PAPER

Quantum simulation of non-trivial topology

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Abstract
We propose several designs to simulate quantum many-body systems in manifolds with a non-trivial topology. The key idea is to create a synthetic lattice combining real-space and internal degrees of freedom via a suitable use of induced hoppings. The simplest example is the conversion of an open spin-ladder into a closed spin-chain with arbitrary boundary conditions. Further exploitation of the idea leads to the conversion of open chains with internal degrees of freedom into artificial tori and Möbius strips of different kinds. We show that in synthetic lattices the Hubbard model on sharp and scalable manifolds with non-Euclidean topologies may be realized. We provide a few examples of the effect that a change of topology can have on quantum systems amenable to simulation, both at the single-particle and at the many-body level.

1. Introduction

The research field of quantum simulation explores, among other goals, the possibility of using well-controlled quantum systems to simulate the behavior of other quantum systems whose dynamics escapes standard theoretical or experimental approaches. As a relevant example, quantum simulators have been used to successfully analyse condensed matter phenomena [1]. Through synthetic gauge fields [2], more ambitious and multidisciplinary problems can be addressed, such as the determination of the phase diagram of (lattice) gauge theories [3–8]. This theoretical progress is supported by vigorous experimental developments with a growing number of platforms available for quantum simulation like cold neutral atoms and molecules [9], trapped ions [10], photonic crystals [11], NV-centers [12], and superconducting qubits [13].

On a different line of research, topological models have attracted great interest as well. Topology is a key feature to understand many physical phenomena, such as the quantum Hall and quantum spin-Hall effects [14], quantization of Dirac monopole charge [15], charge fractionalization and non-perturbative properties of vacua of Yang-Mills theories [16–19], etc. Topology also plays an essential role in engineering novel states of ultracold matter, such as topological insulators [20]. Notably, topological protection has been considered as a resource for quantum computation [21]. Nonetheless, non-trivial topology is not easy to implement in practical systems. For instance, there is no obvious way to manipulate a 2D condensed matter system to be topologically connected as on a higher genus Riemann surface. Experimental limitations are thus an obstacle to analyse the effects of non-trivial topologies on quantum systems.

The reunion of these two topics, namely quantum simulation and topology, is a natural and tantalizing evolution for both sets of ideas. So far, the focus in quantum simulation has been on topological properties emerging in infinite systems due to their dynamics, e.g., in the toric code [22–24] or in periodically driven systems [25–27], and in synthetic quantum Hall [28–40] and quantum spin-Hall [41–45] systems that exhibit edge states when subjected to open boundary conditions. The search for edge states and other topological properties [46–50] includes also theoretical and experimental efforts in understanding Majorana fermions, since they are produced at the boundaries of some quantum systems [51–53]. But, so far, the simulation of systems with non-trivial boundary conditions, with the exception of circle/torus geometry (see theory [54–56],...
and experiments [57–63], and references therein), has been very scantily explored (see also [64, 65], which appeared while this work was in progress).

Geometry and topology have already made their appearance in quantum simulators, specifically in optical lattices with ultracold atoms. Recently, there have been proposals to simulate quantum many-body physics in certain types of curved background spacetimes [66], tailoring the hopping amplitudes of the optical lattice. Moreover, in [67] a protocol was introduced to use the different atomic states as an artificial extra dimension. This latter proposal plays a key role in our approach to the simulation of quantum matter in different topologies. In effect, by managing the internal interactions between the internal states, we will show how to turn an open 1D optical lattice into a system with periodic boundary conditions (PBC), a cylinder, a torus or a Mobius strip. Our proposal can be engineered also using other platforms and/or may be combined with other techniques.

Such as the ones allowing for well-established toroidal compactifications [68–83], or the speckle potentials allowing to simulate in a controlled way disorder [84, 85].

The paper is organized as follows. Section 2 presents the general strategy to simulate non-trivial topology on a quantum system, while the experimental aspects are discussed in section 3. Section 4 is devoted to an analysis of such as the ones allowing for well-established toroidal compactifications [68–83], or the speckle potentials allowing to simulate in a controlled way disorder [84, 85].

The general aim of our work is to build quantum simulators for dynamics in different topologies out of an optical lattice, which naturally have open boundary conditions. In order to illustrate our strategy let us start with the simplest paradigmatic example: simulating quantum dynamics on a ring i.e., a 1D system with PBC. In principle, this can be achieved by embedding it into a plane, bending it into a circumference and creating an effective interaction between the two extremes which is identical to the one in the bulk. Thus, an extra dimension is required, as well as the possibility of bending the system without altering its dynamical properties. Both requirements are difficult to meet. Therefore, we shall explore a different possibility, which amounts to engineering an artificial extra dimension.

For definiteness, let us discuss a bosonic 1D hopping model with $L$ sites whose Hamiltonian is merely kinetic. Let $a_i^+$ create a boson at site $i$. The PBC are obtained by connecting the end points with an extra term

$$H_L = -\sum_{i=1}^{L-1} J a_i^+ a_{i+1} + J a_1^+ a_L + \text{h.c.},$$

where $J$, the closing hopping, should be taken as $J = J$. The problem of simulating an $S^1$ topology is tantamount to generating this closing term which connects both boundaries of the system. There are several generic strategies to create that term:

- embed the system in a plane and bend it until both boundaries touch, thus reducing the boundary term to an ordinary bulk term.
- Induce a long-range hopping through a medium or an intermediate state.
- Use a synthetic dimension.

