Minimal instances for toric code ground states

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A decade ago Kitaev’s toric code model established the new paradigm of topological quantum computation. Due to remarkable theoretical and experimental progress, the quantum simulation of such complex many-body systems is now within the realms of possibility. Here we consider the question, to which extent the ground states of small toric code systems differ from LU-equivalent graph states. We argue that simplistic (though experimentally attractive) setups obliterate the differences between the toric code and equivalent graph states; hence we search for the smallest setups on the square- and triangular lattice, such that the quasi-locality of the toric code Hamiltonian becomes a distinctive feature. To this end, a purely geometric procedure to transform a given toric code setup into an LC-equivalent graph state is derived. In combination with an algorithmic computation of LC-equivalent graph states, we find the smallest non-trivial setup on the square lattice to contain 5 plaquettes and 16 qubits; on the triangular lattice the number of plaquettes and qubits is reduced to 4 and 9, respectively.

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The quest for topological quantum computation [1] relies on well controllable quantum systems exhibiting anyonic statistics. Several theoretical studies have proposed microscopic models featuring such topological phases; the most prominent example being the toric code [2]. While it is generally accepted that the fractional quantum Hall effect is well described as a topological state of matter [3–5], alternative setups such as cold atomic gases [6] and ion traps [7] are considered as suitable building blocks for a quantum simulation of topological matter. However, such systems naturally appear with a small number of involved particles. Regarding actual implementations it is therefore a fundamental question, how many particles the simulated system must involve in order to observe the characteristic properties of the system in question. In this letter, we address this question for the toric code and present setups with a minimal number of qubits in order to obtain non-trivial ground states.

In the last years great experimental progress led to the generation of highly entangled multi-qubit states in realistic settings, such as cold atomic gases in optical lattices [8, 9], Rydberg states [10–12], trapped ions [7, 13–15] and polarized photons [16–18]. On the theoretical side, a powerful tool in characterizing and describing states with multipartite entanglement is given by the concept of graph states [19, 20], and several realistic setups for the generation of graph states have been proposed [21–24].

In contrast to the coherent and measurement based state preparation above, the concept of quantum simulation [25, 26] of complex many-body Hamiltonians provides an alternative approach for the generation of topological phases as a suitable resource for topological quantum computation; the paradigm of which has been established by Kitaev’s toric code [2]. Subsequently several setups for the quantum simulation of the latter were discussed [6, 27–29]. Special focus was laid on the observation of mutual anyonic statistics for the elementary excitations [30–33], and experimental success within the realm of photonic quantum simulation was reported [34, 35].

However, the systems considered so far are small and their ground states tend to be well known multi-qubit states, e.g., the ground state of a single toric code plaquette is a simple GHZ state [36], as shown in Fig. 1 (A). Hence we argue that a proper characterization of toric code systems implies the comparison of its ground states with other known multi-qubit states, such as graph states.

In this letter, we address the question on the minimal

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Figure 1. (Color online) A: The (non-degenerate) ground state of a single toric code plaquette is a GHZ state and equivalent to the shown graph state. Qubits (○) are located on the edges. Vertices (●) denote star operators $A_s$ and grey faces plaquette operators $B_p$ acting on the adjacent qubits. The LC-equivalent graph state connects the qubits directly (—). B: The 8 neighbours (——) of the centered qubit (●) in the toric code on a square lattice. Neighbours and the centered qubit have a star and/or plaquette operator in common.
system required to observe such characteristic properties of the toric code. Let us explain this in more detail: The implementation of the toric code naturally gives rise to the concept of \textit{quasilocal} [37] interactions and nearest neighbour sites: The qubits are located on the links of a graph which is embedded into a surface, see Fig. 1 (B). The qubits connected to the same vertex \( s \) give rise to the quasilocal stabilizer \( A_s = \prod_{i \in s} \sigma_i^z \), while the qubits surrounding a plaquette \( p \) are involved in the many-body stabilizer \( B_p = \prod_{i \in p} \sigma_i^z \): the latter are qubits connected to a common vertex of the dual graph. This leads to a natural definition of nearest-neighbour (or \textit{vicinal}) qubits in the toric code, as qubits involved in the fundamental stabilizers \( A_s \) or \( B_p \). For a square lattice, the number of nearest neighbours is therefore eight, see Fig. 1 (B). It is a fundamental property of the toric code, that it is generated by \textit{quasilocal} stabilizer operators, and therefore, its experimental realization requires only \textit{quasilocal} interactions.

As mentioned above, for small systems the ground state of the toric code is equivalent to well known states: for example, the non-degenerate ground state of the setup with four qubits depicted in Fig. 1 (A) is equivalent to a GHZ state. Here, and throughout the manuscript, two states are termed “equivalent”, if they differ only by a sequence of single qubit rotations, that is, local unitary operations. On the other hand, GHZ states belong to the class of graph states [19, 20] which can be generated efficiently by applying two qubit operations (see Refs. [19, 38] and Chapter 1). As a stabilizer quantum code the toric code ground states are \textit{always} equivalent to a class of graph states. For the above example [Fig. 1 (A)], the depicted equivalent graph state exhibits only connections (i.e. edges) between qubits which are \textit{vicinal} within the toric code. However, this condition will break down for large setups: graph states, which are equivalent to the toric code ground states, will in general exhibit connections between qubits that are \textit{not} nearest-neighbours, and therefore their experimental realization requires non-local interactions. Hence a fundamental property of the toric code is the \textit{quasilocal} nature of its stabilizers – which distinguishes it from the equivalent \textit{non-local} graph states. Many applications of the toric code as topologically protected quantum memory rely on this local character.

A minimal setup for topological matter should inherit this crucial property. In this manuscript, we present minimal setups for the toric code, such that the equivalent graph states are \textit{non-local}, to wit, they require connections between qubits that are \textit{not} nearest-neighbours. A major obstacle is that the equivalence between the toric code ground states and graph states is not unique, since (in general) there are multiple equivalent graph states. As a consequence, we have to exclude the existence of any equivalent graph state with only local connections. We therefore first derive a simple, geometric procedure to construct equivalent graph states from a given toric code setup. In a second step, all equivalent graph states for promising candidates are generated and tested on their property of non-local connections.

The results for the most important setups are shown in Fig. 2: We find that for the common square lattice the minimal number of qubits is \( N = 16 \), see Fig. 2 (A), while for a triangular setup the minimal system reduces to \( N = 9 \) qubits, see Fig. 2 (B).

Finally, we have to point out that our main interest lies in equivalent, that is \textit{local unitary (LU) equivalent} graph states. However, computing LU-classes of graph states turns out to be exceedingly difficult whereas equivalence classes with respect to \textit{local Clifford (LC)} operations can be computed in principle for any number of qubits algorithmically [20, 39–41], though not efficiently. For graph states it is known [19] that up to 7 qubits LU- and LC-classes coincide. Unfortunately, the conjecture that this equivalence holds for arbitrary numbers of qubits (see [42] and references therein) was disproved by means of a 27-qubit counterexample [43]. However, it is believed [43] (but so far not proved) that this counterexample is the...
smallest one. Since the systems considered in this letter are well below this threshold, we will deal with LC-instead of LU-equivalent graph states in order to simplify computations.

This paper is organized as follows. In the first chapter (I) we summarize the mathematical and physical prerequisites used throughout this work: First, we outline the basics of the stabilizer formalism and give a short remark on different notions of “equivalent” states. Subsequently, two important classes of stabilizer states, graph states and surface codes, are reviewed in more detail. In the second chapter (II) we present our first result, proving that there is a simple geometrical procedure for the transformation of surface codes into LC-equivalent graph states. This procedure is illustrated by means of two examples. In the last chapter (III) we present our second result: the smallest toric code setups for which no local graph state exists. Readers familiar with the basics of graph theory, graph states and surface codes may skip the preliminaries and proceed with Chapter II. Experimentalists primarily interested in the proposed non-local setups may omit the second chapter, as it is only of technical interest for the setups in question.

I. PRELIMINARIES

The settings considered in this paper are composed of \( N \) qubits (or spins) and characterized by their coupling interactions. Thus the states are taken to be elements of the \( 2^N \)-dimensional Hilbert space \( \mathcal{H}^N := \bigotimes_{i=1}^N C_2^i \). The Pauli matrices acting on the Hilbert space of the \( i \)th qubit are denoted by \( \sigma_x^i, \sigma_y^i, \sigma_z^i \) and, for the sake of consistency, \( \sigma_0^i = 1 \). As a basis of the 1-qubit Hilbert space \( \mathcal{H}^i \) one chooses usually the computational basis \( \{ |0\rangle, |1\rangle \} \) defined as eigenbasis of \( \sigma^z \). Henceforth we shall use the convention \( \sigma^+ |0\rangle = |0\rangle \) and \( \sigma^+ |1\rangle = |−1\rangle \).

