I. FIBONACCI ANYONS

The degrees of freedom in our microscopic models are so-called Fibonacci anyons, one of the simplest types of non-abelian anyons \([1, 2]\). The Fibonacci theory has two distinct particles, the trivial state \(1\) and the Fibonacci anyon \(\tau\), which can be thought of as a generalization (or more precisely a ‘truncated version’) of an ‘angular momentum’ when viewing the Fibonacci theory as a certain deformation \([3]\) of \(SU(2)\). We will now make this notion more precise and illustrate it in detail. In analogy to the ordinary angular momentum coupling rules, we can write down a set of ‘fusion rules’ for the anyonic degrees of freedom which are analogs of the Clebsch-Gordon rules for coupling of ordinary angular momenta, the pentagon and hexagon relationsthen yield that the corresponding anyonic degrees of freedom which are analogs of the Clebsch-Gordon rules for coupling of ordinary angular momenta, 

\[
1 \times 1 = 1 \quad 1 \times \tau = \tau \times 1 \quad \tau \times \tau = 1 + \tau,
\]

where the last fusion rule reveals what is known as the non-abelian character of the Fibonacci anyon: Two Fibonacci anyons \(\tau\) can fuse to either the trivial particle or to another Fibonacci anyon. In more mathematical terms, these fusion rules can also be expressed by means of so-called fusion matrices \(N_j\) whose entries \((N_j)_{ij}\) equal to one if and only if the fusion of anyons of types \(j_1\) and \(j_2\) into \(j\) is possible. The fusion rules are related to the so-called ‘quantum dimensions’ \(d_j\) of the anyonic particles by

\[
N_j |d_j\rangle = d_j |d_j\rangle,
\]

where \(|d_j\rangle\) is the (‘Perron Frobenius’) eigenvector corresponding to the largest positive eigenvalue of the \(2 \times 2\)-matrix \(N_j\). [The sense in which these numbers are ‘dimensions’ will become apparent in section II A 1 below.] For the particles in the Fibonacci theory the quantum dimensions are \(d_1 = 1\) and \(d_\tau = \varphi \equiv (1 + \sqrt{5})/2\) and the total quantum dimension of the theory is then given by \(D = (\sum_j d_j^2)^{1/2} = \sqrt{1 + \varphi^2}\).

To define our Hamiltonian, some additional ingredients of the theory of anyons are required. In analogy to the \(ij\)-symbols for ordinary \(SU(2)\) spins, there exists a basis transformation \(F\) that relates the different ways three anyons can fuse to a fourth anyon, depicted as

\[
\begin{array}{ccc}
  a & b & c \\
  \downarrow & \downarrow & \downarrow \\
  d & e & f
\end{array}
= \sum_f \left( F_{abc}^d \right)_f \begin{array}{ccc}
  a & b & c \\
  \downarrow & \downarrow & \downarrow \\
  d & e & f
\end{array}.
\]

The left hand side (l.h.s.) represents the quantum state that arises when anyon \(a\) first fuses with anyon \(b\) into an anyon of type \(e\), which, subsequently, fuses with anyon \(c\) into an anyon of type \(d\). Similarly, the right hand side (r.h.s.) denotes the quantum states that arises when anyon \(b\) first fuses with anyon \(c\) into anyon type \(f\) which, in turn, fuses with anyon \(a\) into anyon type \(d\). Whilst keeping all external labels, the types of the three anyons \((a, b, c)\) as well as the resulting anyon type \(d\) fixed, the states on the l.h.s and r.h.s. are fully specified by the labels \(e\) and \(f\), respectively. Eq. (3) says the so-specified states are linearly related to each other by the so-called \(F\)-matrix \([4]\) with matrix elements \(\left( F_{abc}^d \right)\).

In general, the \(F\)-matrix is uniquely defined (up to ‘gauge transformations’) by the fusion rules through a consistency relation called the pentagon equation \([6]\). Similarly, the braiding properties of anyons are given by the so-called \(R\)-matrix (which however is not needed here) that is uniquely determined by the hexagon equation \([6]\).

For the Fibonacci theory, it is straightforward to verify that in most cases there is only one term on the right-hand-side in Eq. (3), e.g. by choosing two or three out of the four anyons \(a, b, c, d\) to be \(\tau\)-anyons. For these cases the consistency with the pentagon and hexagon relations then yields that the corresponding \(F\)-matrix elements equal to 1. There is only one configuration that gives rise to \(F\)-matrix elements that are non-trivial: If all anyons are \(\tau\)-anyons, e.g. \(a = b = c = d = \tau\), both the 1- and the \(\tau\)-fusion channels appear, and the \(F\)-matrix takes the explicit form

\[
F_{\tau\tau\tau} = \begin{pmatrix}
  (F_{\tau\tau\tau})_{1} \quad (F_{\tau\tau\tau})_{2} \\
  (F_{\tau\tau\tau})_{1} \quad (F_{\tau\tau\tau})_{2}
\end{pmatrix} = \begin{pmatrix}
  \varphi^{-1} \quad \varphi^{-1/2} \\
  \varphi^{-1/2} \quad \varphi^{-1}
\end{pmatrix}.
\]

As a final ingredient to explicitly derive our Hamiltonian, we have to introduce the so-called modular \(S\)-matrix that relates the anyon “flux” of species \(b\) through an anyon loop of species \(a\) to the case without any loop by

\[
\begin{array}{c}
  a \\
  \begin{array}{c}
    \overline{b} \\
    \hline
  \end{array}
\end{array} = \frac{S_{ab}}{S_{\tau\tau}^b} \begin{array}{c}
  a \\
  \begin{array}{c}
    \overline{b} \\
    \hline
  \end{array}
\end{array}.
\]
For the case of Fibonacci anyons, the $S$-matrix takes the explicit form

$$S = \begin{pmatrix} S_1^1 & S_1^2 \\ S_2^1 & S_2^2 \end{pmatrix} = \frac{1}{D} \begin{pmatrix} 1 & \varphi \\ \varphi & -1 \end{pmatrix}. \tag{6}$$