This work focuses on the last solution. The introduction of an extra dimension through internal degrees of freedom was proposed in [67]. Indeed, an open 1D line of $L$ sites, each endowed with $M$ internal states, can be regarded as an $L \times M$ synthetic 2D lattice, see figure 1. In geometric terms, we can think of the internal states as a fiber opening at each real-space site. The resulting synthetic lattice would be, therefore, a discrete analogue of a fibre bundle. A generic hopping Hamiltonian for this system can be written as

$$H = -\sum_{\sigma, \sigma'} J_{\sigma, \sigma'}^v b_i^{\dagger(\sigma)} b_i^{(\sigma')} + J_{\sigma, \sigma'}^h b_i^{\dagger(\sigma)} b_{i+1}^{(\sigma')} + \text{h.c.},$$

where $J^v$ and $J^h$ are sets of vertical and horizontal hoppings, in the synthetic lattice view. The vertical term allows us to connect any pair of internal states in the same physical site, while the horizontal term allows us to connect any two internal states in physically neighboring sites.

Figure 2 illustrates the process by which we can convert an open spin chain with $M = 2$ states per site into a system with PBC. In equation (2), simply set $J_{\sigma, \sigma'}^h = J \delta_{\sigma, \sigma'}$ and $J_{\sigma, \sigma'}^v$ to be zero in the bulk, but not in the extremes, $i = 1$ or $i = L$, in which case we have a connecting term between the two species: $J_{1,2}^v = J$. If we
introduce \( l_x = 2L \) virtual particle creation operators, \( a_i = b_i^{(1)} \) and \( a_{2L+1-j} = b_j^{(2)}, j = 1, \ldots, L \), the Hamiltonian reads exactly as a 1D-PBC hopping Hamiltonian.

Let us summarize the idea. By inducing appropriate hoppings on the internal degrees of freedom — whether we call them species, spin values, etc. — we can attain effectively higher-dimensional dynamics, giving them a geometric meaning [67]. This higher dimension can be bent and sewn in different ways, as shown in the previous example. The simplest application consists on turning an open chain with two species into a closed one with a single species. It only needs localized control of the transformations between the species at the boundaries of the open system.

As an additional feature, our synthetic approach allows to control the phases of the induced hoppings. This is equivalent to inducing a magnetic field piercing the chain and, via a gauge transformation, to create boundary conditions which interpolate continuously between periodic and anti-periodic ones. In critical 1D spin models a non-trivial magnetic flux can be regarded as a defect in the associated conformal field theory [86, 87].

The 1D-PBC lattice described above is the basic building block for more interesting 2D models. In the next sections we will discuss more exotic boundary conditions, such as Möbius strips.

### 2.1. Assembling cylinder and torus

A cylinder can be understood as a fiber bundle of segments emerging from each point of a circumference. Let us describe how to create cylindrical synthetic lattice (i.e., a ladder with PBC) of size \( 2L_x \times L_y \) from a 1D open chain of \( L_x \) real sites, with \( M = 2L_y \) internal states per site. Let \( a_{ij} \) create a particle at site \((i, j)\) of the cylinder. Its
correspondent in the synthetic lattice will be \( b_{ij}^{\dagger} \) if \( i \leq L_x \) and \( + b_{ij} \) otherwise. An example with \( L_y = 2 \) is shown in figure 3.

The Hamiltonian of a free bosonic system on a cylinder can be written as

\[
H = -J \sum_{i=1}^{L_x} \sum_{j=1}^{L_y} \left( a_{i,j}^\dagger a_{i,j+1} + a_{i,j+1}^\dagger a_{i,j} \right) + \text{h.c.}
\]  

which can be mapped to the form (2). If \( i \neq 1 \) and \( i \neq L_x \), we have

\[
J_{i,\sigma,\sigma'}^h = J \delta_{\sigma,\sigma'}, \\
J_{i,\sigma,\sigma'} = J B_{\sigma,\sigma'},
\]

where \( B_{\sigma,\sigma'} \) is a \( 2L_y \times 2L_x \) bulk Hermitian matrix of internal hoppings implementing motion in the transverse direction of the cylinder

\[
B_{\sigma,\sigma'} = \delta_{\sigma,\sigma'} \pm 2,
\]

i.e., it is always possible to jump between internal states differing by two units. On the extremes, for \( i = 1 \) or \( i = L_x \), we have to add a new term

\[
J_{i,\sigma,\sigma'}^h = J B_{\sigma,\sigma'} + J C_{\sigma,\sigma'},
\]

where \( C_{\sigma,\sigma'} \) is a closing Hermitian matrix which is responsible for sewing the open edges of the cylinder. Since it corresponds to a pile of circumferences, the non-zero entries of those matrix are of the form

\[
- C_{jj} = - C_{j+1,j} = 1.
\]

The geometrical meaning of that closing matrix is that each horizontal line bends on itself, without mixing.

The synthetic cylinder we just described can be easily turned into a torus by changing matrix \( B_{\sigma,\sigma'} \), with the introduction of new non-zero terms \( B_{1,2L_x-1} = B_{2L_y,1} = B_{2L_x,2} = 1 \) which sew together the upper and lower ends of each fiber, see figure 4. This construction makes sense for \( L_y \geq 3 \). In other terms, each fiber becomes a circumference instead of an open segment. However, while the number of layers, \( L_y \), of the cylinder are limited only by the total number of internal species available, in the case of the torus \( L_y \) can be further restricted by the ability of coupling the internal species cyclically (see section 3 for cold atom implementation).