In this paper we consider two distinguished classes of \( N \)-qubit states known as surface codes and graph states. Both are examples for the much larger and intensively studied class of stabilizer states. Thus we will start with a short survey of the stabilizer formalism.

This formalism is widely used in quantum information theory and describes a certain class of \( N \)-qubit states, called stabilizer states, and their transformations by means of stabilizer groups.

To this end define the Pauli group \( G_1 := \text{span} \{ \sigma_x^i, \sigma_y^i, \sigma_z^i \} \) operating on a 1-qubit Hilbert space and generated by the Pauli matrices. The generalization for \( N \) qubits is defined as the \( N \)-fold tensor product \( G_N := \bigotimes_{i=1}^N G_1^i \) of the 1-qubit Pauli group. Now let \( \mathcal{G} = \{ g_i \}_{i=1}^d \subseteq G_N \setminus \{ −1 \} \) be a given set of independent, pairwise commuting Pauli operators. Then \( \mathcal{S} := \text{span} \mathcal{G} \) is called stabilizer of the protected subspace \( \mathcal{PS} = \{ |\Phi\rangle \in \mathcal{H}^N | \mathcal{S} |\Phi\rangle = |\Phi\rangle \} \). \( \mathcal{G} \) is the generating set of \( \mathcal{S} \) and \( d = |\mathcal{G}| := \text{rank} \mathcal{S} \) is called rank of the stabilizer. The dimension of the protected space is \( \text{dim} \mathcal{PS} = 2^{N−d} \) (for a proof and further reading see [44]), thus there are \( 2^{N−d} \) independent stabilizer states in \( \mathcal{PS} \).

It proves advantageous to describe \( N \)-qubit states \( |\Psi\rangle \in \mathcal{PS} \) by means of their stabilizing operators generated by \( \mathcal{G} \), for one has to track at most \( N \) generators whereas the description of general \( N \)-qubit states requires an overall number of \( 2^N \) complex coordinates. In order to describe unitary transformations of stabilizer states by transformations of their stabilizer, one may use only unitaries that leave \( G_N \) invariant under conjugation.

In this context, one defines the Clifford group \( C_1 := N_{U(2)}(G_1) = \{ U \in U(2) | U G_1 U^\dagger = G_1^1 \} \) as normalizer of the Pauli group in the 1-qubit unitary group \( U(2) \), and canonically extends this definition to the local Clifford group \( C_N^l := \bigotimes_{i=1}^N G_1^i \) operating on an \( N \)-qubit Hilbert space. If the transformations of a given stabilizer state are restricted to \( C_N^l \) (or more generally \( C_N \), see [45]), it is sufficient to transform the generators \( \mathcal{G} \rightarrow CGC^\dagger \) to describe the transformation \( C \in C_N^l \) in the stabilizer formalism since \( CGC^\dagger \) once more meets the requirements of a generating set.

Whereas in the stabilizer framework local Clifford operations play a crucial role, one generally deals with a larger class of local operations, namely local unitaries \( U \in U_N^l := \bigotimes_{i=1}^N U(2)^i \). Since two arbitrary multi-qubit states cannot be transformed into one another by local Clifford operations or even local unitaries in general, the class of \( N \)-qubit states decomposes into equivalence classes with respect to restricted sets of transformations.

To this end let \( |\Psi\rangle \) and \( |\Psi'\rangle \) be two arbitrary \( N \)-qubit states. They are called equivalent under local unitary operations (LU-equivalent or equivalent for short) if there exists \( U \in U_N^l \) such that \( |\Psi\rangle = U |\Psi'\rangle \); write \( |\Psi\rangle \sim_{\text{LU}} |\Psi'\rangle \). Accordingly one calls them equivalent under local Clifford operations (LC-equivalent) if there exists \( C \in C_N^l \) such that \( |\Psi\rangle = C |\Psi'\rangle \); write \( |\Psi\rangle \sim_{\text{LC}} |\Psi'\rangle \).

Since \( C_N^l \) is a proper subgroup of \( U_N^l \), it follows \( \text{LC} \implies \text{LU} \) but as \( C_N^l \neq U_N^l \) the converse cannot be true in general. However, due to the lack of counterexamples it was conjectured [42] that for the smaller class of stabilizer states \( \text{LU}- \) and LC-equivalence coincide: \( \text{LC} \equiv \text{LU} \). Although it was shown that this holds for a large class of stabilizer states [46, 47] the LU-LC-conjecture was disproved by Ji et al. in 2009 [43]. Nevertheless it was shown in Ref. [19] that \( \text{LU} \equiv \text{LC} \) holds for graph states with up to 7 qubits. Furthermore it is believed [43] (but so far not proved) that the counterexample used in Ref. [43] is the smallest one (\( N = 27 \)). Recently the class of stabilizer states for which the LU-LC-conjecture holds was enlarged [48], including a large class of surface codes.

After this brief survey of basic concepts of the stabilizer formalism and the notion of LC-equivalence we may now define a special class of stabilizer states, called graph states.

Graph states were introduced by Hein et al. [19] in 2004 and denote a class of \( N \)-qubit states that can be described by simple graphs. In this context it is shown [19] that the aforementioned LC-equivalence is closely...
related to the concept of local complementations known in graph theory. To state the concepts of graph states more precisely, basics of graph theory are required. Thus we start with a short review of the mathematics that may be skipped by the reader familiar with the concepts of graph theory.

A finite undirected graph is a pair $G = (V, E)$ where $V$ denotes a finite set of vertices and $E \subseteq \{\{a, b\} \mid a, b \in V\}$ a set of edges connecting these vertices. If there are neither loops (i.e. edges connecting a vertex to itself) nor multiple edges (i.e. vertices connected by more than just one edge) $G$ is called a simple graph. In contrast, if there are multiple edges (i.e. $E$ is a multiset) but no loops [49] $G$ is called a multigraph. In the following, simple graphs shall be denoted by upper-case letters $G, H, \ldots$ and multigraphs by calligraphic letters $\mathcal{L}, \mathcal{M}, \ldots$

For a given graph $G = (V, E)$ the functions $\mathcal{V}(G)$ and $\mathcal{E}(G)$ denote the sets of vertices and edges, i.e. $\mathcal{V}(G) = V$ and $\mathcal{E}(G) = E$. The order of a graph $G = (V, E)$ is the number of its vertices and denoted by $|G| := |V|$. One naturally defines the neighbourhood of a vertex $v \in V$ as the subset $N_v := \{w \in V \mid \{w, v\} \in E\} \subseteq V$ of vertices directly connected to $v$. For the sake of brevity we shall denote the complete graph over a vertex set $V$ by $\langle V \rangle := \{\{v, w\} \mid v, w \in V; v \neq w\}$, that is, the unique simple graph with edges connecting any pair of vertices.

Graph theory deals not only with properties of graphs but also with transformations of the latter. A particularly interesting transformation in the context of graph states is known as local complementation. The local complement $\tau_\circ (G)$ of a simple graph $G = (V, E)$ at vertex $v \in V$ is formally defined as $\tau_\circ (G) := G + \langle N_v \rangle$, where $+$ denotes the binary addition of edges. Practically the procedure described by $\tau_\circ$ reads as follows. Focussing exclusively on the neighbours of $v$ one deletes all edges in $E$ connecting these vertices and adds new edges between formerly unconnected neighbours. This yields $\tau_\circ (G)$.

In conclusion we note that two graphs $G = (V, E)$ and $H = (V, F)$ over the same vertex set $V$ are called equivalent under local complementations (LC-equivalent) if and only if there is a sequence of local complementations $\tau = \tau_{v_1} \circ \tau_{v_2} \circ \cdots \circ \tau_{v_n}$ such that $G = \tau (H)$; write $G \sim_{LC} H$. In the context of graph states, overloading “LC-equivalent” proves convenient and is therefore intended; this will become clear below.

We may now give the formal definition of graph states and comment on their properties. To this end let $G$ be an arbitrary simple graph with edges $E$ and vertices $V$. Consider a system of $N = |G|$ qubits – each identified with a vertex $v \in V$ – and define the (independent) operators

$$K^G_v := \sigma^v_u \prod_{w \in N_v} \sigma^w_v \in G_N.$$  
(1)

Then $G[G] = \{K^G_v \mid v \in V\}$ is a generating set of the graph state stabilizer $S[G] = \text{span} G[G]$. If there is no danger of confusion, we shall omit the dependence on $G$, viz. $S \equiv S[G]$. Since there are $N$ independent generators, the protected space $\mathcal{PS}[G]$ is one-dimensional. The state stabilized by $S[G]$, henceforth characterized by $G$, is called graph state and denoted by $|G\rangle$.

