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Joint entanglement of topology and polarization enables errorprotected quantum registers

David S Simon^{1,2,5}, Shuto Osawa² and Alexander V Sergienko^{2,3,4}

- Department of Physics and Astronomy, Stonehill College, 320 Washington Street, Easton, MA 02357, United States of America
- ² Department of Electrical and Computer Engineering, Boston University, 8 Saint Mary's St., Boston, MA 02215, United States of America

Photonics Center, Boston University, 8 Saint Mary's St., Boston, MA 02215, United States of America

Department of Physics, Boston University, 590 Commonwealth Ave., Boston, MA 02215, United States of America Author to whom any correspondence should be addressed.

E-mail: simond@bu.edu, sosawa@bu.edu and alexserg@bu.edu

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Abstract

Linear-optical systems can implement photonic quantum walks that simulate systems with nontrivial topological properties. Here, such photonic walks are used to jointly entangle polarization and winding number. This joint entanglement allows information processing tasks to be performed with interactive access to a wide variety of topological features. Topological considerations are used to suppress errors, with polarization allowing easy measurement and manipulation of qubits. We provide three examples of this approach: production of two-photon systems with entangled winding number (including topological analogs of Bell states), a topologically error-protected optical memory register, and production of entangled topologically-protected boundary states. In particular it is shown that a pair of quantum memory registers, entangled in polarization and winding number, with topologically-assisted error suppression can be made with qubits stored in superpositions of winding numbers; as a result, information processing with winding number-based qubits is a viable possibility.

1. Introduction

States with integer-valued topological invariants, such as winding and Chern numbers, exhibit a variety of physically interesting effects in solid-state systems [1–5], including integer and fractional quantum Hall effects [6–10]. When systems with different values of topological invariants are brought into contact, states arise that are highly localized at the boundaries. These edge or boundary states have unusual properties; for example, in two-dimensional materials they lead to unidirectional conduction at the edges, while the interior bulk remains insulating. Because of the inability to continuously interpolate between discrete values of the topological invariant, these surface states are protected from scattering and are highly robust against degradation, making them prime candidates for use in error-protected quantum information processing.

Optical states with similar topological properties can be produced by means of photonic quantum walks in linear-optical systems [2, 11–23]. Photonic walks have demonstrated topological protection of polarization-entanglement [24] and of path entanglement in photonic crystals [25].

Optical systems are useful laboratories to study topologically-nontrivial states, due to the high level of control: system properties can be varied over a wide range of parameters, in ways not easy to replicate in condensed matter systems. In the quest to carry out practical quantum information processing tasks, it is of great interest to examine more closely the types of topologically-protected states that can be optically engineered. Those that also entangled are of particular interest for quantum information applications.

The goal here is to entangle states associated with distinct topological sectors, and to do so in a way *that allows this entangled topology to be readily available for information processing*. Specifically, linear optics will be used to produce: (i) entangled topologically-protected boundary states, (ii) winding-number-entangled bulk states, and (iii) an entangled pair of error-protected memory registers. To create the states, a source of initial polarization-



Figure 1. Directionally-unbiased threeport unit: three beam splitters form a triangular array with three external ports and three internal edges. One port of each beam splitter passes through a phase shifter and then reflects back onto itself via a mirror. Altering the phase shift imparted at these mirror units allows control of the device properties by altering the interference between paths. Adapted figure with permission from [27], Copyright (2016) by the American Physical Society.

entangled light is necessary, specifically type-II spontaneous parametric down conversion (SPDC) in a $\chi^{(2)}$ nonlinear crystal. Taking this initial state as given, all further processing requires only linear optical elements.

Topological invariants characterize global properties of systems and cannot be easily distinguished by localized measurements. This difficulty in measurement limits their use in many applications. That problem is solved here by linking topology to a more easily-measured variable, polarization. Polarization and winding number will be tightly correlated (and in fact, jointly entangled with each other), but will serve distinct purposes: winding number provides stability against perturbations, while polarization allows easy access and measurement.