### 2.2. Möbius strip and twisted torus

The analogy of the synthetic lattice and the fiber bundle, with the internal states playing the role of the fiber, can be exploited further. We can glue the fibers opening at different sites in a different way, in order to provide a non-trivial topology to the manifold. For example, by gluing the first and last fibers of a cylinder via a reflection we can turn it into a non-orientable manifold, a Möbius strip.
Let us discuss in detail the construction of the artificial Möbius strip, a $2L_x \times L_y$ ladder with twisted boundary conditions, i.e., site $(2L_x, j)$ is connected to site $(1, L_y + 1 - j)$. The free Hamiltonian reads

$$H = J \left( \sum_{i=1}^{2L_x-1} \sum_{j=1}^{L_y-1} a_{i,j}^\dagger (a_{i+1,j} + a_{i,j+1}) + \sum_{j=1}^{L_y-1} a_{2L_x,j}^\dagger a_{1,L_y+1-j} \right) + \text{h.c.}. \quad (7)$$

The corresponding synthetic lattice Hamiltonian corresponds to the general form (2), with the following choice of hoppings, see figure 5. For sites $i \neq 1$ and $i \neq L_x$, they are the same as for the cylinder, equation (4). For the extremes, one of them should be the same as for the cylinder, say $i = L_x$. But $i = 1$ must be twisted and connect the different values of the transverse coordinate

$$J_{i,x}^{\nu\sigma} = J B_{x,\sigma'} + J C_{x,\sigma'}^M, \quad (8)$$
where $B_{\sigma'}$ is the bulk matrix, given by equation (5) and the Möbius closing matrix, $C M_{\sigma'}$ has non-zero terms which revert the site ordering of the extra dimension, i.e., connects sites $y = j$ with $y = L_y + 1 - j$. Therefore, we get that the non-zero elements of $CM$ have the form $-+ - () C_{jLj} M_2, 2 1$ (and symmetric) and $+− − () C_{jLj} M_2, 2 1 1 y$. See the Ly = 2 case exemplified in figure 5, where at site $i = 1$ the (yellow) hoppings glue the two different circles.

Of course, this scheme presents the handicap that the size of the transverse direction is not scalable, i.e., it is limited by the number of internal species available and by our ability to couple them. But, as we will see in the next sections, already with $L_y = 2$ we can obtain substantial differences between the cylinder and the Möbius strip.

We can combine the schemes for the Möbius strip and the torus in order to build a twisted torus. The real space Hamiltonian corresponds to (7) with the extra term connecting the $y = 1$ and $y = L_y$ values of all fibers: $\sum J_{i} (a_{i,j}^+ a_{i,j'} + \text{h.c.})$. This maps into $\sum J_{i} (b_{i}^{(1)} b_{i}^{(2)} L_y - 1) + b_{i}^{(2)} b_{i}^{(1)}$ for the synthetic lattice Hamiltonian. See figure 6 for an illustration.

More general boundary conditions, which do not correspond to a 2D manifold, are related to the application of a general unitary matrix of hoppings between sites at $y = i L_y$ and $i = 1$, which can be parametrized as

$$\sum_{j,j'} U_{i,j'} a_{i,j}^+ a_{i,j'} + \text{h.c.,}$$

where $U \in U (L_y)$. If $U$ is the identity matrix, we obtain the cylinder. Let us now consider the case $L_y = 2$. The Möbius strip corresponds to the $U = \sigma_x$ case, which has determinant $-1$. Therefore, can not be connected continuously to the identity matrix. On the other hand, one can reach a pseudo-Möbius strip using a rotation of $\pi$, $U_{1,2} = -U_{2,1} = 1$.

### 3. Cold atom implementation

In this section, we show how the artificial topologies described previously can be made concrete, for instance, in a cold atom set-up. The basic features that allow to realize the abstract construction of section 2 in a cold atom system are the following:

- the synthetic lattice is obtained by loading in a 1D (spin-independent) optical lattice the atoms, whose hyperfine states, belonging to a unique or few hyperfine manifolds, provide internal species which form the synthetic dimension;
- the hopping term $J_{\sigma,\sigma'}$ of (2) is the free hopping of atoms in the 1D optical lattice and is naturally spin-independent, $J_{\sigma,\sigma'} = J \delta_{\sigma,\sigma'}$, as assumed in the construction of Hamiltonians (3) and (7) and their periodic completions;
the hopping term \( J'_{\nu,\sigma} \) is induced and tailored by laser and radiofrequency couplings which are local in the real-space picture, i.e., are acting on a single site of the 1D chain.

A similar dictionary can be obtained for other platforms. For instance, in the spirit of [67], a synthetic dimension can be achieved also in photonic crystals as in [88, 89], by changing the connectivity of the lattice.

It is worth to notice that, while the real spatial dimension is virtually ‘unlimited’ (or better scaled up to order \( 10^5 \) lattices sites), the synthetic dimension is limited always by the number of atomic internal states available, which is up to 10 for standard atoms like \( ^{40}\text{K} \) or \( ^{87}\text{Sr} \) [90], but can be up to 20 in \( ^{167}\text{Er} \) [91], if just one hyperfine manifold is taken into account. However, by considering more than one hyperfine manifold simultaneously or ultracold molecules, see e.g., [92], this number can be further increased. A limited synthetic dimension translates into a limited transverse dimension of the artificial topology.

Let us start by discussing how to implement the building block of our construction, a (spinless) periodic chain from a (spinful) open one. In cold atoms, model (2) applied to create PBC can be realized for instance by loading atoms with at least two hyperfine (almost degenerate) ground states \((F \geq \frac{l + 1}{2})\) in a spin-independent quasi-1D optical lattice of \( L \) sites. The free tunneling provides the terms in \( J' \), while the terms in \( J \) can be created using Rabi oscillations between the hyperfine states, induced by Raman lasers focused on sites 1 and \( L \), respectively. Thus, the synthetic approach we are proposing is essentially local, since the different species are physically at the same site. Notice the scalability of the procedure: we can build PBC 1D systems of any size \( 2L \), if we can build an associated open system with \( L \) sites and \( M = 2 \) internal states.