In this paper we shall define graph states in this stabilizer representation. However, there is an alternative representation which proposes a simple procedure for the construction of graph states in terms of controlled z-gates $U_{ab} = \mathcal{P}^{z+}_{ab} \otimes \mathbf{1}_b + \mathcal{P}^{z-}_{ab} \otimes \sigma_z^b$. Here $\mathcal{P}^{z\pm}_{v} = \frac{1}{2}[\mathbf{1}_v \pm \sigma^v_z]$ denote the projectors operating on qubit $v$. Then it can be shown [19] that

$$|G\rangle = \prod_{\{a, b\} \in \mathcal{E}(G)} U_{ab} |+\rangle^{\mathcal{V}(G)}$$  
(2)

where $|+\rangle = 1/\sqrt{2} (|0\rangle + |1\rangle)$ and $|+\rangle^{\mathcal{V}(G)}$ denotes the $N$-fold tensor product of states indexed by the vertices $\mathcal{V}(G)$. According to Eq. (2) the empty graph on $N$ vertices $V$ is represented by the product-state $|+\rangle^V$. One may add an edge between two vertices $a, b \in V$ by applying a controlled z-gate $U_{ab}$ on the corresponding qubits. This yields a handy procedure for the construction of any graph state $|G\rangle$ on $N$ qubits – edge by edge – provided an entropy free initial state $|+\rangle^{\mathcal{V}(G)}$.

Apart from their implementation, graph states are particularly interesting multi-qubit states since several physical properties can be related to graph theoretic properties of the underlying graph [19]. One of these properties is the aforementioned local Clifford equivalence: Let $|G\rangle$ and $|H\rangle$ be two graph states with corresponding graphs $G = (E, V)$ and $H = (F, V)$. Then they are LC-equivalent if and only if their graphs are connected by a sequence of local complementations. Formally this reads $|G\rangle \sim_{LC} |H\rangle \Leftrightarrow G \sim_{LC} H$. The notion of LC-equivalent graphs was known in graph theory for a while and it was shown that there is an efficient algorithm to check whether two graphs are LC-equivalent [39, 40]. Thus the aforementioned relation between the LC-equivalence of graph states and graphs provides an efficient algorithm to decide whether two graph states belong to the same LC-class.

For this reason the question arises whether this insight into the nature of LC-equivalence may be extended to a larger class of multi-qubit states. Indeed, it was shown [41, 50] that graph states are universal standard forms of stabilizer states. That is to say, for any $N$-qubit stabilizer state $|\Psi\rangle$ there exists an LC-equivalent graph state $|G_\Psi\rangle$. Since the transformation $|\Psi\rangle \xrightarrow{\text{LC}} |G_\Psi\rangle$ is efficiently computable, the problem of LC-equivalence for stabilizer states is reduced to the class of graph states – where it can be solved by means of mere graph theory.

Besides graph states the stabilizer formalism is widely used for the description of quantum codes. A prominent example we are particularly interested in is the toric code model (tcn) and its generalization, the surface codes.

In 2002 Kitaev proposed the qubit model named toric code [2] showing the inherent capability of quantum error correction on a physical level and a topologically ordered phase at zero temperature. Whereas the tcn is typically
defined on a square lattice embedded into a 2-torus, one may generalize this idea to arbitrary (multi-) graphs on orientable compact manifolds, called surface codes [2, 48, 51].

Aiming at the definition of surface codes, we need some notation regarding graph embeddings. To this end let \( \mathcal{L} \) be a multigraph and \( \Sigma \) a closed, orientable 2-manifold. A **graph embedding** \( \mathcal{X} \equiv \mathcal{X}_{\mathcal{L}, \Sigma} \) of \( \mathcal{L} \) into \( \Sigma \) can be informally understood as a “drawing” of \( \mathcal{L} \) on the surface \( \Sigma \) without intersecting edges. In the following “embedding” has always to be read as *2-cell embedding*, that is, a drawing on the surface, such that every face is homeomorphic to an open disc in \( \mathbb{R}^2 \) [52]. The **genus** \( g[\Sigma] \) of \( \Sigma \) is its number of “holes” or “handles”. The **genus** \( g[\mathcal{L}] \) of a graph is defined as the least genus of surfaces, such that an embedding exists. Each embedding \( \mathcal{X}_{\mathcal{L}, \Sigma} \) induces a set of faces denoted by \( \mathbb{P}_X(\mathcal{L}) \) or simply \( \mathbb{P}(\mathcal{L}) \). Given a graph embedding \( \mathcal{X}_{\mathcal{L}, \Sigma} \) the **dual graph** \( \mathcal{L}^* = (V^*, E^*) \) is defined so that \( V^* := \mathbb{P}_X(\mathcal{L}) \) and two dual vertices \( v^*_1, v^*_2 \in V^* \) are adjacent, i.e. \( \{v^*_1, v^*_2\} \in E^* \), iff there is a common edge in \( E(\mathcal{L}) \) to \( v^*_1 \) and \( v^*_2 \) as faces. A **(co-)cycle** \( \mathcal{C} \subseteq E(\mathcal{L}) \) \((\mathcal{C}^* \subseteq E(\mathcal{L}^*))\) on \( \mathcal{L} \) \((\mathcal{L}^*)\) is a set of edges, such that each vertex is adjacent to an even number of these edges. Are there at most two of these edges joining at each vertex and is \( \mathcal{C} \) \((\mathcal{C}^*)\) connected within the graph, then it is called a **simple (co-)cycle**, namely, a single cycle without self-intersections.

For the definition of surface codes, consider a multigraph \( \mathcal{L} \) 2-cell embedded via \( \mathcal{X}_{\mathcal{L}, \Sigma} \) into a surface \( \Sigma \) of genus \( g[\Sigma] \geq g[\mathcal{L}] \) and a system of \( |E(\mathcal{L})| \) qubits identified with the edges of \( \mathcal{L} \) (cf. graph states!). Define the **star** and **plaquette operators**

\[
A_s := \bigotimes_{i \in s} \sigma_i^x \quad \text{and} \quad B_p := \bigotimes_{i \in p} \sigma_i^z
\]

for each vertex \( s \in V(\mathcal{L}) \) and face \( p \in \mathbb{P}_X(\mathcal{L}) \). Here we identify a face or plaquette \( p \in \mathbb{P}_X(\mathcal{L}) \) with its set of peripheral edges and a vertex \( v \in V(\mathcal{L}) \) with its adjacent edges. Then \( \mathcal{S} \equiv \mathcal{S}[\mathcal{X}] := \text{span} \{A_s, B_p \mid s \in V(\mathcal{L}), p \in \mathbb{P}_X(\mathcal{L})\} \) is the stabilizer of the **surface code** corresponding to the chosen embedding \( \mathcal{X}_{\mathcal{L}, \Sigma} \). Its degeneracy is \( \dim\mathcal{PS} = 4^g = 2^{2g} \) in the case of closed boundary conditions [51] as one can show easily using Euler’s formula for 2-cell embeddings [53]. A possible basis of the protected space \( \mathcal{PS} \) is defined and manipulated by means of **loop operators**

\[
Z_k \equiv Z[\mathcal{C}_k] := \bigotimes_{i \in C_k} \sigma_i^z, \quad X_k \equiv X[\mathcal{C}_k^*] := \bigotimes_{i \in C_k^*} \sigma_i^x, \quad (4)
\]

where the simple (co-)cycles \( \mathcal{C}_k \subseteq \mathcal{L} \) and \( \mathcal{C}_k^* \subseteq \mathcal{L}^* \) are chosen homologically distinct for different \( k \) and non-trivial (see Fig. 3). In algebraic topology, two paths on a compact, orientable 2-manifold \( \Sigma \) are considered **homologous** if their union is the boundary of a surface embedded into \( \Sigma \). In our case, two cycles on the embedded graph \( \mathcal{L} \) are homologous iff their union is the boundary of a union of faces. This defines an equivalence relation \( \sim \) on the group of all cycles \( \mathcal{E}(\mathcal{L}) \). The quotient \( H_1 := \mathcal{E}(\mathcal{L})/\sim \) is called **first homology group** and contains homology classes \([\mathcal{C}]\) of cycles \( \mathcal{C} \in \mathcal{E}(\mathcal{L}) \). Restricted to the protected space \( \mathcal{PS} \), the effect of a loop operators \( Z[\mathcal{C}] \) \((X[\mathcal{C}^*])\) depends solely on the homology class of \( \mathcal{C} \) \((\mathcal{C}^*)\) since they act trivially if the (co-)cycle is homologously trivial, that is, \( \mathcal{C} \) \((\mathcal{C}^*)\) is the boundary of a union of faces (stars, i.e. dual faces). Therefore, by a slight abuse of notation, we may regard \( Z[\mathcal{C}] \) either as a mapping \( Z[\cdot] \) with domain \( \mathcal{E}(\mathcal{L}) \), or as an operator \( Z \) with domain \( H_1 \).