There is an important relationship between the modular $S$-matrix and the matrix encoding the fusion rules, introduced in the paragraph above Eq. (2): the modular $S$-matrix diagonalizes the fusion rules, the ‘Verlinde Formula’,

$$S_b^b (N_a)^{c'}_c S_c^{c''} = \delta_c^b \frac{S_a}{S_b} \tag{7}$$

(repeated indices are summed) where $S_\dagger$ denotes the adjoint of the unitary matrix $S$. The eigenvalues of the matrix $(N_a)$ are thus $\frac{S_a}{S_b}$, and the largest (positive) eigenvalue, the quantum dimension $d_a$, can be seen to be

$$d_a = \frac{S_a}{S_1^1}. \tag{8}$$

Due to the unitarity of the modular $S$-matrix one immediately checks that the total quantum dimension equals

$$D = \frac{1}{S_1^1}. \tag{9}$$

II. THE LADDER MODEL

In this section we will discuss details of the “ladder model” in a one-dimensional geometry, whose Hamiltonian is given by Eq. (2) in the main part of the paper. We start by defining the Hamiltonian in detail, and then discuss the gapped topological phases, critical phases, and the exact solutions.

A. The Hamiltonian

1. Explicit expression

To establish a notation for the basis states we consider the skeleton lattice inside the high-genus ladder geometry as shown in Fig. 1. The basis states are given by all admissible labeling of the edges of the skeleton with 1 or $\tau$ particles, subject to the vertex constraints given by the fusion rules. The number of basis states, $B_L$, of the ladder with $L$ plaquettes and periodic boundary conditions is given by

$$B_L = \sum_{\{a_i, b_i, c_i\}} (N_{c_1})^{a_1}_{a_2} (N_{c_2})^{a_2}_{a_3} \cdots (N_{c_L})^{a_L}_{b_2} (N_{c_1})^{b_1}_{b_2} (N_{c_2})^{b_2}_{b_3} \cdots (N_{c_L})^{b_L}_{b_2} = \sum_{\{i_1, \ldots, i_L\}} [\text{Tr}(N_{i_1} N_{i_2} \cdots N_{i_L})]^2, \tag{10}$$

where $N_i$ are the fusion matrices of Fibonacci theory as introduced above. The largest eigenvalue of the matrix $N_i$ is the quantum dimension $d_i$. Thus, the leading behavior of the traces for large $L$ is,

$$B_L \sim \sum_{\{i_1, \ldots, i_L\}} (d_{i_1} d_{i_2} \cdots d_{i_L})^2 = \sum_{k=0}^L (d_1^2)^{L-k} (d_\tau^2)^k = (1 + \varphi^2)^L = D^2. \tag{11}$$

The Hilbert space thus grows asymptotically, for large $L$, as a power of the square of the total quantum dimension $D^2$.

The Hamiltonian (as given in Eq. (2) of the main part of the paper)

$$H = -J_{r} \sum_{\text{rungs} r} \delta_{\ell(r), 1} - J_{p} \sum_{\text{plaq} p} \delta_{\phi(p), 1}, \tag{12}$$

consists of two non-commuting terms, the rung term which is diagonal in the chosen basis, and the plaquette term which depends on the four edges of the plaquette $p$, and the four adjoining edges. By inserting an additional anyon loop of type $s$ into the center of the plaquette, we can project onto the flux through this additional loop (and hence the flux through the plaquette) by the following procedure (for a derivation see the following subsection)
The duality can be made exact by using "twisted" boundary conditions where the ends of the ladder are connected according to only difference is that the rungs connect two different cylinders, while the plaquettes connect the space (the "outside").

The additional $s$-loop is inserted by performing a sequence of $F$-transformations:

$$
\delta_{\phi(p)_1} \begin{array}{c|cc|c}
\alpha & \beta & \gamma \\
\delta & \delta & \gamma \\
\end{array} = \sum_{s=1, \tau} \frac{d_s}{D^2} \begin{array}{c|cc|c}
\alpha & \beta & \gamma \\
\delta & \delta & \gamma \\
\end{array}.
$$

Using the identity

$$
\delta_{m\delta'} \begin{array}{c|cc|c}
\alpha & \beta & \gamma \\
\delta & \delta & \gamma \\
\end{array} = \begin{array}{c|cc|c}
\alpha & \beta & \gamma \\
\delta & \delta & \gamma \\
\end{array},
$$

we obtain the final expression

$$
\delta_{\phi(p)_1} \begin{array}{c|cc|c}
\alpha & \beta & \gamma \\
\delta & \delta & \gamma \\
\end{array} = \sum_{s=1, \tau} \frac{d_s}{D^2} \sum_{\alpha', \beta', \gamma'} (F_{d\delta s})_{1}^{\alpha'} \begin{array}{c|cc|c}
\gamma' & \beta' & \gamma \\
\delta' & \delta' & \gamma \\
\end{array}.
$$

The ladder geometry has a local duality between the inside and outside: the inside of the rungs is dual to the plaquettes. The only difference is that the rungs connect two different cylinders, while the plaquettes connect the same space (the "outside"). The duality can be made exact by using "twisted" boundary conditions where the ends of the ladder are connected according to $a_1 = b_{L+1}$ and $b_1 = a_{L+1}$ (so that the ladder looks like a Moebius strip). Indeed, our exact diagonalization results confirm that the excitation spectra are identical under exchange of the couplings $J_x$ and $J_y$ for twisted boundary conditions. However, in the case of periodic boundary conditions ($a_1 = a_{L+1}$, $b_1 = b_{L+1}$), which we shall focus on in the following, this duality is only up to degeneracies.
2. Bigger (mathematical) picture

So far our discussion in this ‘Supplementary Material’ has been largely focused on detailed algebraic manipulations. In this subsection we wish to give a brief idea of the general bigger picture of topological field theories which underlies these detailed manipulations. At the same time we will provide a deeper understanding of the so-called ‘Levin-Wen model’ within this context.