### 3.1. Cylinder and torus

Let us extend the above construction to the simulation of a cylinder by layering many circles together as explained in section 2.1. It is easy to realize that the Hamiltonian of (3) can be implemented with up to two-photon transitions. Indeed, the most direct arrangement of the internal degrees of freedom \( \sigma \) is in terms of hyperfine states within a unique hyperfine manifold \( F \geq \frac{l + 1}{2}, \text{e.g.,} |\sigma\rangle = b^{(\sigma)} |0\rangle = |F, m_F = m + \sigma\rangle \). For this ordering of the spins, the synthetic sewing coupling \( C_{\sigma^r} \) applied at real-space sites \( i = 1 \) and \( i = L \) requires \( \Delta m_F = 1 \), while the synthetic transverse coupling \( B_{\sigma^r} \) requires \( \Delta m_F = 2 \) for any \( L_y \). Thus, the only limitation in \( L_y \) is given by the number of available internal states. It is worth to notice that the coupling \( C_{\sigma^a} \) applies only alternatively, i.e., it connects only odd and even spin values. This implies that the hyperfine states have to be spectroscopically distinguishable, for instance, through a quadratic Zeeman splitting.

The spin arrangement considered above is not the only possible one. Furthermore, two or more (meta)stable hypermanifolds can be considered. Such a construction is particularly favorable in Earth-alkali like atoms like Yb (see e.g., [93–96]) where the optically connected \(^{1S}\) and the \(^{3P}\) may be used. In this case, a convenient arrangement is to place odd (even) \( \sigma \)'s in the first (second) manifold, i.e., \( |2\sigma\rangle = b^{(2\sigma)} |0\rangle = |I, F, m_F = m + \sigma\rangle, \quad (|2\sigma + 1\rangle = b^{(2\sigma + 1)} |0\rangle = |I, F, m_F = m + \sigma\rangle) \). Thus, the Hamiltonian (3) involves in this scheme just \( \Delta m_F = 1 \) transitions. As further discussed below (see section 3.3), the two-manifold construction allows for a richer interaction pattern than single-manifold one. Both schemes are depicted in figure 3.

Let us now turn to the implementation of a torus geometry. As described in section 2.1, further couplings are needed, which connect the top and the bottom circles. In the synthetic-lattice basis, this is equivalent to connecting the last of the odd (even) spins with the first odd (even) one, for any real-space site. Such a construction makes sense when \( L_y \) is at least 3, the case whose implementation is detailed in figure 4. For simplicity, let us focus on the two-manifold construction. Here, the additional coupling requires just \( \Delta m_F = 2 \) and, similarly to the PBC engineered in [35], can be achieved for instance via a 3-photon transition. For generic \( L_y \), the needed transition has \( \Delta m_F = L_y - 1 \).

### 3.2. Möbius and twisted torus

Let us now discuss the cold atom implementation of a Möbius strip. As explained in section 2.2, we can get a Möbius from the cylinder by replacing the synthetic coupling \( C_{\sigma^a} \), at (e.g.) real-space site \( i = 1 \) with the coupling \( C_{\sigma^M} \). This is equivalent to connecting the internal states \( |\sigma = 2l\rangle \) with \( |2\rangle (L_y - 1 - l) \), and the states \( |\sigma = 2l - 1\rangle \) with \( |2\rangle (L_y - l + 1) \), for \( l = 1, \ldots, L_y \). It is immediate to realize that for any arrangement of the internal states as hyperfine states (both in the one- and two-manifold scenarios) this implies that the maximum \( \Delta m_F \) needed to engineer \( C_{\sigma^M} \) scales with \( L_y \). For instance, in the two-manifold scheme with the arrangement for the \( \sigma \)'s described above, the maximal \( \Delta m_F \) is exactly \( L_y \), see figure 5(c). Thus, the feasible transverse dimension of the strip is technically limited, let us say to \( L_y = 4 \), value which requires at least four-photon transitions.

The step to the implementation of a twisted torus is quite easy and requires the addition of a coupling with \( \Delta m_F = L_y - 1 \), as described above.
3.3. Interactions

The constructions we have presented produced the kinetic terms of the Hamiltonians, which are indeed the right part for the connectivity of the model (including boundary conditions) and, thus, for the hoppings. Cold atom implementation provides a natural way of including interactions. In the synthetic-lattice picture, ordinary on-site interactions due to collisions of atoms with different spins appear as long-ranged.

The pattern of such long-range interactions can be partially controlled. For instance, it is potentially very different for in the (i) one-manifold and in the (ii) two-manifold schemes. Interactions between hyperfine states of the same manifold may change a lot from atom to atom, but the non-spin-changing ones are in general all of the same order of magnitude and, for Earth-alkali atoms, they are equal. The spin-changing ones are naturally suppressed (p-wave) and can be enhanced without inducing too high three-body losses by using optical Feshbach resonances [97–99]. An alternative route is given by Raman–induced interactions near s-wave Feshbach resonance [100–102]. Collisions between atoms in different manifolds are not affected by such constraint but are in general lossy.

Let us start discussing in details the case (i) with the assumption of SU (2F + 1)-symmetric interactions

\[ H_I = \frac{U}{\sqrt{2}} \sum \hat{n}_j (\hat{n}_j - 1), \]

where \( \hat{n}_j = \sum_{\sigma=\pm1,2L} b_j^{\sigma\dagger} b_j^{\sigma} \) is the total occupation on site \( j \) of the physical 1D-chain. As the interaction is invariant under reordering of the spins, the final Hubbard model on the synthetic cylinder and the Möbius strip looks the same for any of the arrangements chosen to represent the \( \mathbf{σ} \) in terms of \( m_F \).

Indeed, supposing that we selectively fill only the spin-states needed i.e., 2L, the local occupation at site \( j \) of the chain becomes the sum of local occupations at sites \( r = j \) and \( r' = 2L + 1 - j \) of the synthetic lattice,

\[ \hat{n}_j = \sum_{l=1}^{2L} (a_j^{l\dagger} a_j^l + a_j^{l+1\dagger} a_j^{l+1}), \]

and the Möbius strip looks the same for any of the arrangements chosen to represent the \( \mathbf{σ} \) in terms of \( m_F \).