If the indices \( k \) are chosen properly, the loop operators fulfill the algebra \( \{Z_k, X_k\} = 0 = [Z_k, X_l] \) for \( k \neq l \). Then \( \{v \} \in \mathcal{PS} \mid Z_k[v] = v_k[v], 1 \leq k \leq 2g \} \) is a basis of \( \mathcal{PS} \), where the components of \( v \in \{-1, 1\}^{2g} \) are called **topological quantum numbers** and represent the logical qubits encoded in the protected space. The topological quantum numbers are flipped by the 2g distinct \( X_k \) loop operators via \( X_k\{v_1, \ldots, v_k, \ldots, v_{2g}\} = \{v_1, \ldots, -v_k, \ldots, v_{2g}\} \) due to the aforementioned (anti-)commutation relations.

To conclude this section, we note that in the special case of a square lattice \( \mathcal{L} \) and \( \Sigma = \mathbb{T}^2 \) a 2-torus, as depicted in Fig. 3, this construction yields the tcm [2].

II. MAPPING SURFACE CODES TO GRAPH STATES

A. Results

In the following, let \( |v\rangle \) be the state of a surface code with topological quantum numbers \( v \), correspond-
Let \( \mathcal{L} \) and show the consistency with well known facts. The in subsection \( \langle A \rangle \) are product states, symbolically \( |G\rangle \) yields the \( |\mathcal{X}\rangle \) \( \otimes |\mathcal{Y}\rangle \). Given the surface code \( |\mathcal{X}\rangle \) is LC-equivalent to disjoint graphs and GHZ states \( |\mathcal{B}\rangle \) are transformed to star graphs. A description of the graphical notation is given in Subsection II.B.

To this end a further notion of graph theory is required. Let \( \mathcal{L} \) and \( \mathcal{L}' \) be (multi-)graphs. \( \mathcal{L}' \) is called a spanning tree of \( \mathcal{L} \) if it is simple, connected, devoid of cycles and \( V[\mathcal{L}'] = V[\mathcal{L}] \) as well as \( E[\mathcal{L}'] \subseteq E[\mathcal{L}] \). That is to say, spanning trees are distinguished subgraphs without cycles that cover all vertices of the original graph. However, they are not unique in general. Spanning trees of a given graph may be constructed by deleting a set \( \mathcal{E} \subseteq E[\mathcal{L}] \) of edges such that there is no cycle left and the graph remains connected. For an example of this procedure see Fig. 4. Let us now introduce a mapping that converts multigraphs to a particular class of simple graphs which proves crucial for the LC-transformation of surface codes to graph states.

**Definition 1** \( (\varphi_{\mathcal{L}'} : \mathcal{L} \rightarrow G) \). Let \( \mathcal{L} \) be a multigraph and \( \mathcal{L}' \) one of its spanning trees. Then \( G = \varphi_{\mathcal{L}'}(\mathcal{L}) \) is a simple graph defined by the following construction: Set \( V(G) := E(\mathcal{L}) \) as the vertex set of \( G \). For \( r, s \in \mathcal{V}(G) \) the set \( \{r, s\} \) is an edge of \( G \) if and only if there is a path \( C_{pq} \) from \( p \) to \( q \) in \( \mathcal{L}' \) (\( p, q \in \mathcal{V}(\mathcal{L}') = V(\mathcal{L}) \)) such that \( p = \{r, q\} \) and \( s \in C_{pq} \).

The physical relevance of this mapping from multigraphs to simple graphs is given by the following

**Theorem 1** (LC-equivalence). Given the surface code state \( |\mathcal{V}\rangle \) on \( \mathcal{L} \) and any spanning tree \( \mathcal{L}' \). Then the graph state \( |G\rangle = |\varphi_{\mathcal{L}'}(\mathcal{L})\rangle \in \mathcal{H}^N \) is LC-equivalent to \( |\mathcal{V}\rangle \).

As a final remark note that the combination of the LC-rule for graph states \([19]\) and Theorem 1 yields the purely graph theoretic statement that for arbitrary spanning trees \( \mathcal{L}_1 \) and \( \mathcal{L}_2 \) it holds \( \varphi_{\mathcal{L}_1}(\mathcal{L}) \sim_{\text{LC}} \varphi_{\mathcal{L}_2}(\mathcal{L}) \). For the sake of consistency, we derived a direct graph theoretic proof for this very statement (which shall be omitted here).

![Figure 4](image)

**Example 1** (Product states). It is easy to see that protected states of two one-point connected surface code systems (Fig. 5 (A)) are product states, symbolically \( |\mathcal{C}\rangle = |\mathcal{C}\rangle \otimes |\mathcal{D}\rangle \). As a consequence of the one-point connection, any LC-equivalent graph state turns out to be disconnected – independent of the spanning tree used for the construction. Since disconnected graphs denote the tensor product of graph states \([20]\), we confirmed the statement \( |\mathcal{C}\rangle = |\mathcal{C}\rangle \otimes |\mathcal{D}\rangle \) in a purely geometric way, for local unitaries cannot alter the degree of entanglement.

**Example 2** (GHZ states). One can show easily that the protected state of a single plaquette bounded by \( N \) qubits is LC-equivalent to the \( N \)-qubit GHZ state \( |\text{GHZ}_N\rangle \) since it has the form \( |\mathcal{O}\rangle = \frac{1}{\sqrt{2}} \left( |+\rangle^\otimes \mathcal{N} + |\rangle^\otimes \mathcal{N} \right) \). Consider the hexagonal plaquette depicted in Fig. 5 (B). Constructing spanning trees is easy since “deleting” an arbitrary edge yields one. Consequently any graph state constructed this way must be a star graph. Indeed, in \([20]\) it is shown that star graphs (and complete graphs) are LC-equivalent to GHZ states.

![Figure 5](image)
C. Comments on the structure of $\varphi_{\mathcal{C}}(\mathcal{L})$

The geometrically constructed graph $\varphi_{\mathcal{C}}(\mathcal{L})$ exhibits an interesting graph theoretical structure, which in turn furnishes insight into the entanglement structure of the corresponding surface code state. To this end, we introduce the cycle space $\mathcal{E}(\mathcal{L})$ and its dual, the cut space $\mathcal{D}(\mathcal{L})$, as linear subspaces of the edge space $\mathcal{E}(\mathcal{L})$; the latter being the power set of edges $\mathcal{E}(\mathcal{L})$ equipped with binary addition and multiplication. The cycle space $\mathcal{E}(\mathcal{L})$ is defined as set of all cycles in $\mathcal{L}$ (as sets of edges) equipped with the binary structure inherited by $\mathcal{E}(\mathcal{L})$. A cut of $\mathcal{L}$ is a bipartition of its vertices; the cut-set of a given cut is the set of edges in $\mathcal{L}$ connecting these two disjoint sets of vertices (often the cut-set is just called cut). The cut space $\mathcal{D}(\mathcal{L})$ is then defined as set of all cut-sets on $\mathcal{L}$ equipped with the usual binary structure. Bases of both subspaces may be constructed by means of an arbitrary spanning tree $\mathcal{L}'$ as follows. For each edge $e = \{p, q\} \in \mathcal{E}(\mathcal{L}) \setminus \mathcal{E}(\mathcal{L}^e)$ choose the cycle $\mathcal{C}_e := \mathcal{C}_{pq} + e$ where $\mathcal{C}_{pq}$ denotes the unique path on the spanning tree connecting $p$ and $q$. $\mathcal{C}_e$ is called fundamental cycle on $\mathcal{L}$ with respect to $\mathcal{L}'$ and one can show easily that the set of all fundamental cycles $\{\mathcal{C}_e | e \in \mathcal{E}(\mathcal{L}) \setminus \mathcal{E}(\mathcal{L}^e)\}$ is a basis of the cycle space. Conversely, each edge $e' \in \mathcal{E}(\mathcal{L}^e)$ induces a bipartition $\mathcal{V}_e^+ \cup \mathcal{V}_e^- = \mathcal{V}(\mathcal{L})$ of the vertices, according to the component of the spanning tree they belong to if $e'$ is deleted. The corresponding cut-set $\mathcal{D}_{e'}$ is called fundamental cut of $e'$ with respect to $\mathcal{L}'$. Analogously the set of all fundamental cuts $\{\mathcal{D}_{e'} | e' \in \mathcal{E}(\mathcal{L}^e)\}$ yields a basis of the cut space.