We confine ourselves to one-dimensional systems. After reviewing directionally-unbiased optical multiports, it is shown how arrays of such multiports can produce entangled pairs of bulk states associated with Hamiltonians of different winding number, as well as entangled pairs of topologically-protected edge states localized near boundaries between regions of different topology.

A variation of the same idea then allows single qubits or entangled qubit pairs to be stored in a linear optical network as winding numbers. Topological protection of boundary states is well-known, but less widely recognized is the fact that bulk wavefunctions also have a degree of resistance to changes in winding numbers [26]. This effect is discussed in the appendix and will be used to reduce polarization-flip errors of qubits stored in the optical register, greatly reducing the need for additional error correction.

2. Directionally-unbiased multiports and topological states

In standard beam splitters and multiports, photons cannot reverse course to exit back out the input port. In [27], a generalized multiport was introduced which allows exit with some probability out any port, including the input. These directionally-unbiased multiports have adjustable internal parameters (reflectances and phase shifts) that allow the exit probabilities at each port to be varied. The three-port version is shown schematically in figure 1.

Directionally-unbiased multiports are linear optical devices with the input/output ports connected via beam splitters to vertex units of the form shown in the inset of figure 1. Each such unit contains a mirror and

phase shifter. The beam splitter-to-mirror distance $\frac{d}{2}$ is half of the distance d between the vertex units in the

multiport. The phase shifter provides control of the properties of the multiport, since different choices of phase shift at the vertices affect how the various photon paths through the device interfere with each other. These devices and some of their applications have been studied theoretically in [27–29] and experimentally demonstrated in [30].

If the unit is sufficiently small (quantitative estimates of the required size and other parameter values may be found in [27]) then its action can be described by an $n \times n$ unitary transition matrix \hat{U} whose rows and columns correspond to the input and output states at the ports. If the internal phase shifts at all the mirror units are known, then an explicit form of the unitary transition matrix \hat{U} can be given. Here, we restrict ourselve to the





three-port and assume that all three vertices are identical, in which case [28]:

$$\hat{U} = \frac{e^{i\theta}}{2 + ie^{i\theta}} \begin{pmatrix} 1 & ie^{-i\theta} - 1 & ie^{-i\theta} - 1 \\ ie^{-i\theta} - 1 & 1 & ie^{-i\theta} - 1 \\ ie^{-i\theta} - 1 & ie^{-i\theta} - 1 & 1 \end{pmatrix},$$
(1)

where θ is the total phase shift at each mirror unit (including both the mirror and the phase plate). The rows and columns refer to the three ports *A*, *B*, *C*.

Arrays of three-ports and phase shifters can simulate a range of discrete-time Hamiltonian systems [29], including some with distinct topological phases [28]. The array acts as a lattice through which photons propagate, leading to photonic analogs of Brillouin zones and energy bands. By altering multiport parameters, systems can be simulated [28] in which Hamiltonians have different winding numbers as one wraps around a full Brillouin zone. As is well known from solid-state physics [1, 3–5], at the boundaries between regions with distinct winding number localized boundary or edge states appear. These edge states are highly stable due to topological protection; different discrete winding numbers on the two sides prevent the state from being destroyed by small, continuous perturbations.

The basic building block for the optical systems described below is the diamond-graph structure [28, 31–33] formed by two unbiased three-ports and an additional phase shifter. This is shown in figure 2, where the unbiased three-port is represented by a vertex connecting three edges. The unit cell for the lattice structures will be formed by two such diamond graphs (figure 3), and so each cell will contain four multiports. The phase shifts in each of the two diamonds may be different, $\phi_a \neq \phi_b$. When a string of these unit cells are connected end-to-end, photons inserted into the chain exhibit quantum walks. The resulting system is governed by a Hamiltonian which can have nontrivial topological structure: depending on the values of the phases ϕ_a and ϕ_b , the winding number ν can take a value of either 0 or 1 [28]. When a chain of winding number $\nu = 1$ is connected to a chain with $\nu = 0$, localized topologically-protected edge states appear at the boundary [28].

Additional discussion of topological aspects of one-dimensional models is given in the appendix, along with numerical simulations displaying these properties in systems formed by chains of unbiased multiport.