In the main text we have given a physically motivated description of the Levin-Wen model in Figs. 1 and 2, leading to the Hamiltonian in Eq. (1) of the main text. Let us now give a more abstract description of it. The most general Levin-Wen Hamiltonian has two kinds of terms: the vertex type (not discussed so far as a term of the Hamiltonian) and plaquette type. Let us consider a surface $\Sigma$, and a trivalent graph $\Gamma$ (which we called ‘skeleton in the main text) embedded in that surface. (The sole role of the surface $\Sigma$, which in the leftmost picture of Fig. 1 of the main text is just a parallel plane sitting in between the two depicted sheets, is to give a well defined meaning to the notion of a ‘plaquette’; namely, all complimentary regions of $\Sigma\setminus\Gamma$, i.e. the complimentary regions of the graph $\Gamma$ within the surface $\Sigma$, are plaquettes.) We always enforce strictly the condition that three labels meeting at a vertex must satisfy the fusion rule. (This is another way of saying that we have set the coupling constant of the ‘vertex term’ in the most general Levin-Wen Hamiltonian to infinity.) As a result, we obtain a Hilbert space called $L(\Gamma, \Sigma)$ consisting of the Hilbert space spanned by all admissible labelings of the trivalent graph $\Gamma$: a labeling of $\Gamma$ is an assignment of a label in a label set $I$ to each edge of the graph, $I_C = \{1, \tau\}$ in the previous subsection I, and the labeling is admissible if the three labels around each vertex satisfy the fusion rules.

Now, there exists another vector space, which brings about the connection with the actual surfaces that were drawn in Figs. 1 and 2 of the main text. In particular, when $C$ denotes a so-called modular category (for a precise definition, which we do not need at the moment, see e.g. Ref. 7) which basically denotes a theory of ‘anyons’ and their corresponding ‘fusion rules’ such as the one described in the previous subsection I, then the vector space $L(\Gamma, \Sigma)$ is the same as the Hilbert space $V_C(S_T)$ (for a definition see e.g. Ref. 7) of an associated Topological Quantum Field Theory (TQFT) corresponding to the ‘modular category’ $C$: specifically let $N_T$ be the thickening of the graph $\Gamma$ to a handle-body (drawing a cylinder around each edge of the graph), and $S_T$ be the boundary surface of $N_T$, then $L(\Gamma, \Sigma) \cong V_C(S_T)$. In the language of TQFT, any ‘pants-decomposition’ of the surface $S_T$ is known to lead to a basis of $V_C(S_T)$, which corresponds to the vector space spanned all possible fusions of the labelings on $\Gamma$.

This interpretation of the Hilbert space $L(\Gamma, \Sigma)$ gives rise to a transparent derivation of the plaquette term, Eq. (16), in the Levin-Wen model. To derive this expression, we use the identification of $L(\Gamma, \Sigma)$ with $V_C(S_T)$. The $\omega$th row of the modular $S$-matrix of the modular category $C$ can be used to construct a projector $\omega_c$ that projects out the particle with a label $c$ through a plaquette. In other words total flux $e$ through a plaquette $p$ can be enforced by inserting $\omega_e$ into a plaquette $p$. The projector turns out to be of the form

$$\omega_c = \frac{1}{D} \sum_a S^a_c [a],$$

where $[a]$ denotes a loop labeled by $a$ as the one drawn in Eq. (5). In order to see that this performs the task let us insert a flux with label $b$ thought the loop $[a]$, resulting in the figure drawn on the l.h.s. of Eq. (5), which we denote in symbols by $[a][b]$. When we now perform the sum in Eq. (17) we obtain, upon making use of Eq. (5),

$$\omega_c(b) := \frac{\delta^b_c}{D} \sum_a S^a_c \cdot [a][b] = \frac{1}{D} \sum_a S^a_c S^b_{S_T} \frac{1}{S_T} [b] = \frac{1}{D} \frac{1}{S_T} \delta^b_c \cdot [c]$$

where we have used the unitarity (plus reality and symmetry) of the modular $S$-matrix, as well as Eq.s (8,9).

Therefore, the plaquette term $\delta_{\omega(p),1}$ is implemented by inserting the projector $\omega_1 = \sum_a \frac{S^a_{S_T}}{S_T} \cdot [a]$ into the plaquette $p$. Now the detailed steps leading to Eq. (16) are easy to understand: The insertion of $\omega_1$ into the plaquette is written explicitly in Eq. (13). In the subsequent equation, first an $F$-move is applied to the two lines connected by the dotted line, and subsequently four more $F$ moves counterclockwise around $p$ are implemented as drawn; finally removing the resulting bubble, we obtain the explicit form of the plaquette term written in Eq. (16).

The mathematical context for the Levin-Wen model is the Drinfeld center $Z(C)$ or quantum double of a unitary fusion category $C$. The label set $I_C$ for the Levin-Wen Hamiltonian is the isomorphism classes of simple objects of $C$. It is known that a unitary fusion category is always spherical. By a theorem of M. Müger [8], the Drinfeld center of any spherical category is always modular. It follows that the Drinfeld center of any unitary fusion category is always modular. Moreover, if the spherical category $C$ itself is modular, then $Z(C)$ is isomorphic to the direct product of the conjugate $C^*$ and $C$, where $C^*$ is obtained from $C$ by complex conjugating all data. Our main example is one of those special cases, where $C$ is the Fibonacci theory.
FIG. 2: Possible basis configurations in the presence of one plaquette excitation. The two ground states (without a flux through the plaquette) and three excited states (with a $\tau$-flux through the plaquette) are linear combinations of these basis states. Solid lines denote $\tau$-anyons while dashed lines symbolize the trivial particle 1.

The decomposition of $Z(C)$ hints directly at the appearance of Dynkin diagram $D_6$ at the critical point in one-dimensional geometry: indeed, the two phases connected by the critical point are the Fibonacci theory and the doubled Fibonacci theory with label sets $\{1, \tau\}$ and $\{(1, 1), (1, \tau), (\tau, 1), (\tau, \tau)\}$, respectively. Based on this, it is natural to expect that the two sets of fusion rules will fit together in a compatible way at the critical point, which is nicely illustrated by the structure of the $D_6$ Dynkin diagram in Fig. 11 which underlies the exact solution of this critical point (Section IID below).