Within this graph theoretical framework, the structure of $\varphi_{\mathcal{C}}(\mathcal{L})$ reads as follows. First, we notice that $\varphi_{\mathcal{C}}(\mathcal{L})$ is bipartite, that is, there is a bipartition of its vertices, so that edges interconnect vertices of different classes only. This bipartition is given by the set of vertices which belong to the spanning tree $\mathcal{L}'$ as edges of $\mathcal{L}$, and its complement. Therefore we may draw $\varphi_{\mathcal{C}}(\mathcal{L})$ schematically as depicted in Fig. 6. It is now straightforward to see that the neighbourhood $N_e(\mathcal{N}_e)$ of vertices $e \in \mathcal{E}(\mathcal{L}) \setminus \mathcal{E}(\mathcal{L}^e)$ $(e' \in \mathcal{E}(\mathcal{L}^e))$ on $\varphi_{\mathcal{C}}(\mathcal{L})$ joined with $e$ $(e')$ equals the fundamental cycle $\mathcal{C}_e$ $(\mathcal{D}_{e'})$ on $\mathcal{L}$ with respect to $\mathcal{L}'$. We conclude that $\varphi_{\mathcal{C}}(\mathcal{L})$ encodes the whole cycle space $\mathcal{E}(\mathcal{L})$, and the cut space $\mathcal{D}(\mathcal{L})$ as well. In Fig. 6 the highlighted subgraphs represent a fundamental cycle $\mathcal{C}_e$ and a fundamental cut $\mathcal{D}_{e'}$ indexed by $e$ and $e'$, respectively. That both, $\mathcal{E}(\mathcal{L})$ and $\mathcal{D}(\mathcal{L})$, are encoded in the same graph is not a coincidence but rather a necessity since it is well known that there is a duality between fundamental cycles and fundamental cuts $[54]$: $\varphi_{\mathcal{C}}(\mathcal{L})$ is a geometrical illustration of this duality. Since entanglement within the framework of graph states is characterised by edges between the qubits/vertices, the LC-equivalence of $|\varphi_{\mathcal{C}}(\mathcal{L})\rangle$ to a given surface code ground state on $\mathcal{L}$ illustrates that the entanglement pattern of a surface code is basically determined by its cycle and cut space structure.

D. Proof of Theorem 1

In this subsection the proof of Theorem 1 is presented in detail. Readers primarily interested in the minimal non-local toric code settings may skip this rather technical part and proceed with Section III.

Proof of Theorem 1. First note that the basis states $|v\rangle$ of the protected space are LC-equivalent since the topological quantum numbers $v_i$ are controlled by loop operators $X_k \in \mathcal{C}_k^{(S)}$. Therefore, from the viewpoint of LU/LC-classification, w.l.o.g. assume $v = (1, \ldots, 1)$. It follows that for each cycle in $\mathcal{L}$ there is a z-type loop operator in the stabilizer $S[\mathcal{X}]$ (Fig. 7 (A)). Let $\mathcal{L}'$ be an arbitrary spanning tree and $\mathcal{E}$ the set of edges such that $\mathcal{L}' = \mathcal{L} - \mathcal{E}$ (Fig. 7 (B)). Apply Hadamard transformations $H_e \in \mathcal{C}_N^{(S)}$ to the qubits identified with edges $e \in \mathcal{E}$. Formally this transformation reads $|v\rangle \mapsto H^\mathcal{E} |v\rangle$ where $H^\mathcal{E} = \otimes_{e \in \mathcal{E}} H_e$. We now argue that $H^\mathcal{E} |v\rangle = |\varphi_{\mathcal{C}}(\mathcal{L})\rangle$ by showing that the new stabilizer $S' := H^\mathcal{E} S[\mathcal{X}] (H^\mathcal{E})^{-1}$ is the graph state stabilizer described by $\varphi_{\mathcal{C}}(\mathcal{L})$.

In the stabilizer picture, a Hadamard transformation is represented by the substitution $\sigma^+ \leftrightarrow \sigma^-$. Consider one of these transformed qubits (8 in Fig. 7 (C)) and its edge $e = \{p, q\} \in \mathcal{E}(\mathcal{L}) (p, q \in \mathcal{V}(\mathcal{L}))$. The adjacent vertices $p$ and $q$ belong to both $\mathcal{L}$ and $\mathcal{L}'$. Since $\mathcal{L}'$ is a tree, there exists one and only one path $\mathcal{C}_{pq}$ on $\mathcal{L}'$ connecting $p$ and $q$ (----- in Fig. 7 (C)). Let $\sigma_{pq}^+ \otimes Z (C_{pq})$ be a combination of plaquette operators in $S[\mathcal{X}]$ (note that $e + \mathcal{C}_{pq}$ is a cycle in $\mathcal{L}$). This stabilizer operator is transformed under the Hadamard transformation as follows

$$K_G^{(e)} = H_e \sigma_{pq}^+ \otimes Z (C_{pq}) \quad H_e = \sigma_{pq}^+ \otimes \otimes_{i \in C_{pq}} \sigma_i^+ .$$

Obviously this is a graph state generator of $S[\varphi_{\mathcal{C}}(\mathcal{L})]$ since in $\varphi_{\mathcal{C}}(\mathcal{L})$ vertex $e$ is connected to all edges in $\mathcal{L}$ that belong to $\mathcal{C}_{pq}$ (Fig. 7 (D)). This argument holds for
any edge in $\mathcal{E}$. Therefore we find that the transformed stabilizer $\mathcal{S}'$ contains at least the graph state generators of $\varphi_{\mathcal{L}}(\mathcal{L})$ for vertices in $\mathcal{E}$. It remains to show that the graph state generators for vertices $e' \in E(\mathcal{L}')$ can be found in $\mathcal{S}'$ as well.

To this end consider an arbitrary qubit $e' \in E(\mathcal{L}')$ in the spanning tree ($\otimes$ in Fig. 7 (E)). The corresponding edge of the spanning tree cuts the latter into two parts. This construction yields a bipartition of the vertex set $V(\mathcal{L}') = V'_r \cup V'_b$ (red $\triangle$ and blue $\square$ in Fig. 7 (E)). Thus there are two types of edges in $\mathcal{E}$: The first type connects vertices of the same partition and the second type interconnects both sets $V'_b$ and $V'_r$. By definition of $\varphi_{\mathcal{L}'}(\mathcal{L})$ a vertex in $\mathcal{E}$ is connected to $e'$ iff it is a type-two vertex.

We claim that the following operator in $\mathcal{S}'$ is the graph state generator for $e'$

$$K^{(e')}_{\mathcal{G}} = \prod_{s \in V'_r} A'_s \text{ where } A'_s = H^{E} A_s (H^{E})^\dagger. \quad (6)$$

To see this, note that $\prod_{s \in V'_r} A'_s$ may affect a qubit $f$ in three different ways:

First, if $f$ belongs to an edge with adjacent vertices of the same partition, nothing happens since $\sigma^z \sigma^z = 1 = \sigma^+ \sigma^-$. The second case occurs if the adjacent vertices belong to different subsets and $f \neq e'$ ($\otimes$ in Fig. 7 (E)). Since this case can only apply if $f \in \mathcal{E}$, a $\sigma^z$ operation is performed on this qubit. The last case resembles the previous one, except for the fact that $f = e'$. Thus $\prod_{s \in V'_r} A'_s$ acts as $\sigma^x$ on $f$.

To sum up: we found that $K^{(e')}_{\mathcal{G}}$ acts as $\sigma^x$ on $e'$, as $\sigma^+$ on all vertices $f$ such that $\{e', f\} \in E(\varphi_{\mathcal{L}'}(\mathcal{L}))$, and trivially otherwise. Therefore we showed that $\mathcal{S} [\varphi_{\mathcal{L}'}(\mathcal{L})] \leq \mathcal{S}'$. To show $\mathcal{S} [\varphi_{\mathcal{L}'}(\mathcal{L})] = \mathcal{S}'$ we note that $\text{rank} \mathcal{S} [\varphi_{\mathcal{L}'}(\mathcal{L})] = N = \text{rank} \mathcal{S} [\mathcal{X}] = \text{rank} \mathcal{S}'$.

### III. MINIMAL NON-LOCAL TORIC CODE INSTANCES

**Setting the scene.** In this section, we present minimal instances for the toric code on square and triangular lattices which are not LC-equivalent to any local graph state. We already provided a brief explanation and motivation for the notion of “local” in the introductory paragraphs. At this point, we shall give a more formal definition and relate the idea of locality to the toric code hamiltonian.