3. Jointly-entangled topologically-protected bulk states

Start with a polarization-entangled photon source, type-II SPDC in a nonlinear crystal. With appropriate filtering and phase shifts, the two-photon output can be taken to be an entangled Bell state

$$|\psi^{\pm}\rangle = \frac{1}{\sqrt{2}} (|H\rangle_1 |V\rangle_2 \pm |V\rangle_1 |H\rangle_2), \tag{2}$$



where 1 and 2 refer to two spatial modes. We wish now to create from this a state of entangled winding number. Polarization entanglement should remain intact, to use for control and measurement purposes.

It should be noted that we are making a slight abuse of terminology here: the winding number is associated with the Hamiltonian, not strictly speaking with the state. But as discussed in the appendix, transitions of bulk states between spatial regions or parameter values with Hamiltonians of different winding number can be arranged to be strongly suppressed. Therefore, as a practical matter, under appropriate conditions one may to a high degree of accuracy think of the winding number as being associated with the state as well.

Consider two chains of unit cells like those of figure 3, distinguishing the upper (*u*) and lower (*l*) chains (figure 4). Using the states $|\Psi^{\pm}\rangle$ as input, each down conversion photon is directed into one of the two chains, so that the labels 1 and 2 in equation (2) now correspond to *u* and *l*. The photons may be coupled into the system via a set of optical circulators and switches, as described in [29]. The circulators are used only to couple photons into the system initially, and to couple them out for measurement at the end; they play no role in the actual operation of the system in between. We take ϕ_b to be polarization-dependent, but we may assume that the action of the phase shifters producing ϕ_a are independent of polarization. In this way, it is arranged for *H* states to encounter equal phases $\phi_a = \phi_b$, while *V* states see $\phi_a \neq \phi_b$. The polarization-dependent phase shifts are easily implemented with thin slices of birefringent material or, if real-time control of the phase shifts is desired, with Pockels cells. In the visible and near infrared, it is easy to find crystals with low absorption and strong birefringence; calcite is one example. Other materials with similar properties can be found for other spectral ranges. So it should be relatively easy to produce the necessary phase shifts with negligible effect on performance. The use of electro-optical effects can enable fine adjustment of the phase shift in each cell if necessary.

If ϕ_b is chosen correctly, then the *H* part is put into a state with winding number $\nu = 0$ and the *V* part into a state with $\nu = 1$. Then the vertically- and horizontally-polarized states will be eigenstates of Hamiltonians with respective winding numbers $\nu_V = 1$ and $\nu_H = 0$. The final state is therefore of the form

$$\frac{1}{\sqrt{2}}(|0_H\rangle_u|1_V\rangle_l \pm |1_V\rangle_u|0_H\rangle_l),\tag{3}$$

where the 0 and 1 represent winding number values of the Hamiltonians that govern their time evolution, while u and l denote the spatial modes in the upper and lower chains. The state is written in condensed form here; a more explicit expression, including the spatial dependence of the wavefunction is given in the appendix. The photons now form winding number-entangled qubit pairs. More generally, both ϕ_a and ϕ_b may both be allowed to be polarization dependent; all that matters is that the polarization-dependent combination (ϕ_a , ϕ_b) leads to each polarization state seeing a Hamiltonian of different winding number.

Usually, the global property of winding number is difficult to determine by local measurements. This is especially true for single-photon states which are usually destroyed by the measurement process, so that multiple measurements cannot be carried out to determine the global state. But here polarization and winding number remain coupled. This jointly-entangled structure allows the variables to play disparate roles: the discrete winding number leads to topologically-enforced stability, while polarization, being defined locally, makes the photon state easy to measure. Polarization can be determined by a single local measurement, allowing the global winding number to be inferred from its value.

