frac{T}{\pi \sqrt{L}} \int_0^\infty du \exp \left[ -\frac{u^2}{2\beta\sigma} \right]
\]
\[
= \sqrt{\frac{\beta\sigma}{2\pi \sqrt{L}}}
\]
where we again assumed \( T \gg \sqrt{L} \) throughout.
Summarizing, we have (compare Eq. (22))
\[
F_{CW}(Q) = \begin{cases} 
\beta\sigma L - \ln T + \text{const.}, & \text{open b.c.} \\
\beta\sigma L - \ln \frac{T}{\sqrt{L}} + \text{const.}, & \text{periodic b.c. when } T \gg \sqrt{L}.
\end{cases}
\]
(B17)

Similarly to domain walls with pinned endpoints, the subleading logarithmic term can again be understood as coming from thermal entropies of transverse fluctuations. The \( \ln \sqrt{L} \) difference is the extra endpoint entropy in open b.c.

[1] P. W. Shor, Phys. Rev. A 52, R2493 (1995).
[2] A. M. Steane, Phys. Rev. Lett. 77, 793 (1996).
[3] C. H. Bennett, D. P. Divincenzo, J. A. Smolin, and W. K. Wootters, Phys. Rev. A 54, 3824 (1996), arXiv:quant-ph/9604024 [quant-ph].
[4] E. Knill and R. Laflamme, Phys. Rev. A 55, 900 (1997).
[5] A. R. Calderbank and P. W. Shor, Phys. Rev. A 54, 1098 (1996), arXiv:quant-ph/9602002 [quant-ph].
[6] A. Steane, Proceedings of the Royal Society of London Series A 452, 2551 (1996), arXiv:quant-ph/9601029 [quant-ph].
[7] D. Gottesman, Physical Review A 54, 1862 (1996), arXiv:quant-ph/9604038 [quant-ph].
[8] A. R. Calderbank, E. M. Rains, P. W. Shor, and N. J. A. Sloane, Phys. Rev. Lett. 78, 405 (1997), arXiv:quant-ph/9605005 [quant-ph].
[9] D. Gottesman, Stabilizer codes and quantum error correction, Ph.D. thesis, California Institute of Technology (1997), arXiv:quant-ph/9705052 [quant-ph].
[10] A. Y. Kitaev, Annals of Physics 303, 2 (2003), arXiv:quant-ph/9707021 [quant-ph].
[11] M. Levin and X.-G. Wen, Phys. Rev. Lett. 96, 110405 (2006), arXiv:cond-mat/0510613 [cond-mat.str-el].
[12] A. Almheiri, X. Dong, and D. Harlow, Journal of High Energy Physics 2015, 163 (2015), arXiv:1411.7041 [hep-th].
[13] B. Skinner, J. Ruhman, and A. Nahum, Physical Review X 9, 031009 (2019), arXiv:1808.05953 [cond-mat.stat-mech].
[14] A. Chan, R. M. Nandkishore, M. Pretko, and G. Smith, Phys. Rev. B 99, 224307 (2019), arXiv:1808.05949 [cond-mat.stat-mech].
[15] Y. Li, X. Chen, and M. P. A. Fisher, Physical Review B 98, 205136 (2018), arXiv:1808.06134 [quant-ph].
[16] Y. Li, X. Chen, and M. P. A. Fisher, Physical Review B 100, 134306 (2019), arXiv:1901.08092 [cond-mat.stat-mech].
[17] S. Choi, Y. Bao, X.-L. Qi, and E. Altman, arXiv e-prints, arXiv:1903.05124 [quant-ph].
[18] M. J. Gullans and D. A. Huse, arXiv e-prints (2019), arXiv:1905.05195 [quant-ph].
[19] Y. Bao, S. Choi, and E. Altman, Phys. Rev. B 101,