The latter is usually written as

$$H_{\text{tcm}} = -J_A \sum_{s \in V(\mathcal{L})} A_s - J_B \sum_{p \in E(\mathcal{L})} B_p \quad (7)$$

where $J_A$ and $J_B$ define the excitation energies for electric and magnetic charges, respectively. Obviously the aforementioned protected space $\mathcal{P} \mathcal{S}$ is also the ground state space of the above hamiltonian and the orthogonal states $|\psi\rangle$ constitute a basis of the latter. In the preliminaries the toric code was described as a quantum code within the stabilizer framework. However, considering the hamiltonian (7), the toric code gains significance as an actual physical system that may be implemented directly or by means of a quantum simulator. Crucial for the physical relevance of $H_{\text{tcm}}$ is its quasi-locality, that is, only quasi-local interactions – represented by star and plaquette operators – are required. Therefore, from a physical point of view, the star and plaquette operators are distinguished by their quasi-locality, whereas from a mathematical point of view all generating sets of $\mathcal{S}$ are equivalent.

Consequently, the set of star and plaquette operators gives rise to a physically motivated adjacency relation on the set of qubits – namely, two qubits $p$ and $q$ are vicinal if both belong to the support of at least one star or plaquette operator, write $p \sqsubset q$. On the other hand, the relevance of graph states in experimental physics is due to their scalability which in turn relies on their successive implementation by means of 2-qubit operations (see Eq. (2)). Therefore it seems appropriate to call two qubits vicinal in a graph state if they are connected by...
an edge, write \( p_{\text{lg}} q \). The introduced symmetric relations \( i \) encode physically motivated adjacencies with respect to a particular class of multipartite entangled states.

After introducing the notion of vicinity, we may now ask the following question: Given the ground state \( |v\) of a toric code, is there an LU-equivalent graph state \(|\text{gs} G_{\text{loc}}\rangle\), such that for all pairs of qubits \( p_{\text{lg}} q \Rightarrow p_{\text{lc}} q \) holds. Provided such a graph state exists, we may apply those implementation techniques for graph states which gave rise to \( \text{gs} \) (within the setup defined by the toric code) in order to generate its ground states. Conversely, if there is no such state, the considered toric code setup yields ground states which are fundamentally different from its equivalent graph states, that is, the toric code ground states cannot be implemented by means of graph states without breaking the spatial structure of the toric code setup. At this point, the quasi-local property of \( H_{\text{tc}} \) becomes relevant. Thus we call toric code setups that exhibit at least one graph state \(|\text{gs} G_{\text{loc}} \rangle\) local and \( G_{\text{loc}} \) (\(|\text{gs} G_{\text{loc}} \rangle\)) itself a local graph (state) with respect to the considered toric code setup. Hence a toric code is local if one can draw an LU-equivalent graph state without “long-range” edges, that is, connecting qubits which cannot interact via star or plaquette operators in the toric code regime. E.g. see Fig. 2 for non-local and Fig. 5 for local instances of toric code systems.

Small toric code systems tend to be local, e.g. the single toric code plaquette in Fig. 1 (A) is represented by a local star graph. However, this property is likely to break down for more complex setups. This motivates our search for the smallest non-local toric code systems. In doing so, we restrained the analysis to small systems on square and triangular lattices, since experimental implementations are likely to use such tesselations.

A. Methods

Our approach is as follows. Due to the restriction to a particular tesselation (let us consider the square lattice exemplarily) we are looking for connected subsets of squares on the square lattice. To this end we may consider only setups with edge-connected squares since vertex-connected structures yield separable ground states or (if there is a loop through the vertices) degenerate ground states; the latter requiring a reduction to a non-degenerate setup by “filling” the loop, anyway [55]. Such structures are known in combinatorics as polyominos (on the square lattice) and polyiamonds (on the triangular lattice). The polyominos for up to \( n = 5 \) squares are depicted in Fig. 8. Application of Theorem 1 yields local graph states for all depicted setups devoid of “inner” squares – provided a suitable spanning tree is used for the construction. Actually, by means of Theorem 1 this can be worked out mentally. Subsequently there is a smallest setup left, such that the geometric transformation yields only non-local graphs. For the square lattice this would be the \( + \)-shaped pentomino \((n = 5)\) with dark coloured inner square, as depicted in Fig. 8.

However, the class of geometrically constructed graph states is generally a proper subset of the whole LC-class containing all LC-equivalent graph states [56]. To fix this issue, we computed the whole LC-class and searched for local representatives. To this end, an algorithm described in [20] and originally derived in [39, 40] was employed, according to which two graphs \( G \) and \( G' \) with adjacency matrices \( \Gamma \) and \( \Gamma' \) are equivalent under local complementations iff there exist diagonal matrices \( \textbf{A, B, C, D} \in \mathbb{F}_{2}^{N \times N} \), such that the non-linear condition

\[
(\textbf{TB + D}) \Gamma' + (\textbf{TAC + C}) = 0
\]

are satisfied over the Galois field \( \mathbb{F}_{2} \) (\( \textbf{E} \) denotes the identity matrix). This condition was implemented using Mathematica and facilitated the generation of the whole LC class \([G]_{\text{LC}} \) for a given graph \( G \). However, this algorithm is inefficient in the sense that computation time increases exponentially with the number of vertices. To our knowledge, no efficient (i.e. polynomial) algorithm has yet been found. Unfortunately, the exponential scaling behaviour renders the LC-class generation for the \( + \)-shaped setup (i.e. \( N = 16 \) vertices/qubits) unfeasible on a regular personal computer, whereas the triangular tetramond depicted in Fig. 2 (B) (i.e. \( N = 9 \) vertices/qubits) requires only few minutes to generate the whole LC-class and seek for local graphs. To deal with this (rather technical) problem, we conceived a method to deduce the non-locality of certain toric code setups from smaller ones. The details concerning this method are technical and purely graph theoretic. We therefore omit them at this point and refer the interested reader to Appendix A.
B. Results

As described above, the smallest setup on the square lattice without geometrically producible local graph state is the +−shaped pentomino (N = 16), as depicted in Fig. 2 (A). The aforementioned reduction method yields the smaller (N = 8) setup depicted in Fig. 2 (C). We proved the non-locality of the latter by generating the whole LC-class and probing for local graphs. Since the result was negative, we conclude that the system depicted in Fig. 2 (C) as well as the 5-plaquette setup in Fig. 2 (A) are non-local. Therefore, on the square lattice, the +−shaped pentomino with N = 16 qubits is the smallest setup without (LC-)equivalent local graph state.

On the triangular lattice the triangle shaped tetramond in Fig. 2 (B) is the smallest one yielding only non-local, geometrically constructed graph states. Furthermore, the generation of all LC-equivalent graphs yields only non-local ones. Thus, on the triangular lattice, the triangle-shaped tetramond with N = 9 qubits is the smallest setup without (LC-)equivalent local graph state.

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More concrete: Provided an embedding of a graph into a surface, then “deleting” the graph decomposes the surface into several pieces (the faces of this embedding). If each of these pieces can be deformed continuously into a disc, the embedding is called a 2-cell embedding. For instance, the common square-shaped plaquettes of a toric code system are homeomorphic to open discs; therefore the toric code is based on a 2-cell embedding of the square lattice into the 2-torus.

If $G$ is a graph on $n$ vertices, the line graph $L(G)$ contains $\binom{n}{2}$ edges. This is equivalent to the choice $\binom{n^2}{2}$ of all local complementations on vertices $a, b$.

This can be seen easily since any non-trivial LC-class $G_{\text{LC}}$ contains graphs with cycles. This is a bipartite graph by construction, and therefore features only cycles with an even number of edges.

That is, it is easy to see that all changes regarding edges between vertices in $V \setminus \{a\}$ occur in the same way, whether there is an edge to $a$ or not. Therefore the only differences between $\tau G_{ab}$ and $\tau (G_{ab} - a)$ affect edges from vertices in $V \setminus \{a\}$ to $a$ itself. Since these edges are deleted, $(\tau G_{ab} - a)$, there is no difference left and it follows $\tau G_{ab} - a = \tau (G_{ab} - a)$.

To prove the following statement.

Lemma 2 (Deletion of leaves). Let $G_{ab} \in \mathcal{G}(V)$ be a simple graph with at least one leaf, $a$ denotes the outer and $b$ the inner vertex. Consider the subgroup

$$\mathcal{T}_a := \{\tau_a \in \mathcal{G}(V) | E(V \setminus \{a\}) \}$$

of all local complementations on vertices $V \setminus \{a\}$. Then it holds

$$\forall \tau \in \mathcal{T}_a : \tau G_{ab} - a = \tau (G_{ab} - a) \tag{A3}$$

In words: Deleting the leaf $(a$ and its adjacent edge) from $G_{ab}$ and transforming this graph by a sequence of local complementations $\tau$ acting on vertices $V \setminus \{a\}$ yields the same graph as if these operations were performed on $G_{ab}$ (without deleting the leaf) and deleting $a$ with all its adjacent edges (which may be more than one) subsequently.