Suppose that a perturbation occurs to the system. Normally, this might cause the photon's polarization to change (a polarization-flip error). For example, a horizontally-polarized state of winding number ν might try to flip into a vertical state: $|\psi_{\nu}\rangle_H \rightarrow |\psi_{\nu'}\rangle_V$, where ν' is the final winding number. But as discussed in more detail in [26] and in the appendix, unless the perturbation is strong enough to severely and globally alter the entire system, transitions from eigenstates of one bulk Hamiltonian to those of a Hamiltonian of different winding number are suppressed. If the hopping parameters are chosen appropriately, the amplitudes for these transitions



can be made arbitrarily small. This means that, to a high level of certainty, the initial and final winding numbers can be assumed to be equal: $\nu' = \nu$. However, due to the way the system was constructed, there are no allowed states that have V polarization and which propagate according to a Hamiltonian of winding number ν , so the polarization flip is suppressed.

Thus, barring extreme alterations of the system, polarization-flip errors are greatly reduced. The suppression of polarization errors occurs without loss of photons, and so there is no damage to any coherence or entanglement present in the system.

4. Topologically-protected quantum memory registers

A basic ingredient needed for quantum computing is a quantum memory unit capable of storing a logical qubit $\alpha |0\rangle + \beta |1\rangle$. Such units need to have read/write capability and should be resistant to bit-flip errors. This can be achieved by a variation on the strategy above. Once again, topological stability suppresses errors, with polarization used for reading and writing stored values.

A schematic depiction of the memory register is shown in figure 5. As before, assume that $\phi_b = \phi_a$ for horizontal polarization and $\phi_b \neq \phi_a$ for vertical. When a horizontally-polarized photon enters the ring it is associated with winding number $\nu = 0$, but for appropriate values of ϕ_b a vertically polarized photon will have $\nu = 1$. The winding number then serves as the logical bit being stored. Readout of bit values requires only simple polarization measurements. Since the input photon may be in any arbitrary superposition of polarization states, the ring can be used to store any possible qubit value. In general, an input polarization qubit $\alpha |H\rangle + \beta |V\rangle$ is stored in a winding-number/polarization qubit, $\alpha |0_H\rangle + \beta |1_V\rangle$, where 0 and 1 are winding number.

Since photons at normal energies do not mutually interact to a significant degree, multiplexing is possible. Multiple qubits can be stored on a single ring by using photons of different frequency; addressing the desired qubit then simply requires opening an exit channel from the ring and using a filter or dichroic mirror with the appropriate frequency-transmission range. Reading out a qubit value consists of measuring the polarization.

5. Entangled quantum memory registers

Notice that the register of figure 5 consists of one strand of the structure of figure 4 wrapped into a circle; the compactification to a circle makes its use in a larger system more practical, but is not necessary for operation. Each strand of figure 4 is already capable of serving as a quantum memory. One could use *both* strands of figure 4 (either in the original linear configuration or compactified to a double-circle structure); for the polarization entangled input of equation (2) the result would be two entangled quantum memory registers, with error suppression provided by the winding number entanglement.

Such an entangled quantum register could provide novel applications in quantum computing. As one example, using an entangled pair of memory registers would enhance security; by using a subset of the registers for security checks instead of for computation, a data breach would be detectable as a sudden loss of entanglement. Similarly, if a register malfunctions then the location of the malfunction should be easy to track down through the drop in the degree of entanglement of the corresponding pair.





6. Topologically-protected entangled boundary states

The setup of figure 4 can be generalized to produce one further effect. First, the polarization-dependent phase can be made different in the upper and lower branches (ϕ_b in upper branch and ϕ_d in lower). Then, a boundary plane can be introduced perpendicular to the chains, such that the polarization-dependent phase will change suddenly when the plane is crossed ($\phi_b \rightarrow \phi_c$ in upper branch and $\phi_d \rightarrow \phi_f$ in lower), as shown in figure 6. As shown in [28], if the phase values on the two sides of the boundary are chosen correctly, then topologically-protected states appear that are tightly localized on the boundary point. Unlike the approximate winding number preservation in the bulk, the topological protection of the boundary state is exact and has been demonstrated in a number of different solid state and optical systems.