B. Topological phases

We start the detailed discussion of the phase diagram with the two distinct gapped non-abelian topological phases: the ‘single torus’ phase where all plaquettes are closed at $\theta = 0$ and the ‘two tori’ phase with closed off rungs at $\theta = \pi/2$. A finite-size scaling analysis of the splitting of the ground state degeneracies and the energy gap shows that the phases extend over a wide range of parameter space as illustrated in the phase diagram (Fig. 6a of the main part of the paper). In this section we discuss the low-lying excited states in these phases, give their explicit wavefunctions at the exactly solvable points, and a perturbative expansion for their dispersion away from these points.

1. The ‘single torus’ phase at $-\pi/2 < \theta < \pi/4$

To describe the lowest excited states we consider the trivially solvable point $\theta = 0$ where $J_r = 0$. In the ground state there is no flux through any of the plaquettes, and all they can be closed, thus reducing the high-genus ladder to a single torus (see Fig. 5a in the main part of the paper). There are two degenerate ground-states configurations with either no flux or a $\tau$-flux through this torus.

Similarly, we can deduce the degree of degeneracy for the lowest excited state by considering the topology of this state. In the lowest excited state, one plaquette flux is present which yields the reduced topology (as compared to the high-genus ladder) and the associated skeleton shown in Fig. 5a of the main part of the paper. Closing all but one plaquette this skeleton allows for 5 different $1, \tau$ coverings, illustrated schematically in Fig. 2. In order to obtain the anyon-fluxes through the excited plaquette, a basis transformation (consisting of a $F$- and a $S$-transformation) of the reduced basis is performed which yields that there are three $\tau$-fluxes through each plaquette. Thus, the lowest excited state at $\theta = 0$ is $3L$-fold degenerate. Tuning away from $\theta = 0$ these $3L$ excitations delocalize and form a three-fold degenerate band.

2. The ‘two tori’ phase at $\pi/4 < \theta < \pi$

At the point $\theta = \pi/2$ (trivially solvable) the ground state has no $\tau$-anyons on the rungs of the ladder. The rungs can hence be cut which yields an effective topology of two separate tori. Of the four degenerate ground states three are symmetric and one is antisymmetric under $y$-reflection. The lowest excited state is a $\tau$-anyon flux through a single rung. The fusion rules then require a flux through both of the two tori, and this state is hence only $L$-fold degenerate. Tuning away from $\theta = \pi/2$, these states delocalize into a non-degenerate band.

3. Perturbation expansion for the quasiparticle bands

Over a broad range of parameters the quasi-particle excitations are well described (see Fig. 6a of the main part of the paper) by a second order perturbative expansion around $\theta = \pi/2$, with a dispersion given by

$$\Delta E(J_p, J_r, k_x) = J_r - \frac{2J_r}{D^2} \cos(k_x) - \frac{J_r^2}{D^4} \left[1 + 2 \cos(k_x)\right] - \frac{J_r^2}{2D^4} \cos(2k_x).$$

Due to duality, this result equally applies for coupling parameters $\theta$ close to $\theta = 0$, with $J_r$ and $J_p$ interchanged.
C. Gapless theories

In this section, we discuss the critical points \((\theta = \pi/4, 5\pi/4)\) and the extended critical phase in the ladder model. We first discuss the gapless theories in terms of numerical results and then present analytical arguments leading to an exact solution for the two critical points \((\theta = \pi/4, 5\pi/4)\).

1. Critical point at \(\theta = \pi/4\)

(numerical findings from exact diagonalization)

At equal positive values of the two coupling constants \((J_p = J_r, \theta = \pi/4)\), the system has a linear energy-momentum dispersion relation with the finite-size spacing between energy levels vanishing linearly in \(1/L\). This indicates that the two adjacent, gapped topological phases are separated by a continuous quantum phase transition and a critical point that is described by a 2D conformal field theory (CFT). To characterize this CFT, we rescale and match the finite-size energy spectra obtained numerically by exact diagonalization for systems with up to \(L = 36\) anyons to the form of the spectrum of a CFT,

\[ E = E_1 L + \frac{2\pi v}{L} \left( -\frac{c}{12} + h + \bar{h} \right), \]

(21)

where the velocity \(v\) is an overall scale factor, and \(c\) is the central charge of the CFT. The scaling dimensions \(h + \bar{h}\) take the form \(h = h^0 + n, \bar{h} = \bar{h}^0 + \bar{n}\), with \(n\) and \(\bar{n}\) non-negative integers, and \(h^0\) and \(\bar{h}^0\) are the holomorphic and antiholomorphic conformal weights of primary fields in a given CFT with central charge \(c\). The momenta (in units \(2\pi/L\)) are such that \(k_x = h - \bar{h}\) or \(k_x = h - \bar{h} + L/2\). Using this procedure, we find that for the critical point at \(\theta = \pi/4\) the rescaled energy spectrum matches the assignments (21) of part of the Kac-Table of the \(m = 9\) unitary Virasoro minimal CFT of central charge \(c = 14/15\), as shown in Fig. 3. In Fig. 4 we list all relevant primary fields of this CFT which appear and their corresponding scaling dimensions. It turns out that only the Kac-Table primary fields \(\phi_{r,s}\) with \(s = \text{odd}\) appear, and those with \(s = 5\) have multiplicity two (the associated states on the ladder being in the bonding/antibonding sectors of ‘transverse momenta’ \(k_y = 0, \pi\)), all others having multiplicity one. These are precisely those Kac-table primary fields which occur in the so-called \((D,A)\)-modular invariant \([9]\) of the \(m = 9\) Virasoro minimal CFT of central charge \(c = 14/15\).

To illustrate how the ground-state degeneracy changes at this critical point from a two-fold degeneracy for the ‘single cylinder’ limit \((J_r = 0)\) to a four-fold degeneracy for the ‘two cylinders’-limit \((J_p = 0)\), we can follow the evolution of eigenenergies in

FIG. 3: Exact diagonalization: Energy spectrum at the critical point \((\theta = \pi/4)\) for a ladder with \(L = 12\) holes and 36 anyons. The energies have been rescaled so that the two lowest eigenvalues match the CFT scaling dimensions.