The situation is quite different in scenario (ii), even under the assumption that the interactions are SU (2F + 1)-invariant in each hyperfine manifold. To be definite let us consider Earth-alkali like atoms and assume that interactions are negligible in each hyperfine manifold with respect to the inter-manifold ones, which we model to be just density–density. The final \( H_I \) term in the synthetic lattice is strongly dependent on the chosen spin arrangement. For instance, we can engineer a model where only the term

\[ \frac{U}{2} \sum_j \hat{N}_j \hat{N}_{2L+1-j}, \]

appears.

4. Topology signatures

Let us discuss possible proof-of-principle experimental signatures of the topology of the underlying manifold showing up in quantum many-body dynamics which are amenable to experimental observation in our synthetic lattices. This section is not intended as a review of the broad subject of topological effects on quantum systems and has no relation to topological quantum field theory [103].

We start by discussing the simplest paradigmatic example of a line with two species which can be designed to mimic a single species Hamiltonian on a circumference, as described in section 2. In order to illustrate this idea in a simple way, let us consider the following Hamiltonian for a double spin chain of length \( L \), with tunable connecting terms at the boundaries:

\[
H = \sum_{i=1}^{L-1} \sigma_{i,1}^x \sigma_{i+1,1}^x + \lambda \sum_{i=1}^{L} \sigma_{i,1}^z \\
+ \sum_{i=1}^{L-1} \sigma_{i,2}^x \sigma_{i+1,2}^x + \lambda \sum_{i=1}^{L} \sigma_{i,2}^z \\
+ \lambda \sigma_{1,1}^z \sigma_{2,2}^z + \lambda \sigma_{L,1}^z \sigma_{L,2}^z,
\]

where \( \sigma_{i,m}^k \) denotes the \( k \)th component of the spin on site \( i \), rung \( m \). The two closing interactions \( J_1 \) and \( J_2 \) are responsible for turning the two open chains into a single one on a circumference. As a consequence, boundary conditions are dictated by these coefficients. A signature of the artificial boundary conditions can be measured by the correlation between spins on different chains at a boundary, namely:

\[
B \equiv \langle \sigma_{1,1}^z \sigma_{L,2}^x \rangle - \langle \sigma_{1,1}^z \rangle \langle \sigma_{L,2}^x \rangle.
\]

Observable \( B \) should be zero for disconnected chains. Its value for fixed \( J_1 = 1 \) and \( J \equiv \lambda \in [-1, 1] \) is shown in figure 7 for \( L = 4 \) and \( \lambda = 1 \). Notice that \( J = 1 \) corresponds to the PBC case, which maximally entangles both chains and gives the highest correlator. For \( J = 0 \), we obtain a single open BC chain. More interestingly, for \( J = -1 \), the twist in the boundary condition induces a perfect cancellation in the correlator. This effect is, indeed, a scalable signature of the non-triviality of the topological effects.

Nonetheless, realistic simulations should model the underlying geometry by tuning the hoppings of fermions or bosons. We shall now address such cases, looking for both single-particle and interacting signatures.
4.1. Single-particle signatures

A natural single-particle playground where we can observe the effect of the topology is to consider synthetic magnetic fluxes, which boil down to hoppings with non-trivial phases. In our synthetic lattice it is very easy to control such phases, in particular to make them linearly dependent with the position on the chain if the synthetic links are induced through Raman lasers.

Let us start with a 1D Hamiltonian with PBC, as in equation (1), either for spinless fermions or bosons, with an arbitrary closing phase \( \phi \mathcal{J} \). Its single-particle spectrum, as a function of \( \phi \pi \in [0, 2\pi] \), is shown in figure 8. Thus, the left and right extremes are PBC, while the center corresponds to anti-periodic ones. Notice that the gap of a fermionic system at half-filling will evolve continuously, presenting a maximum at \( \phi \pi = \frac{\pi}{2} \). Of course, this gap scales as \( -L^{-1} \), but for a fixed value of \( \lambda \) the topological signature can be clearly observed.

The more involved case of a 2-rung ladder is shown in figure 9. The upper panels show the single-particle spectrum for a free Hamiltonian either for spinless fermions or bosons, such as equation (3) with \( L_y = 2 \). The closing link between the two extremes can be chosen to be a generic unitary matrix, as shown in equation (9). In all cases we have selected a one-parameter family of unitary matrices with special properties. Thus, in the left panel of figure 9 it is a rotation of angle \( \phi \).

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**Figure 7.** Two four-spin Ising chains can be turned into a single chain of eight spins by tuning the boundary couplings, as shown in equation (11). The plot shows the correlation of spins of each species at the different boundaries, \( B \), as a function of the coupling \( J^x \in [-1, 1] \), for \( J^y = \lambda = 1 \). Note the cancellation of correlations in the case of an artificially frustrated boundary.

**Figure 8.** Hofstadter-like single-particle spectrum of a 1D system with \( L = 40 \) sites under a continuous change in the boundary conditions, as in equation (1), with \( \mathcal{J} = e^{i\phi} \). The X-axis is labeled by the phase \( \phi \in [0, 2\pi] \), and color corresponds to eigenvalue index.
\[ U_{+1}(\phi) = \begin{pmatrix} \cos \phi & -\sin \phi \\ \sin \phi & \cos \phi \end{pmatrix} \]  

where the +1 stands for the value of the determinant. Thus, for \( \phi = 0 \) we have \( U_{+1}(0) = I \), the identity matrix, which implies cylindrical boundary conditions. Meanwhile, for the central panel we have used a different one-parameter family:

\[ U_{-1}(\phi) = \begin{pmatrix} \sin \phi & \cos \phi \\ \cos \phi & -\sin \phi \end{pmatrix} \]  

Although those transformations are also unitary, they all have determinant \(-1\). For \( \phi = 0 \) we obtain \( U_{-1}(0) = \sigma_z \), which denotes a Möbius strip. Notice that the single-particle spectrum is slightly different in both cases, and thus the energy gap at half-filling, shown in the lower panel of figure 9, constitutes a clear topological signature, even if it reduces as \( L_x \) increases. Remarkably, the Möbius ground state is exactly degenerate, since the gap is exactly zero.