Taking the simplest case, suppose $\phi_b = \phi_d$ and $\phi_c = \phi_f$, so that the upper and lower chains are identical. Then for polarization-entangled input (equation (2)), the boundary state will be in a superposition of two positions (points *A* and *B*), as a Schrodinger cat state. These entangled boundary states will be verticallypolarized and would be topologically protected. Considering just the state at the boundary, let $|e\rangle$ and $|\emptyset\rangle$ represent, respectively, the presence or absence of a localized edge state. Then the state on the boundary plane will be of the form

$$\frac{1}{\sqrt{2}}(|e_V\rangle_u|\varnothing_H\rangle_l \pm |\varnothing_H\rangle_l|e_V\rangle_u),\tag{4}$$

where we assume as before that vertical polarizations see different winding numbers on the two sides of the boundary and horizontal polarizations do not. Here, as before, *u* and *l* label whether the spatial mode is in the upper and lower branch. Note that this entanglement is distinct from path entanglement; photons exist simultaneously in *both* branches, even if edge states are absent from a given branch.

Another possibility is to take $\phi_a \neq \phi_b$ for the *vertical* polarization in the upper chain, but in contrast to take $\phi_a \neq \phi_b$ for *horizontal* polarization in the lower chain; in this case, there would be an entangled state which is a superposition of having either localized boundary states at *both* A and B or at *neither*:

$$\frac{1}{\sqrt{2}}(|e_V\rangle_u|e_H\rangle_l \pm |\varnothing_H\rangle_u|\varnothing_V\rangle_l).$$
(5)

The states of equations (4) and (5) are maximally entangled, with entropy of entanglement equal to 1, and may be thought of as topologically-stable implementations of Bell states; these states can also be used to store entangled qubits.

Edge states appear due to interference between various amplitudes for quantum walks through each chain; they should survive as long as the photons remain contained inside the system, coherent and unmeasured. Small perturbations in the refractive index or in path lengths along the photon trajectories should not disturb the results. For example, in the appendix numerical results are displayed (figure A6) that show that the boundary state persists over a wide range of *v* and *w* parameters, as long as the winding numbers do not change.

7. Conclusion

We have proposed a hybrid strategy for quantum information processing, in which local and global properties (polarization and winding number) are jointly-entangled, allowing one to simultaneously exploit the benefits of both: discreteness of global, topological properties affords stability and error suppression, while local properties are easy to manipulate and measure. This has applications in producing entangled topological states and in designing quantum registers (possibly in entangled pairs) with topologically-assisted reduction of bit-flip errors.

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two chains will support states of different winding number, with stable localized states appearing at the boundary.

Besides reducing bit-flip errors, the use of discrete topological quantities also helps maintain loss of entanglement through the same mechanism: if there are no nonentangled joint states that a photon pair can scatter into, then the entanglement will remain robust. This can help avoid some of the problems that occur in many approaches to quantum computing as a result of the fragility of entangled states.

Efficient measurement of topological quantum numbers has been a longstanding problem. Although other methods of measuring topological variables in photonic systems have been proposed or carried out [34–39], they require determination of probability amplitudes by measurements on multiple photons. The method given here has the advantage of being able to operate at the single photon level.

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Appendix. Winding number, wavefunctions, and topological state protection

In [28] it was shown that the chain of directionally-unbiased multiports in figure A1(a) is a photonic equivalent of the Su–Schrieffer–Heeger (SSH) system used to model the behavior of polymers. In this appendix, we briefly review the topological properties of this system and verify via numerical simulations that these properties hold for chains of unbiased multiports; in particular, we numerically demonstrate the resistance of the wavefunction to enter regions of different winding number.

The one-dimensional SSH system is composed of a periodic string of unit cells, with two alternating subcells *A* and *B* of distinct reflection and transmission amplitudes within each cell. This forms a two-level system, with a momentum-space Hamiltonian of the form

$$\hat{H} = \int_{-\pi}^{\pi} dk \ E(k) \ \frac{\boldsymbol{d}(k) \cdot \sigma}{|\boldsymbol{d}(k)|} \otimes |\boldsymbol{k}\rangle \langle \boldsymbol{k}|, \tag{A.1}$$

where *k* is the quasi-momentum and the integral is over a full Brillouin zone. Due to the existence of a chiral sublattice symmetry with generator $\Gamma = e^{-i\frac{\pi}{2}\sigma_z}$ that anticommutes with the Hamiltonian, the spectrum is symmetric, with energies coming in opposite-sign pairs, $\pm E(k)$. The gap between the two energies only closes when *A* and *B* have equal transmission amplitudes.