The open boxes indicate the primary fields of the 7th minimal model with central charge \(c = 14/15\). The topological symmetry sectors are indicated with symbols \(1 \equiv y_{1,1}, \tau \equiv y_{-1,-1}\) and \(\tau + 1 \equiv y_{1,-1}\).

FIG. 4: CFT fields: Scaling dimensions \(h_{(r,s)} + \bar{h}_{(r,s)}\) of the primary fields in the \((D,A)\) modular invariant of the 7th minimal model with central charge \(c = 14/15\). On the right, we give momentum and topological symmetry assignments of these primary fields for our microscopic model.

\[ (r, s) \mid h_{(r,s)} + \bar{h}_{(r,s)} \mid k_x k_y \]

\begin{align*}
(1, 1) & \quad 0 \quad 0 \quad 0 \quad \varphi^0 \\
(3, 3) & \quad \frac{4}{15} \quad 0 \quad 0 \quad \varphi^2 \\
(5, 5) & \quad \frac{8}{15} \quad 0 \quad 0 \quad -1 \\
(7, 7) & \quad \frac{12}{15} \quad 0 \quad 0 \quad \varphi^0 \\
(2, 1) & \quad \frac{2}{15} \quad 0 \quad 0 \quad \varphi^0 \\
(4, 3) & \quad \frac{4}{15} \quad 0 \quad 0 \quad \varphi^0 \\
(6, 5) & \quad \frac{8}{15} \quad 0 \quad 0 \quad -1 \\
(6, 5) & \quad \frac{8}{15} \quad 0 \quad 0 \quad \varphi^{-1} \\
(8, 7) & \quad \frac{2}{15} \quad 0 \quad 0 \quad \varphi^{-2}
\end{align*}
FIG. 5: The energies of the lowest lying energy states around the critical point ($\theta = \pi/4$) as a function of the ‘dimerization’ $\theta$. Results are shown for system size $L = 10$. 

the vicinity of this critical point as shown in Fig. 5. Moving away from the critical point ($\theta = \pi/4$) corresponds to a dimerization of the model: in an alternative basis choice, discussed in detail in section II D, it becomes apparent that the rung and plaquette terms alternatingly act on even and odd ‘sites’. For $\theta \searrow \pi/4$, the four-fold ground-state degeneracy is lifted with one of the four ground states approaching the field with rescaled energy $2/45$ ($k_y = 0$), and two degenerate ground states moving to a rescaled energy $2/15$ ($k_y = 0$ and $k_y = \pi$). The single first excited state in this gapped phase softens towards the rescaled energy $4/15$ at the critical point. As we move into the adjacent gapped phase for $\theta < \pi/4$ only the field with rescaled eigenenergy $2/45$ moves back towards the ground-state, while the two other fields move upwards in energy and form a three-fold degenerate excited state.

2. Extended critical phase for $\theta \in (\pi, 3\pi/2)$

For negative coupling parameters $J_p, J_r < 0$, we find an extended critical phase around the point of equal coupling strength which in our circle phase diagram is opposite to the critical point discussed above. For the whole extent of this critical phase we can match the finite-size energy spectra to the $Z_8$ parafermion CFT with central charge $c = 7/5$. This theory is part of the sequence of $Z_k$ parafermion CFTs with conformal weights $\Delta_{\phi,j} = \frac{j(j+1)}{k+2} - \frac{m^2}{k}$, where $j = 0, \frac{1}{2}, 1, ..., k/2, |m| \leq j$ and $j - m =$ integer), in the notation of [10]. The details of the assignments for $k = 8$ can be found in Fig. 6 and Table II.

In order to verify that the critical phase around the exactly soluble point $\theta = 5\pi/4$ extends to the vicinity of the decoupling points $\theta = \pi$ and $\theta = 3\pi/2$, we consider an effective model where we fix all rung occupations to $\tau$-anyons. This assumption is exactly true at the decoupling point $\theta = 3\pi/2$. Implementing this constraint significantly reduces the size of the Hilbert space and allows us to numerically study this effective model for larger system sizes with up to 48 anyons.

The effective Hamiltonian in the reduced Hilbert space is given by

$$H^{\text{eff}} = -LJ_r - J_p \sum_{\text{plaq}} \delta_{\phi(p),1}. \quad (22)$$

The first term is a constant, and can thus be omitted which then turns the actual value of $J_p$ irrelevant. A positive $J_p$ corresponds to the limit $\theta \searrow 3\pi/2$, while a negative $J_p$ allows to study the limit $\theta \nearrow 3\pi/2$. 


For positive $J_p$, we find that the splitting of the ground state degeneracies goes to zero for $1/L \to 0$, and the energy gap approaches a finite value as shown in Fig. 8. This further supports the stability of the gapped topological phases up to, but excluding, the points $\theta = \pi$ and $\theta = 3\pi/2$ in our phase diagram.

For negative $J_p$, the rescaled energy spectrum of this effective model is critical and again matches (with much higher accuracy than at $\theta = \pi$) the $Z_8$ parafermion conformal field theory with central charge $c = 7/5$ as shown in Fig. 6. We can hence conclude that the whole quadrant $\theta \in (\pi, 3\pi/2)$ is occupied by an extended critical phase described by the same conformal field theory as the exactly solvable point $\theta = 5\pi/4$.

Approaching the endpoints of this extended critical phase at $\theta = \pi$ and $\theta = 3\pi/2$, the low-energy spectrum collapses into a flat band resulting in an extensive ground state degeneracy below an energy gap of size 1 at the points $\theta = \pi$ and $\theta = 3\pi/2$. Moving beyond these ‘decoupling points’ where one of the terms in the Hamiltonian vanishes, this extensive ground-state degeneracy is split again and a gap opens for $\theta < \pi$ and $\theta > 3\pi/2$, respectively, as the system enters the two gapped, topological phases discussed above.
FIG. 8: Energy gap $\Delta E(1/L)/J_p$ between the first excited state and the ground state, as well as the splitting of the ground state degeneracy, $\delta E(1/L)/J_p$, for the effective model Eq. 22. The two ground states become precisely degenerate only in the thermodynamic limit. The results indicate that the energy gap extrapolates to a finite value. Since the effective model is valid in the limit $\theta \to 3\pi/2+$, the gapped topological phase extends all the way up to this point.