Proof. It is easy to see that all changes regarding edges between vertices in $V \setminus \{a\}$ occur in the same way, whether there is an edge to $a$ or not. Therefore the only differences between $\tau G_{ab}$ and $\tau (G_{ab} - a)$ affect edges from vertices in $V \setminus \{a\}$ to $a$ itself. Since these edges are deleted, $\tau G_{ab} - a$, there is no difference left and it follows $\tau G_{ab} - a = \tau (G_{ab} - a)$.

Lemma 2 allows statements about the LC-class $[G_{ab} - a]_{\text{LC}}$. More precisely: All graphs in $[G_{ab}]_{\text{LC}}$ that are reachable from $G_{ab}$ by concatenations of local complementations $\tau$ yield graphs in $[G_{ab} - a]_{\text{LC}}$ by deleting $a$ and its adjacent edges. If one can show that all graphs in $[G_{ab}]_{\text{LC}}$ may be linked to each other by operations in such subgroups, this yields a procedure to compute LC-classes of special subgraphs.

This idea leads to the following proposition for simple graphs with leaves:

Proposition 1 ($\varepsilon$-symmetric LC-classes). Let $G_{ab}$ be a simple graph with at least one leaf $a$ and $b$ denote the outer and inner vertex, respectively.

Then for all $H \in [G_{ab}]_{\text{LC}}$ there exists $H'_a \in [G_{ab} - a]_{\text{LC}}$ or $H'_b \in [G_{ba} - b]_{\text{LC}}$ such that $H = a = H'_a$ or $H - b = H'_b$, respectively.
In other words: If an LC-class features a leaf-graph representative with outer and inner vertex $a$ and $b$, then deleting one of the latter (in some cases it does not matter which one is deleted) yields a graph in $[G_{ab} - a]_{LC}$ or $[G_{ba} - b]_{LC}$. Consequently, any graph in $[G_{ab}]_{LC} = [G_{ba}]_{LC}$ features at least one subgraph in $[G_{ab} - a]_{LC}$ or $[G_{ba} - b]_{LC}$.

**Proof.** Let $G_{ab}$ be a representative with outer and inner vertex $a$ and $b$. Then one can partition the LC-class $[G_{ab}]_{LC}$ in the following way:

- All graphs with a leaf belong to $A$ iff $a$ is the outer and $b$ the inner vertex. Therefore $G_{ab} \in A$.
- All graphs with a leaf belong to $B$ iff $b$ is the outer and $a$ the inner vertex. Therefore $G_{ba} \in B$.
- All graphs $H$ where $\{a, b\} \in E(H)$ and for all $v \in \V(H) \backslash \{a, b\}$ it holds
  \[\{a, v\} \in E(H) \Leftrightarrow \{b, v\} \in E(H)\]
  belong to $C$. Due to the second statement, these graphs are totally symmetric with respect to $a$ and $b$. $\varepsilon_{ab}H = H$, that is.
- All graphs $H$ where $\{a, b\} \notin E(H)$ and for all $v \in \V(H) \backslash \{a, b\}$ it holds
  \[\{a, v\} \in E(H) \Leftrightarrow \{b, v\} \in E(H)\]
  belong to $D$. These graphs are totally symmetric; the only difference to type-$C$ graphs is the missing edge between $a$ and $b$.

These four types of graphs are depicted in Fig. 9. Note that the number of edges between $a$ and $b$ and the vertices $\V(G) \backslash \{a, b\}$ (denoted by light grey discs) is not constant within one class in general. That the described classification of graphs is a partition, i.e. $[G_{ab}]_{LC} = A \cup B \cup C \cup D$ has to be shown. To this end, we introduce a classification of local complementations $\tau_a \in T$ which depends on the graph they operate on. First, we call $\tau_a$ and $\tau_b$ an $a$- and $b$-operation, respectively. Secondly, local complementations $\tau_v$ (acting on $G$) where $v \in \V(G) \backslash \{a, b\}$ and $\{a, v\}, \{b, v\} \notin \E(G)$ are called $\gamma$-operations and LC-operations $\tau_v$ (acting on $G$) where $v \in \V(G) \backslash \{a, b\}$ and $\{a, v\} \in \E(G)$ or $\{b, v\} \in \E(G)$ are called $\xi$-operations. Obviously every local complementation $\tau_v$, $v \in \V(G)$, belongs (depending on the graph it is applied to) to one of these four classes. Let us examine how these four classes of local complementations and the four classes of graphs introduced above collude:

- First, note that $\gamma$-operations cannot alter the class a graph belongs to $(A, B, C$ or $D)$ since all edges used for the classification are unaffected by $\gamma$-operations. In Fig. 9 this is denoted by magenta arrows pointing from each set into itself.
- Clearly, $a$-operations act trivially on $A$. Applied to an element of $C$, it deletes all edges between $b$ and $\V(G) \backslash \{a, b\}$ and yields a graph that belongs to $B$. Since $\tau_a^2 = 1$, this is true for the opposite direction as well. Finally, graphs that belong to $D$ cannot leave this set since no edge between $a$ and $b$ is created. These connections are denoted by blue arrows in Fig. 9.
- Due to the symmetry with respect to $a$ and $b$, all statements for $\tau_a$ hold for $\tau_b$ (i.e. $b$-operations) as well. These relations are denoted by red arrows in Fig. 9. Note that $\tau_a$ equals $\tau_b$ if they are applied to elements in $D$.
- $\xi$-operations act on vertices denoted by dark grey circles in Fig. 9 (vertices that are connected to $a$ or $b$ or both of them). Obviously such operations cannot manipulate the leaf. Therefore $\xi$-operations cannot leave the sets $A$ and $B$ (but they may alter the number of edges connecting $a$ or $b$ to vertices in $\V(G) \backslash \{a, b\}$). Acting on elements in $C$, a $\xi$-operation deletes the edge between $a$ and $b$, hence converting type-$C$ graphs to type-$D$ graphs and vice versa. These relations are denoted by green arrows in Fig. 9.

Consider the following situation: Starting from $G_{ab} \in A$ one tries to generate as much elements in $[G_{ab}]_{LC}$ as possible by using local complementations in $\tau_a$. Therefore one is allowed to use $b$-, $\gamma$- and $\xi$-operations to generate graphs. In terms of Fig. 9 the “path” cannot use the blue arrows. Note that the only $a$-operation that cannot be omitted connects $C$ and $B$ (in $A$ it acts trivially and in $D$ it may be replaced by a $b$-operation). We call graphs in this orbit $T_a G_{ab}$ $a$-accessible. In conclusion it holds $T_a G_{ab} \subseteq [G_{ab}]_{LC} \setminus B$. Analogue arguments yield $T_b G_{ba} \subseteq [G_{ba}]_{LC} \setminus A$ for $b$-accessible graphs starting from $G_{ba} \in B$.

We are now going to show that the more stricter relations $T_a G_{ab} = [G_{ab}]_{LC} \setminus B$ and $T_b G_{ba} = [G_{ba}]_{LC} \setminus A$ hold. To this end, consider an arbitrary “path” jumping from class to class, w.l.o.g. starting at $G_{ab}$ and ending at $H \in [G_{ab}]_{LC} \setminus B$. Formally described by
\[ H = \prod_t \tau_v G_{ab} \text{ where } \tau_v \in T \]
as depicted in Fig. 10 (A). In general there will be some $a$-operations in this chain. Since neither $G_{ab}$ nor $H$ are in $B$ there is an even number of them (in Fig. 10 (A) there are two, denoted by blue jumps). We are restricted to non-$a$-operations in $T_a$, though. We can adhere to this restriction by “swapping” the $a$-jumps and the corresponding $\gamma$- and $\xi$-operations in $B$ to $b$-jumps and the same $\gamma$- and $\xi$-operations in $A$. This is shown in Fig. 10 (B).
III Let in order to show that the surface code system can enable the very graphs describing the same one as formerly obtained by the second application of, due to symmetry, each $H$ is permuted. However, since the second application of $C$ before jumping back to $A$ showed that $\tau \text{LC} \gamma = \gamma \text{LC} \tau \gamma$. Choose $\tau G_{ba} = [G_{ba} \text{LC}] \backslash A$ and $\tau G_{ba} = [G_{ba} \text{LC}] \backslash B$.

To conclude the proof, recall Lemma 2. Consider $H \in [G_{ab} \text{LC}]$. Assume without loss of generality that $H \notin B$ (the following argument holds for $H \notin A$ analogously). Then we showed that $H \in \mathcal{T}_a G_{ab}$ where $G_{ab}$ is a leaf-graph with outer vertex $a$. Choose $\tau \in \mathcal{T}_a$ such that $H = \tau G_{ab}$. According to Lemma 2 it follows that $H - a = \tau G_{ab} - a = \tau (G_{ab} - a)$. Set $H'_a := \tau (G_{ab} - a) \in [G_{ab} - a]_{\text{LC}}$ and it follows $H - a = H'_a \in [G_{ab} - a]_{\text{LC}}$. Which was to be proven.