The Hamiltonian is determined by the vector

$$d(k) = (v + w \cos k)\hat{x} + (w \sin k)\hat{y},$$
 (A.2)

where v(k) and w(k) are, respectively, the intracell hopping amplitude between A and B and the intercell hopping amplitude. \hat{x} and \hat{y} are basis vectors in the two-dimensional Hilbert space spanned by the A and B substates. In the simplest SSH model, v and w are constants, and d(k) traces out a circle in k space; for the unbiased multiport chains, v and w are continuous functions of k, so that the circular paths become continuously deformed. Since topological properties of systems are unchanged by continuous deformations of the parameters, the unbiased multiport system has the same topological properties as the SSH model, as is verified numerically below. d(k) must remain orthogonal to the chiral symmetry generator σ_z in order to preserve the symmetry. However, the direction of the vector d(k) becomes undefined at $k = \pi$ when v = w. The gap between the energy levels closes at the parameter values for which this occurs. For other parameter values, d(k) must avoid the origin, leading to a distinction between values at which the path traced out by d(k) encloses the origin and those for which it does not. The latter cases are topologically trivial, with bulk winding number v = 0, while the former cases have nontrivial winding number v = 1. The winding number cannot change without d(k) crossing the singular point and the energy gap closing.

When the Hamiltonian changes abruptly from one topological state to different one (say from winding number 0 to winding number 1), highly-localized states appear at the boundary between the two topological regions. It has been demonstrated in a number of different physical contexts that there is a form of topological protection attached to these states: no continuous localized disturbance can destroy the state or cause a change in the winding number on the two sides of it. In particular, this has been demonstrated experimentally for a number of photonic systems [20, 21], including systems based on photonic quantum walks [2, 16].

Further, it can be shown [26] that a wavefunction initially present in the bulk on one side of the boundary tends to resist transmission into the second, topologically distinct, bulk region. The wavefunction instead shows a tendency to reflect back into the original region when it encounters the boundary. This tendency can be made nearly complete by a wise choice of the parameters in the Hamiltonian, as will be discussed below in the context of the SSH system.

This topologically-assisted suppression of transitions is the key to why the systems in sections 3–6 are of interest. The polarization will be linked with winding number, so that the suppression of transitions between different winding number states will suppress polarization changes.

Label each unit cell of the lattice by an integer position label *n*. Each such unit cell has two subunits or 'substates', labeled *A* and *B*. We take the coordinate system such that the center of each cell is at x = n, with the *A* and *B* subcells located at $x = n - \frac{1}{4}$ and $x = n + \frac{1}{4}$, respectively. We insert a photon at some initial site and then let it undergo a quantum walk. We assume the insertion point is at an initial *A* subsite; corresponding expressions for insertion at *B* are similar. The Hamiltonian can be expressed [26] in the form

$$H(k) = E_k \begin{pmatrix} 0 & e^{i\theta_k - ik/2} \\ e^{-i\theta_k + ik/2} & 0 \end{pmatrix},$$
 (A.3)

where

$$E_k = (v^2 + w^2 + 2vw\cos k)^{1/2}$$
(A.4)

and

$$\theta_k = \tan^{-1} \left(\frac{(\nu - w)}{(\nu + w)} \tan \frac{k}{2} \right). \tag{A.5}$$

The form of equation (A.3) shows clearly the winding of the matrix elements of *H* in the complex plane as the angle θ_k , which is essentially a Berry phase, changes. The eigenvectors, of energies $E_{\pm} = \pm E_k$, are of the form

$$|\pm\rangle = \frac{1}{\sqrt{2}} \begin{pmatrix} 1\\ \pm e^{-i\left(\theta_k - \frac{k}{2}\right)} \end{pmatrix},\tag{A.6}$$

where the two components represent the amplitudes of being in the A and B states.