FIG. 9: Rescaled energy spectrum of the effective model with $J_p$ negative ($L = 16$), and $Z_8$ parafermion CFT assignments. The topological symmetry sectors are indicated with symbols $1$ ($y_{1,1} = \varphi^2$), $\tau$ ($y_{1,\tau} = \varphi^{-2}$) and $\tau + 1$ ($y_{1,\tau} = -1$).
3. Topological stability of the critical phases

Both critical theories have a large number of rescaled energies (21) that are smaller than two. These eigenenergies are associated with operators whose correlation functions decay with scaling exponent $h + \bar{h} < 2$. Such operators are relevant in the renormalization group sense. This means that any operators $O$ with scaling dimensions (rescaled energies) $h + \bar{h} < 2$ which is invariant under all symmetries of the Hamiltonian may appear as an additional term in the latter and can thus drive the system out of the critical phase into a gapped phase or a different critical phase. For a critical phase to be stable there must hence exist a symmetry in the model such that the identity field (associated with the ground state) belongs to a different symmetry sector than all other fields $\phi$ with $h + \bar{h} < 2$ and $k_\phi = 0$ (fields at $k_\phi \neq 0$ do not obey the translational symmetry of the Hamiltonian).

Indeed, our model has an additional topological symmetry operator (11) that can stabilize the critical phases: There can be either no flux (denoted as 1-flux) or a $\tau$-flux entering the periodic ladder from one side, and a 1- or a $\tau$-flux leaving the ladder as illustrated in Fig. 10. There are hence three possibilities for possible flux assignments: (i) no flux is entering from above, and no flux is leaving [Fig. 10a], (ii) a $\tau$-flux is entering and leaving [Fig. 10b], or, (iii) a $\tau$-flux is entering from one side, and leaves through one or several plaquettes as shown in Fig. 10c). For each operator, one of the three scenarios applies and we can explicitly determine the topological sectors by considering the following hermitian symmetry operator (which commutes with the Hamiltonian)

$$Y[a, b, c] = \sum_{a', b'} \prod_{i=1}^{L} (F_{a_i}^{b_i} F_{a_{i+1}}^{b_{i+1}}) (F_{c_{i+1}}^{b_i} F_{c_i}^{b_{i+1}}) |a', b', c\rangle,$$

where $|a, b, c\rangle = |a_1, b_1, c_1, a_2, b_2, c_2, ..., a_L, b_L, c_L\rangle$ are labels according to Fig. 1. This operator inserts additional $\tau$-loops parallel to the two 'spines' of the ladder. As in the case of the plaquette term Eq. (16), this is done by connecting them to the ladder with 1-particles. The flux through each of these two additional $\tau$-loops can be either 1 or $\tau$, where a 1-flux yields a factor of $S_i^x/S_{i+1}^x = \varphi$, and a $\tau$-flux gives $S_i^x/S_{i+1}^x = -\varphi^{-1}$ (note that a $S$-transformation has to be performed in order to obtain the flux through the additional $\tau$-loops). Hence there are three possible eigenvalues of $Y$: $y_{1,1} = \varphi^2$ (scenario i), $y_{\tau, \tau} = \varphi^{-2}$ (scenario ii) or $y_{1, \tau} = -\varphi^{-1}\varphi = -1$ (scenario iii).

We numerically evaluate the topological symmetry sectors in the two critical phases (see Tables I and II, and Figs. 3, 6 and 9). At the critical point separating the topological phases ($\theta = \pi/4$), we find that the relevant operators can be classified according to $s = 1 \leftrightarrow y_{1,1}$, $s = 3, 7 \leftrightarrow y_{\tau, \tau}$, $s = 5$ $\leftrightarrow y_{1, \tau}$. In particular, only one operator, $\phi_{(2,1)}$, is in the same topological symmetry sector as the ground state, i.e. the identity field $\phi_{(1,1)}$. It is this field that drives the system out of the critical phase when varying the coupling constant $\theta$. With the scaling dimension of this operator being $x = 2/3$ the gap opens as $\Delta E(\theta) \propto |\theta - \pi/4|^\nu$ on either side of the critical point, where $\nu = 1/(2 - 2/3) = 3/4$. In the second critical phase, $\theta \in (\pi, 3\pi/2)$, the topological symmetry assignments of the relevant operators are given by $r = 0 \leftrightarrow y_{1,1}$, $r = 2, 6 \leftrightarrow y_{\tau, \tau}$, $r = 4 \leftrightarrow y_{1, \tau}$. In particular, there is no relevant field in the same topological symmetry sector as the ground state, which implies that there is no symmetry-allowed relevant operator in this gapless theory and the critical point must be part of an extended gapless phase. This observation demonstrates that our observation (from exact diagonalization studies) that the extended critical phase in the quadrant $\theta \in (\pi, 3\pi/2)$ is described by the same $Z_8$ parafermion CFT with central charge $c = 7/5$ is correct.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig10.png}
\caption{Topological symmetry sectors: a) No $\tau$-flux enters or leaves the ladder. b) A $\tau$-flux enters from one side and leaves at the other side. c) A $\tau$-flux enters from one side and leaves through a plaquette.}
\end{figure}
D. Analytical solution

Our ladder model defined by the Hamiltonian in Eq. (2) in the main part of the paper can be solved exactly at the two critical points $\theta = \pi/4$ and $\theta = 5\pi/4$ (see the phase diagram in Fig. 5 of the main text). The key observation leading to this exact solution is that the topological structure of our model implies that its Hilbert space is in fact built on the so-called $D_6$-Dynkin diagram, which is drawn below in Fig. 11.