2. Application to the locality problem

In order to apply the graph theoretic knowledge about $\varepsilon$-symmetric LC-classes introduced above, we have to define in graph theoretic terms the notion of “local surface codes” and “local graph states”.

Definition 2 (Adjacency relation, Local graph). Let $V$ be an arbitrary finite set. An adjacency relation $\Lambda$ on $V$ is an irreflexive and symmetric relation, i.e. there is no $v \in V$ such that $v \sim v$ and for each pair $v, w \in V$ it holds $v \sim w \Rightarrow w \sim v$. An adjacency relation is represented by an adjacency matrix $H$ where $H_{v,w} = 1$ if $v \sim w$ and $H_{v,w} = 0$ otherwise. Therefore $\Lambda$ defines a simple graph $G_\Lambda$ with adjacency matrix $H$. A simple graph $G_{\text{loc}}$ is called local with respect to $\Lambda$ if it is a subgraph of $G_\Lambda$.

Given a surface code $\mathcal{L}$ (more precisely: a 2-cell embedding $X_{\mathcal{L},\Sigma}$ of $\mathcal{L}$ in $\Sigma$), the stabilizer generators $A_\gamma$ and $B_\varepsilon$ induce an adjacency relation $\Lambda$ on the set of edges (or qubits) $E(\mathcal{L})$ as follows. Two qubits $e, f \in E(\mathcal{L})$ are adjacent, $e \sim f$, if and only if there is a common face $p \in \mathcal{P}(\mathcal{L})$ or a common vertex $s \in \mathcal{V}(\mathcal{L})$ such that $e, f \in p$ or $e, f \in s$, respectively. Note that by this definition of $\Lambda$, local graphs $G_{\text{loc}}$ in the sense of Def. 2 are the very graphs describing local graph states $|G_{\text{loc}}\rangle$ according to the definition given in Section III.

Since we want to reduce surface code setups to smaller ones by removing edges/qubits, a method to compare two setups with respect to their adjacency relations is needed. Therefore we introduce the notion of strictness.

Definition 3 (Strictness). Let $\mathcal{L}_1$ and $\mathcal{L}_2$ be two surface codes with edges (qubits) $E(\mathcal{L}_1) \subseteq E(\mathcal{L}_2)$. Furthermore denote by $\Lambda_1$ and $\Lambda_2$ the adjacency relations as defined above. They define simple graphs $G_{\text{loc}}$ and $G_{\text{loc}}$, with vertices $E(\mathcal{L}_1)$ and $E(\mathcal{L}_2)$, respectively. $\Lambda_1$ is called stricter than $\Lambda_2$ (write $\Lambda_1 \succ \Lambda_2$) if $G_{\text{loc}}[E(\mathcal{L}_2)] \triangleleft G_{\text{loc}}[E(\mathcal{L}_1)]$.

Here $G_{\text{loc}}[E(\mathcal{L}_2)]$ denotes the subgraph of $G_{\text{loc}}$ on the vertices $E(\mathcal{L}_2) \subseteq E(\mathcal{L}_1)$. Definition 3 and Proposition 1 enable us to state and prove the following theorem, which is the main result of this section:

Theorem 2 (Surface code reduction). Let $\mathcal{L}$, $\mathcal{L}_a$ and $\mathcal{L}_b$ be surface code systems with the property that there exist qubits $a, b \in E(\mathcal{L})$ such that $E(\mathcal{L}_a) = E(\mathcal{L}) \setminus \{a\}$ and $E(\mathcal{L}_b) = E(\mathcal{L}) \setminus \{b\}$. $\Lambda$, $\Lambda_a$ and $\Lambda_b$ denote the induced adjacency relations. In addition we require

1. system $\mathcal{L}$ to be stricter than the other systems, i.e. $\Lambda \succ \Lambda_a$ and $\Lambda \succ \Lambda_b$.
2. that there is a leaf-graph $G_{ab}$, describing an LC-equivalent graph state of system $\mathcal{L}$, such that $G_{ab} - a$ and $G_{ba} - b$ describe LC-equivalent graph states of $\mathcal{L}_a$ and $\mathcal{L}_b$, respectively.
3. the surface code systems $\mathcal{L}_a$ and $\mathcal{L}_b$ to be nonlocal. Then the surface code system $\mathcal{L}$ is nonlocal.

Proof. In order to show that the surface code system defined by $\mathcal{L}$ is non-local, one has to find at least one non-local edge for each element in $[G_{ab}]_{\text{LC}}$ with respect to the adjacency relation $\Lambda$. Let $H \in [G_{ab}]_{\text{LC}}$ be an arbitrary graph representing an LC-equivalent graph state. According to Proposition 1, each element in $[G_{ab}]_{\text{LC}}$ has at least one subgraph in $[G_{ab} - a]_{\text{LC}}$ or $[G_{ba} - b]_{\text{LC}}$. W.l.o.g. let $H'_a \in [G_{ab} - a]_{\text{LC}}$ be such a subgraph, i.e. $H'_a \subseteq H$. Since $H'_a$ describes an LC-equivalent graph state of system $\mathcal{L}_a$ (see 2.) there is at least one edge $e \in \{v_1, v_2\} \in E(H'_a) \subseteq E(H)$, such that the vertices are not adjacent with respect to $\Lambda_a$, i.e. $e \notin E(G_{\Lambda_a})$ (see 3.). Since system $\mathcal{L}$ is stricter than $\mathcal{L}_a$ (see 1.) it follows $G_{\Lambda_a}[E(\mathcal{L}_a)] \triangleleft G_{\Lambda_a}$ and consequently

Figure 10. (Color online) An arbitrary path from $G_{ab}$ (red dot) to $H$ (green dot) is depicted in (A). The “forbidden” (blue) paths using $A$-operations are swapped to $A$ (red jumps) using $B$-operations instead, see (B). Now the whole path from $G_{ab}$ to $H$ is $a$-accessible. A detailed description is given in the text.

It is easy to see that jumping to $A$ by $b$-operations instead and subsequently applying the same $\gamma$- and $\varepsilon$-operations yields a graph in $A$ that resembles the graph obtained in $B$ before jumping back to $C$ with the only difference that $a$ and $b$ are permuted. However, since the second application of $\tau_2$ yields a totally symmetric graph in $C$, this graph has to be the same one as formerly obtained by the second $\tau_2$ operation.

Consequently we found a procedure to reach any graph $H \in [G_{ba}]_{\text{LC}} \backslash B$ by local complementations in $T_a$ from $G_{ab}$ and, due to symmetry, each $H' \in [G_{ba}]_{\text{LC}} \backslash A$ by exclusive application of $T_b$ operations from $G_{ba}$ [58]. Thus we showed $T_a G_{ba} = [G_{ba}]_{\text{LC}} \backslash A$ and $T_b G_{ba} = [G_{ba}]_{\text{LC}} \backslash B$.
Figure 11. Example for the reduction procedure based upon Theorem 2. A: The +-shaped pentomino as presented in Fig. 2 (A). B-C: By reducing the upper square twice one obtains successively smaller systems. D-F: This procedure is repeatedly applied to the remaining squares and leads to the system depicted in Fig. 2 (C). The “removal” of each edge/qubit is justified by Theorem 2; thus the non-locality of the smallest setup (F) implies the non-locality of the whole chain of larger setups.

e \not\in E(G_{\Lambda_{a}}) \Rightarrow e \not\in E(G_{\Lambda} \mid E(L_{a}))$. Since $v_{1}, v_{2} \in E(L_{a})$ it follows furthermore $e \not\in E(G_{\Lambda})$. Thus we found an edge $e$ in $E(H)$ that represents a non-local interaction in the larger system $\mathcal{L}$. Since $H$ was chosen arbitrarily, this shows the non-locality of $\mathcal{L}$.

Using Theorem 2, the +-shaped pentomino (see Fig. 2 (A)) can be reduced to the 8-qubit setup depicted in Fig. 2 (C). The procedure applied in this particular case is depicted in Fig. 11. The peripheral edges are contracted successively until we end up with the 8-qubit setup in Fig. 11 (F). The contractions are justified by Theorem 2. Each step requires an LC-equivalent graph state with leaf-graph $G_{ab}$ as depicted in Fig. 2 (A), such that $G_{ab} - a$ describes an LC-equivalent graph state of the next (reduced) setup. One can perform these steps easily by hand (they are not very exciting!) and, as a consequence, verify the non-locality of the +-shaped pentomino using the non-locality of the 8-qubit setup.