Given a state initially localized at subsite A of site n_0 , the spatial wavefunction at later time t will be of the form [26]

$$\psi_{v,w}(r, t) = \frac{1}{N} \sum_{kn} e^{ik(n-n_0)} \bigg(\phi \bigg(r - n + \frac{1}{4} \bigg) \cos(E_k t) + i e^{-i\theta_k + i\frac{k}{2}} \phi \bigg(r - n - \frac{1}{4} \bigg) \sin(E_k t) \bigg).$$
(A.7)

The ϕ functions are the Wannier functions [40–42], defined as the Fourier transform of the momentum-space Bloch wavefunctions

$$\phi_n(r) = \phi(r-n) = \frac{1}{\sqrt{N}} \sum_k e^{-ikn} \psi_k(r).$$
(A.8)

The Wannier functions are tightly localized near the lattice sites (labeled by integer *n*), are orthonormal, and form a complete basis set for the allowed position-space wavefunctions. *t* here is some integer multiple of the time between steps of the discrete quantum walk, t = mT, with m = 0, 1, 2, ... The winding number gives the number of times that the phase ϕ_k winds around the circle as *k* traverses a complete Brillouin zone.



Figure A2. When all of the unit cells of a chain have the same parameter values, a state inserted into the system at any point exhibits standard quantum walk behavior, with ballistic spreading of probability. (a) shows the evolution of the spatial probability distribution versus time, while (b) shows the spatial distribution at a fixed time (after 50 time steps). Here the parameter values used are $\phi_A = -\pi/2$ and $\phi_B = 0$, corresponding to a winding number of 1. The photon starts initially at position 68.

The entangled states in the main text of the paper can be written more explicitly in terms of these wavefunctions. For example, the states of equation (3) can be written

$$\frac{1}{\sqrt{2}}(|\psi_{\nu_{1},w_{1}},H\rangle_{u}|\psi_{\nu_{2},w_{2}},V\rangle_{l}\pm|\psi_{\nu_{2},w_{2}},V\rangle_{l}|\psi_{\nu_{1},w_{1}},H\rangle_{u}),$$
(A.9)

where (v_1, w_1) are a pair of hopping parameters corresponding to winding number 0 and (v_2, w_2) correspond to winding number 1. The wavefunctions of equation (A.7) correspond to the overlap between the state vectors and position eigenstates: $\psi_{v,w}(r, t) = \langle r | \psi_{v,w} \rangle$.

We now numerically display the existence of some of the properties mentioned above for the case of quantum walks on chains of directionally-unbiased multiports. First consider a single chain of such multiports (figure A1(a)) arranged into alternating pairs of diamond graphs, as in section 3. The amplitudes v and w in this case are functions of the adjustable phases ϕ_A and ϕ_B in the two diamond graphs making up each unit cell. The Hamiltonian and the associated vector d(k) can be readily calculated as functions of these phases [28].

When a photon is inserted into the system at a given location, it will begin a quantum walk. After a given number of time steps, the probability distribution can be calculated for the location of the photon. In a classical random walk, the distribution would be expected to have Gaussian form, with a width proportional to \sqrt{N} , where *N* is the number of steps. However, a quantum walk exhibits ballistic behavior, with a distribution that spreads linearly in time. Calculation of the distribution for the unbiased multiport chain shows such ballistic behavior, as seen in figure A2. The bias of the walk (left or right) can be altered by adjusting the values of *v* and *w*.

Consider now two chains lined up end-to-end, and connected at their mutual boundary, as in figure A1(b). Adjust the diamond graph phases so that the winding number is 0 on the left side of the boundary and 1 on the right side. As shown in figures A3 and A4, a state beginning in one region tends not to cross into the other region, even if the available energy levels are the same on both sides. The mismatch of winding numbers leads to a mismatch of eigenstates on the two sides, which in turn reduces propagation from one side to the other. Figure A4 also clearly shows the accumulation of the localized state at the boundary, a feature that is absent from the topologically uniform case of figure A2(b). The peak that accumulates at the boundary remains fixed at that location for all time.