![Dynkin diagram $D_6$.](Fig. 11)

The Dynkin diagram indeed appears very naturally: let us make a change of basis for our Hamiltonian as illustrated in Fig. 12. This new choice of basis (drawn on the left), which arises from a different decomposition of the high-genus surface, is related to the original one (drawn on the right) by a simple $F$-transformation. In particular, consider the new basis in the left part of Fig. 12: with the even-numbered 'sites' (which correspond to the original rungs) we associate a label $d_i = 1$ or $d_i = \tau$ (the flux through that cross-section of the surface). With the odd-numbered 'sites' (which correspond to the original plaquettes) we associate a variable consisting of a pair of labels, $(a_i, b_i)$ which can assume four values, i.e., $(a_i, b_i) = (1, 1), (1, \tau), (\tau, 1), (\tau, \tau)$, and denotes the pair of fluxes through the two cross-sections of the surface at the position of the plaquette. The allowed fusion channels at the vertices where variables $(a_i, b_i)$ and $d_{i\pm k}$ meet then correspond precisely to the condition that they be adjacent nodes on the Dynkin diagram of the $D_6$ Lie algebra, as illustrated in Fig. 11 above. For example, a local label $(a_i, b_i) = (\tau, \tau)$ at an odd-numbered 'site' $i$ allows for labels $d_{i-1} = 1$ and $d_{i+1} = \tau$ at the neighboring even-numbered sites, which is reflected in the fact that label $(\tau, \tau)$ is connected by a line to both labels 1 and $\tau$ in the Dynkin diagram.

![Two possible basis choices corresponding to different decompositions of the high-genus surface.](Fig. 12)

In summary, the elements of this new basis of the Hilbert space on which the Hamiltonian acts are of the form

$$|\vec{a}\rangle := |\ldots, \alpha_{i-1}, \alpha_i, \alpha_{i+1}, \ldots\rangle,$$

where $\alpha_j = d_j$ if $j$ is even, and $= (a_j, b_j)$ if $j$ is odd] denotes a point on the $D_6$-Dynkin diagram representing the flux through the high-genus surface at the 'site' $j$ of the chain. The sequence of $\alpha_j$ must satisfy the condition that $\alpha_{j+1}$ is a nearest neighbor site of $\alpha_j$ on the $D_6$-Dynkin diagram.

In this new basis, the rung and plaquette terms $H^R$ and $H^P$ of our ladder Hamiltonian

$$H = -J_p \sum_{i \text{ odd}} H^P_i - J_r \sum_{i \text{ even}} H^R_i,$$

take on the following form [12]

$$H^P_{i}|a_i, b_i\rangle = \sum_{k=1, \tau} \frac{d_i}{d_k} \sum_{a_i' b_i'} (F^b_{b_i a_i a_i} a_i') (F^b_{d_{i+1} a_i b_i} a_i') |a_i', b_i'\rangle,$$

$$H^R_i|d_i\rangle = \sum_{d_i'} (F^b_{b_i a_i a_i + 1} d_i') (F^b_{d_{i-1} a_i b_i + 1} d_i') |d_i'\rangle.$$

![Phase diagram and Dynkin diagram](Fig. 12)
In fact, these terms can be seen to form a representation of the Temperley-Lieb algebra [13] which arises from the \( D_v \)-Dynkin diagram, and has “d-isotopy” parameter \( D = \sqrt{1 + \varphi^2} = 2 \cos(\pi/10) \), the total quantum dimension of our Fibonacci theory. Specifically, consider the operators \( e_i \) constructed from the components \( v_\alpha = \sin(\alpha \pi/10) \) \( (\alpha = 1, \ldots, 6, v_1 = v(1,1), v_2 = v(1), v_3 = v(2), v_4 = v(2), v_5 = v(3,1) = v(3,3)) \) of the (‘Perron Frobenius’) eigenvector corresponding to \( D \), the largest positive eigenvalue of the adjacency matrix of the \( D_v \)-Dynkin diagram [14],

\[
e_i \mid \ldots, \alpha_{i-1}, \alpha_i, \alpha_{i+1}, \ldots \rangle := \sum_{\alpha_i} \left( (e_i)^{\alpha_{i+1}}\right)^{\alpha_i} \mid \ldots, \alpha_{i-1}, \alpha_i', \alpha_{i+1}, \ldots \rangle, \tag{27}\]

where \( \left( (e_i)^{\alpha_{i+1}}\right)^{\alpha_i} = \delta_{\alpha_i-1, \alpha_i+1} \sqrt{v_{\alpha_i} v_{\alpha_i'}} / v_{\alpha_i-1} v_{\alpha_i+1}. \)

These operators form a known representation [15] of the Temperley-Lieb algebra with “d-isotopy” parameter \( D \), i.e.

\[
e_i^2 = D e_i, \quad e_i e_{i \pm 1} e_i = e_i, \quad [e_i, e_j] = 0 \quad \text{for} \quad |i - j| \geq 2. \tag{28}\]

Now one can check that the rung and plaquette terms, Eq. (26), of the Hamiltonian in the new basis, Eq. (25), are proportional to these operators, i.e.

\[
H_i^P = \frac{1}{D} e_i \quad \text{for} \quad i \text{ odd}, \quad H_i^R = \frac{1}{D} e_i \quad \text{for} \quad i \text{ even}. \tag{29}\]

The Hamiltonian Eq. (26) is in fact that corresponding to the (integrable) restricted-solid-on-solid (RSOS) statistical mechanics lattice model based on the \( D_v \)-Dynkin diagram [15]. Specifically, the two-row transfer matrix \( T := T_2 T_1 \) of this lattice model

\[
\begin{array}{c}
\alpha_{2n-1}^{'} \\
W[2n] \\
\alpha_{2n+1}^{'} \\
\alpha_{2n-1} \\
W[2n+1] \\
\alpha_{2n+1} \\
\end{array}
\]

is written in terms of Boltzmann weights \( W[i] \) assigned to a plaquette \( i \) of the square lattice

\[
T_1 := \prod_n W[2n], \quad \text{and} \quad T_2 := \prod_n W[2n+1] \tag{30}\]

with

\[
W[i]^{\alpha}_{\alpha'} = \left\{ \begin{array}{c}
\sin \left( \frac{n}{10} - u \right) / \sin \frac{n}{10} \left[ 1^{\alpha'}_{\alpha} + \sin u / \sin \frac{n}{10} \right] \\
\delta_{\alpha', \alpha} \end{array} \right\}. \tag{31}\]