Even more, our synthetic construction allows to smoothly interpolate between the two by considering a \( U \) matrix of the form:

\[ U_{+1 \rightarrow -1}(\phi) = \begin{pmatrix} \cos \phi & -\sin \phi e^{i2\phi} \\ \sin \phi & \cos \phi e^{i2\phi} \end{pmatrix} \]  

this hopping matrix is also unitary, but its determinant is \( e^{2i\phi} \). For \( \phi = 0 \) it is +1, and we have the cylinder, while for \( \phi = \pi/2 \) it becomes \(-1\), and we obtain the Möbius strip. The single-particle spectrum of this \( L_x \times 2 \) ladder is shown in the third panel of figure 9. Notice that the gap vanishes for \( \phi = \pi/2 \), since that point corresponds to a Möbius band.
where $H_K$ is the kinetic Hamiltonian described in equation (2), and $U$ is the strength of the nearest-neighbor interaction. $\mu_i$ is a local chemical potential and $n_i$ is the local particle number. The sum in the second term is over nearest neighbors of a certain adjacency structure, which need not be the same as the one employed for the kinetic term. We take $\mu_i$ to be slightly random, in order to remove exact degeneracy in the ground state. The topology of the underlying lattice are totally encoded in the kinetic Hamiltonian $H_K$, which is affected by a global hopping constant $J$.

Let us start by considering a bosonic system with Hamiltonian (15) and focus on the local particle-number fluctuations in the ground state, $\sigma^2 = \sum_i (\langle n_i^2 \rangle - \langle n_i \rangle^2) / N$, where $N$ is the total particle number. It can be employed to distinguish the different phases. Mean-field calculations cannot distinguish between different topologies, since they are local in character so, a fortiori, it will give the same estimate for $\sigma^2$ for all boundary conditions. Using exact diagonalization, on the other hand, different topologies can be told apart by inspecting the behavior of $\sigma^2$ as a function of $J/U$. For a large $J/U$ the bosons are in a superfluid state with large particle-number fluctuations, since each particle is delocalized over the whole lattice. For small $J/U$ the bosons are localized in a checkerboard pattern and the particle-number fluctuations are small.

We consider different boundary conditions for compact lattices (torus and Klein bottle, which is a Möbius strip with its boundaries glued together) and open lattices (cylinder and Möbius strip) for different sizes.

In figure 10 we plot $\sigma^2$ for the ground state of the Bose–Hubbard model on a torus and on a Klein bottle. The data shows that $\sigma^2$ can tell the different boundary conditions apart in this case for intermediate values of $J/U$. In figure 11 we plot $\sigma^2$ as a function of $J/U$ for the normal strip and the Möbius strip for different strip lengths. As expected, in the limits $J/U \to 0$ and large $J/U$ the ground state has the same boson number fluctuations, which is explained by the fact that in both limits the ground state is a product state in the site basis $|104\rangle$. In the latter limit this is not exactly the case due to finite size effects. The data shows that for intermediate values of $J/U$, where the ground state is entangled, $\sigma^2$ is sensitive to the different boundary conditions. Despite the fact that the dependence of physical quantities on the boundary conditions may be construed as finite-size effect, in a large system with $N$ lattice sites the effects of the different boundary conditions can be made to be as important as for...
small systems by rescaling the links between boundaries as $J \rightarrow J\sqrt{N}$, that is, $J$ times the number of twisted links modulo a factor of order one.

Let us now consider a fermionic system with two species per site and on-site repulsion making up a synthetic lattice described by a Hamiltonian of the form (15):

$$H = H_0 + U \sum_{i=1}^{L_x} n_i^{(1)} n_i^{(2)} + \text{h.c.,}$$

where $n_i^{(j)}$ stands for the occupation operator on site $i$ of the $j$th rung. The topology of the underlying manifold plays a substantial role on the low energy physics of Hamiltonian (16). On a cylinder, the dynamics is characterized by the independent motion of particles on both rungs, while on the Möbius band the ladder presents a crossing between them, where the species transmute. Thus, both rungs become a unique circuit, thus changing dramatically the dynamics.

Some results are shown graphically in figure 12 and $L_x = 6$ and $U = J$. From top to bottom we see panels (a) and (b), which depict the ground state and first excited state for the cylinder, and panels (c) and (d) which show the corresponding Möbius states. The color of the circles represent the density, $\langle n_i \rangle$, while the colored arcs represent the density–density correlator, $\langle n_i n_j \rangle - \langle n_i \rangle \langle n_j \rangle$. The dashed lines are the correlators $(a_i^\dagger a_j)$, normalized to be in $[-1, 1]$ to share the same color code.

Figure 12. Representation of the ground state and first excited state of the fermionic Hubbard model (16) on a cylinder and a Möbius strip of size $L_x = 6$ and $L_y = 2$, with repulsion $U$ only taking place vertically and equal to the hopping rate $J$. The color code is marked at the rightside colorbar. The color of each node represents the expected value of $\langle n_i \rangle$. The color of the curved links represent the density–density correlator, $\langle n_i n_j \rangle - \langle n_i \rangle \langle n_j \rangle$. The dashed lines are the correlators $(a_i^\dagger a_j)$, normalized to be in $[-1, 1]$ to share the same color code.
hopping correlators. The physical picture can be described as follows. The particles move along their horizontal lines in counter-phase, i.e., with highly negative density–density correlator between the two lines. This does not lead to frustration because $L_x$ is even and the lanes never cross.