Unlike the protection of the boundary states, the reduced level of bulk state transitions is partial and depends continuously on the hopping parameters. The preservation of the bulk state is 'topologically-assisted' in the sense that it occurs because of reflection at the boundary between topological phases, but it not of a topological





nature in itself, since the transmission and reflection coefficients remain continuous functions of the hopping parameters within each topological sector. The reflection at the boundary is not total: a small transmission amplitude into the second region exists, but it can be arranged to be negligible. For the specific case where the two hopping amplitudes v and w interchange the values when the boundary is crossed, $v \leftrightarrow w$, it is shown analytically in [26] that the degree of leakage across the boundary depends on the difference |v - w| between the two hopping amplitudes. The transmission amplitude is 100% when v = w, but drops toward zero as $|v - w| \rightarrow 1$. So the leakage into the second region can be made as small as desired by taking |v - w| sufficiently large. Qualitatively, the principle reason for this is that the change in topology of the Hamiltonian forces a sudden, discontinuous change in the eigenstates on the two sides of the boundary, making it hard for the rightward-propagating solutions on the two sides to be consistently patched together: the net result is an increased likelihood of reflection.

Reflection at points where there are sudden changes in the dynamics are a very general occurrence. Not only do they occur at points of sudden potential energy change, as described in every quantum mechanics text, and at







points of topological phase change as considered here (see also [43] for an alternative approach), but something very similar happens at many other types of sudden inhomogeneities, including boundaries between regions governed by nonrelativistic (Schrödinger) and relativistic (Dirac) dynamics [44–46].

Finally, consider a disturbance to the system. Again suppose a two-chain system with different winding numbers on the two chains, but now we perturb the system. In figure A5, the result is shown when a state of winding number $\nu = 0$ is perturbed after time-step 30. The disturbance consists of altering the phase shifts to those characteristic of winding number $\nu = 1$ for one time step, then returning to the original values on subsequent steps. The disturbance is therefore localized in time. Normally, such a sudden jolt to the system would create a scattered state capable of propagating away to infinity in both directions. But once again, we see negligible propagation into the right-hand region, indicating that no scattered state associated with the Hamiltonian of 'wrong' winding number appears. Other types of disturbances (localized in space, rather than time, for example, or with different values for the perturbed phases) lead to similar results.

The plots above demonstrate the resilience of the bulk state: the state remains largely unaffected by brief disturbances that temporarily flip the winding number of the Hamiltonian, and tends to reflect to avoid entering regions of opposite winding number. Taken together, this indicates a resistance to states that evolve according to one Hamiltonian making transitions to states evolving according to a Hamiltonian of different winding number; since the Hamiltonians are determined by the polarization states, this indicates by extension a reduced rate of polarization flips.

The resistance to transitions between regions of different winding numbers is quantified in [26], where transmission and reflection coefficients at the boundary are calculated. Assuming the simplest case, where the hopping amplitudes are interchanged at the boundary ($v_{\text{left}} = w_{\text{right}}$ and $w_{\text{left}} = v_{\text{right}}$) the transition rate decreases as |v - w| grows. The transition probability per encounter with the boundary can be made to drop

below 10⁻³, for example, by choosing the parameters such that $\left|\frac{v-w}{w}\right| > 0.96$.



Figure A6. In figures (a)–(e), a variety of phase settings are used, all of which lead to different winding numbers on the two sides of site 86. Despite the details of the walk changing with each change in settings, the bound state remains stable and fixed at the boundary. However, when the winding numbers are made equal on the two sides, the bound state abruptly disappears. (All of these figures were computed after 100 time steps.)

The boundary state, of course, is well known to be stable against perturbations of the Hamiltonian; this can be easily demonstrated. In figures A6(a)–(e), a range of different phase settings are applied. As long as the winding numbers remain different on the two sides, the edge state at the boundary (site 86) remains. Everything else about the walk dynamics may vary, but existence of the bound state remains highly stable. Only when the singular point of the Hamiltonian is crossed and the winding number becomes equal on the two sides (figure A6(f)) does the boundary state disappear.

ORCID iDs

David S Simon b https://orcid.org/0000-0003-4716-9244 Shuto Osawa b https://orcid.org/0000-0003-0542-1405

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