The parameter \( u > 0 \) is a measure of the lattice anisotropy, \( 1 \) is the identity operator, and

\[
e_i^{[\alpha]} := \left[ \prod_{m \neq i} \delta_{\alpha', \alpha_m} \right] \left( e_i \right)^{\alpha_{i+1}}_{\alpha_i}. \tag{32}\]

The Hamiltonian of the so-defined lattice model is obtained from its transfer matrix by taking, as usual [16], the extremely anisotropic limit, \( 0 < u \ll 1 \),

\[
T = \exp \{ -a (H + c_1) + O(a^2) \}, \quad a = \frac{u}{D \sin \pi/10} \ll 1,
\]

yielding

\[
H = - \sum_i \frac{1}{D} e_i. \tag{33}\]
Since, due to Eq. (29), the operators ‘$\frac{1}{2} e_i$’ are nothing but the rung and plaquette operators, we have thus demonstrated that the Hamiltonian of the RSOS statistical mechanics model based on the Dynkin diagram $D_6$ coincides with the Hamiltonian, Eq. (25), of our ladder model.

The RSOS model based on $D_6$ is known [17, 18] to provide an (integrable) lattice realization of the $(D, A)$ modular invariant [9] of the 7th unitary minimal CFT of central charge $c = 14/15$. In particular, the Hamiltonian of Eq. (2) of the main text at angle $\theta = \pi/4$ will yield the spectrum of that CFT. This exact analytical result is borne out precisely by our numerical (exact diagonalization) studies reported in subsection (II C 1). This CFT with central charge $c = 14/15$ describes the quantum critical point of a 1 + 1 D quantum system, our ladder model. While we cannot make an exact statement for the related 2 + 1 D quantum model, we note that Fendley has recently discussed this 2 + 1 D quantum critical point from a 2 + 0 D perspective [19] by considering a one-parameter family of wavefunctions connecting the ground-state wavefunctions of the two extreme limits of the 2 + 1 D model in Eq. (1) of the main manuscript. For a certain value of the parameter $\theta$ he finds a conformal quantum critical point whose ground-state correlators are written in terms of this same $c = 14/15$ CFT.

Another version of this lattice model yielding in the anisotropic limit the negative, $-H$, of the Hamiltonian in Eq. (33) is also integrable and provides [20] a lattice realization of the $Z_8$ parafermionic CFT of central charge $c = 7/5$. In particular, the Hamiltonian of Eq. (2) of the main text at angle $\theta = 5\pi/4$ will yield the spectrum of that CFT. Again, this exact analytical result is borne out precisely by our numerical (exact diagonalization) studies reported in subsection (II C 2).

### III. THE HONEYCOMB LATTICE MODEL

In this section, we discuss details of the “honeycomb lattice model” whose Hamiltonian is given by Eq. (1) in the paper. We first define the plaquette term of the model and then discuss two limiting phases of the model.

#### A. The Hamiltonian

In analogy to the plaquette term in the ladder model, Eq. (16), the plaquette term of the honeycomb lattice model (Eq. (1) in the main text) is defined by

$$\delta \phi_{(p),1} \left| \begin{array}{cc} a & b \\ e & f \\ \zeta & \gamma \\ \delta & \zeta \\ \epsilon & \gamma \\ d & c \end{array} \right| = \sum_{s=1,3} \frac{d_s}{D^2} \sum_{\alpha',\beta',\zeta'} (F_{\gamma,\alpha'} s')_\delta (F_{\beta,\gamma'} s')_\epsilon \left( F_{\gamma,\alpha'} s' \right)_\delta (F_{\beta,\gamma'} s')_\epsilon \left( F_{\gamma,\alpha'} s' \right)_\delta (F_{\beta,\gamma'} s')_\epsilon$$

where the additional two edges of a plaquette are reflected in two additional $F$-transformations. Again, we can parametrize the coupling constants on a circle as $J_p = \cos(\theta)$ and $J_e = \sin(\theta)$.

#### B. Excitations

We briefly mention the elementary excitations of this model. In the ‘two-sheets’ phase, which corresponds to couplings $\theta = 0$ ($J_p = 1$, $J_e = 0$), the elementary excitation is a single plaquette with a $\tau$-flux giving rise to a single ‘hole’ as illustrated on the left in Fig. 13. These excitations are gapped with a gap size of $J_p$ and will delocalize for small couplings $J_e \neq 0$ forming quasiparticle bands. Similar to the ladder model the dispersion of this quasiparticle band can be calculated perturbatively around the ‘two-sheets’ limit.

In the opposite limit of ‘decoupled spheres’, which corresponds to couplings $\theta = \pi/2$ ($J_e = 1$, $J_p = 0$), the elementary excitation is a ‘plaquette ring’ where all edges around a given plaquette have $\tau$-fluxes, as illustrated on the right in Fig. 13. Again, a perturbative analysis allows to qualitatively describe the quasiparticle band.

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[2] For a pedagogical introduction, we refer to J. Preskill, Lecture notes on quantum computation, available online at http://www.theory.caltech.edu/~preskill/ph229.

[3] A quantum group deformation with deformation parameter $q$ being a root of unity.
The $F$-matrix is an extension (truncation) of the $6-j$ symbol of $SU(2)$ to the quantum group $SU(2)_q$ for deformation parameter $q$ at root of unity [5].

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The plaquette term is formulated in the same manner as the one in Eq. 16. The rung term is a projector of the fusion product of anyons $a_{i-1}$, $b_{i-1}$, as well as of anyons $a_{i+1}$ and $b_{i+1}$ onto the trivial particle.

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This is the matrix whose only non-vanishing matrix elements are $A_{\alpha,\alpha'} = 1$ when $\alpha$ and $\alpha'$ are nearest neighbors on the Dynkin diagram.

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**FIG. 13**: The elementary excitations above the extreme ground states illustrated in Fig.1 of the main part of the paper: a single plaquette flux in the ‘two-sheet’ phase, and a single plaquette in the ‘multi-sphere’ phase.