Panels (c) and (d) show the situation for the Möbius topology. The ground state is degenerate, and both states are depicted there. The local density now shows a checkered pattern, and also the density–density correlators. The vertical hopping correlators show also an interesting pattern, alternating positive and negative values. The physical picture is as follows. The lane crossing induced by the topology frustrates the homogeneous ground state found in cylindrical topology. The two lanes have become one, and the only possibility to reduce vertical repulsion is to freeze the system into a charge-density wave. Particles can not move as fast as they would like to reduce their kinetic energy, which is an analogue of a traffic jam. That is the reason for the lane changing correlators.

This results combines remarkably well with the information represented in figure 12, showing that the Mott transition takes place at different values of the $J/U$ parameter for different topologies, independently of the system size. This effect is related in a non-trivial way to frustration and does not scale with the system size.

5. Conclusions

We have shown that non-trivial topologies can be simulated by a combination of two techniques, namely the use of several species at every spatial degrees of freedom and the generation of couplings among these species only at the boundaries of the system. In other words, species work as an extra dimension that allows for the generation of topological transformations from localized interactions.

In particular we have presented explicit proposals for the realization of the following geometries:

- a circle,
- a cylinder,
- a torus,
- a Möbius strip,
- a twisted torus.

We have discussed different possibilities of experimental realization of the proposed schemes, extending significantly the ideas of [67]. Finally, we have presented several signatures of the underlying lattice topology both on free and interacting systems. These examples involve synthetic gauge fields and synthetic dimension, including:

- a two-species open Ising chain with localized interactions among them can be converted in a double-length single-species chain with a synthetic magnetic field.
- Hofstadter-like spectra can be obtained for a circle, a cylinder and a Möbius strip.
- Hubbard systems of moderate size can be engineered on a torus, a Klein bottle, a cylinder and a Möbius strip.

It is worth noticing that by switching on the hopping terms connecting the sites at the bases of the cylinder among themselves, we can also simulate a lattice with the topology of a two-sphere. In the cold atom construction this amounts to closing off the top and bottom openings of the cylinder, thus creating a compact surface with no holes.

Our findings open paths to further investigations of both free and weakly interacting, as well as strongly correlated systems in optical lattices with non-trivial topology. Combining such lattice geometries with synthetic gauge fields leads to various spectacular effects that are within the reach of current experiments.

Acknowledgments

We acknowledge useful discussions with S Iblisdir, P Massignan and L Tagliacozzo, and financial support from FIS2013-41757-P, FIS2012-33642, ERC AdG OSYRIS, EU IP SIQS, EU STREP EQuaM, and Fundació CELLEX. O B acknowledges support from Fundação para a Ciência e a Tecnologia (Portugal), namely through programmes PTDC/POPH and projects PEst-OE/EGE/UI0491/2013, PEst-OE/EEI/LA0008/2013, UID/EEA/50008/2013, IT/QuSim and CRUP-CPU/CQVibes, partially funded by EU FEDER, and from the EU FP7 projects LANDAUER (GA 318287) and PAPETS (GA 323901).
References

[1] Lewenstein M, Sanpera A and Ahufinger V 2012 Ultracold Atoms in Optical Lattices: Simulating Quantum Many-Body Systems (Oxford: Oxford University Press)
[2] Dalibard J, Gerbier F, Juzeliūnas G and Öhberg P 2011 Rev. Mod. Phys. 83 1523
[3] Zohar E, Cirac J I and Reznik B 2012 Phys. Rev. Lett. 109 125302
[4] Banerjee D, Dalmonte M, Müller M, Rico E, Stebler P, Wiese U J and Zoller P 2012 Phys. Rev. Lett. 109 175302
[5] Tagliacozzo L, Celi A, Zamora A and Lewenstein M 2013 Ann. Phys., NY 330 160
[6] Zohar E, Cirac J I and Reznik B 2013 Phys. Rev. Lett. 110 125304
[7] Banerjee D, Bogli M, Dalmonte M, Rico E, Stebler P, Wiese U J and Zoller P 2013 Phys. Rev. Lett. 110 125303
[8] Tagliacozzo L, Celi A, Orland P, Mitchell M W and Lewenstein M 2013 Nat. Commun. 4 2615
[9] Bloch I, Dalibard J and Nascimbène S 2012 Nat. Phys. 8 267–76
[10] Blatt R and Roos C 2012 Nat. Phys. 8 277–84
[11] Aspuru-Guzik A and Walther P 2012 Nat. Phys. 8 285–91
[12] Cal J, Retzker A, Jeledec F and Plenio M B 2013 Nat. Phys. 9 168–73
[13] Houghton A A, Türeci H E and Koch J 2012 Nat. Phys. 8 292–9
[14] Haldane F D M 1988 Phys. Rev. Lett. 61 2015
[15] Sakurai J J and Napolitano J 2011 Modern Quantum Mechanics (Reading, MA: Addison-Wesley)
[16] Su W, Schrieffer J R and Heeger A J 1979 Phys. Rev. Lett. 42 1698
[17] Jackiw R and Rebbi C 1976 Phys. Rev. D 13 3398
[18] Witten E 1979 Nucl. Phys. B 156 269–83
[19] Veneziano G 1979 Nucl. Phys. B 159 213–24
[20] Hasan M Z and Kane C L 2010 Rev. Mod. Phys. 82 3045
[21] Nayak C, Simon S H, Stern A, Freedman M and Sarma S D 2008 Rev. Mod. Phys. 80 1083
[22] Kitagawa T 2003 Ann. Phys., NY 303 2–30
[23] Weimer H, Müller M, Lesanovsky I, Zoller P and Büchler H P 2010 Nat. Phys. 6 382–8
[24] Barreiro JT, Müller M, Schindler P, Nigg D, Monz T, Chwalla M, Hennrich M, Roos C F, Zoller P and Blatt R 2011 Nature 470 486–91
[25] Kitagawa T, Berg E, Rudner M and Demler 2010 Phys. Rev. B 82 235114
[26] Kitagawa T, Broome M A, Fedrizzi I, Rudner M S, Berg E, Kassel I, Aspuru-Guzik A, Demler E and White A G 2012 Nat. Commun. 3 882
[27] Abdöth J K and Obuse H 2013 Phys. Rev. B 88 121406
[28] Aidalsherburg M, Atala M, Lobse M, Barreiro J, Paredes B and Bloch I 2013 Phys. Rev. Lett. 111 185301
[29] Miyake H, Siviloglou G, Kennedy C, Burton W and Ketterle W 2013 Phys. Rev. Lett. 111 185302
[30] Goldman N, Beugnon J and Gerbier F 2012 Phys. Rev. Lett. 108 255303
[31] Goldman N, Dalibard J, Dauphin A, Gerbier F, Lewenstein M, Zoller P and Spielman I B 2013 Proc. Natl. Inst. Sci. 110 6736–41
[32] Dauphin A and Goldman N 2013 Phys. Rev. Lett. 111 135302
[33] Zhao E, Bray–Ali N, Williams C, Spielman I and Satija I 2011 Phys. Rev. A 84 063629
[34] Hügel D and Paredes B 2014 Phys. Rev. A 89 023619
[35] Celli A, Massignan P, Ruseckas J, Goldman N, Spielman I B, Juzeliūnas G and Lewenstein M 2014 Phys. Rev. Lett. 113 040401
[36] Atala M, Aidalsherburg M, Lobse M, Barreiro J T, Paredes B and Bloch I 2014 Nat. Phys. 10 586–93
[37] Buchhold M, Cocks D and Hofstetter W 2012 Phys. Rev. A 85 036414
[38] Liu X J, Liu X, Wu C and Sinova J 2010 Phys. Rev. A 81 033622
[39] Källi M, Trotszyk S and Parmenekanti A 2012 Phys. Rev. A 86 063632
[40] Burrello M, Fulga I, Alba E, Lepori L and Trombettoni A 2013 Phys. Rev. A 88 053619
[41] Goldman N, Satija I, Nikolic P, Bermudez A, Martin-Delgado M, Lewenstein M and Spielman I 2010 Phys. Rev. Lett. 105 255302
[42] Hauke P et al 2012 Phys. Rev. Lett. 109 145301
[43] Beeler M, Williams R, Jimenez-Garcia K, LeBlanc L, Perry A and Spielman I 2013 Nature 498 201–4
[44] Iotzu G, Messer M, Desbuquois R, Lebrat M, Uehlinger T, Greif D and Esslinger T 2014 Nature 515 237–40
[45] Struck I, Simonet J and Sengstack K 2014 Phys. Rev. A 90 031601
[46] Alba E, Fernandez-Gonzalvo X, Mur–Petit I, Pachos J and Garcia-Ripoll J 2011 Phys. Rev. Lett. 107 235301
[47] Liu X J, Law K, Ng T and Lee P 2013 Phys. Rev. Lett. 111 120402
[48] Wang L, Soluyanov A and Troyer M 2013 Phys. Rev. Lett. 110 166802
[49] Abanin D, Kitagawa T, Bloch I and Demler E 2013 Phys. Rev. Lett. 110 165304
[50] Hauke P, Lewenstein M and Eckardt A 2014 Phys. Rev. Lett. 113 045303
[51] Fu L and Kane C L 2008 Phys. Rev. Lett. 100 096407
[52] Mourik Y, Zau K, Flores S, Plassard S, Bakker E and Kouwenhoven L 2012 Science 336 1005–7
[53] Jiang L, Kitagawa T, Alicea J, Akhmerov A, Pekker D, Refael G, Cirac J I, Demler E, Lukin M D and Zoller P 2011 Phys. Rev. Lett. 106 220402
[54] Kanamoto R, Carr L D and Ueda M 2009 Phys. Rev. A 79 063616
[55] Carr L D, Clark C W and Reinhardt W P 2000 Phys. Rev. A 62 063610
[56] Carr L D, Clark C W and Reinhardt W P 2000 Phys. Rev. A 62 063611
[57] Ramanathan A, Wright K C, Muniz S R, Zelan M, Hill W T III, Lobb C J, Helmersson K, Phillips W D and Campbell G K 2011 Phys. Rev. Lett. 106 130401
[58] Eckel S, Jendrezejewski F, Kumar A, Lobb C and Campbell G 2014 Phys. Rev. X 4 031052
[59] Jendrezejewski F, Eckel S, Murray N, Lanier C, Edwards M, Lobb C J and Campbell G K 2014 Phys. Rev. Lett. 113 045305
[60] Wright K C, Blakestad R B, Lobb C J, Phillips W D and Campbell G K 2013 Phys. Rev. A 88 063633
[61] Murray N, Kyriakos M, Edwards M, Wright K C, Campbell G K and Clark C W 2013 Phys. Rev. A 88 053615
[62] Wright K C, Blakestad R B, Lobb C J, Phillips W D and Campbell G K 2013 Phys. Rev. Lett. 110 025302
[63] Ramanathan A, Wright K C, Muniz S R, Zelan M, Hill W T, Lobb C J, Helmersson K, Phillips W D and Campbell G K 2011 Phys. Rev. Lett. 106 130401
[64] Beugeling W, Quelle A and Smith C M 2014 Phys. Rev. B 89 235112
[65] Quelle A, Beugeling W and Smith C M 2014 Solid State Commun. at press doi:10.1016/j.ssc.2014.10.024
[66] Boada O, Celi A, Latroyer J I and Lewenstein M 2011 New J. Phys. 13 035002
