Tunable Acoustic Valley–Hall Edge States in Reconfigurable Phononic Elastic Waveguides

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We investigate the occurrence of acoustic topological edge states in a 2D phononic elastic waveguide due to a phenomenon that is the acoustic analog of the quantum valley Hall effect. We show that a topological transition takes place between two lattices having broken space-inversion symmetry due to the application of a tunable strain field. This condition leads to the formation of gapless edge states at the domain walls, as further illustrated by the analysis of the bulk-edge correspondence and of the associated topological invariants. Interestingly, topological edge states can also be triggered at the boundary of a single domain, when boundary conditions are properly selected. We also show that the static modulation of the strain field allows us to tune the response of the material between the different supported edge states. Although time-reversal symmetry is still intact in this material system, the edge states are topologically protected when intervalley mixing is either weak or negligible. This characteristic enables selective valley injection, which is achieved via synchronized source strategy.

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I. INTRODUCTION

Topological acoustics has rapidly emerged as a recent and fascinating branch of physical acoustics. Following the groundbreaking research in solid-state physics, which showed the existence of topological states of matter [1,2], researchers have been able to formulate the acoustic analog of selected mechanisms [3-12,12]. During the last decade, several studies have investigated topological materials based on broken space-inversion symmetry (SIS). In electronic materials, SIS was broken using different methodologies including graphenelike lattices with staggered sublattice potentials [13–17], strained graphene [18], and multilayered graphene under externally applied electric fields [19–25]. The effect of the asymmetry is that of opening a gap at the original Dirac cone, associated with the hexagonal lattice structure, in which edge states are supported. These edge states cannot be explained by the quantum Hall effect (QHE) mechanism. In fact, as timereversal symmetry (TRS) is intact, the lattice still possesses a trivial topology within the context of QHE [1,26,27]. However, owing to the large separation of the two valleys in k space, valley-dependent topological invariants can be defined and used to classify the topological states of the different lattices. This approach, usually referred to as quantum valley Hall effect (QVHE), was recently investigated for application to fluidic acoustic waveguides [9,12], as well as for elastic plates with local resonators [28,29].

In this paper, we consider the dispersion and propagation behavior of a topological elastic phononic waveguide assembled from a trusslike unit cell [Fig. 1(a)]. Compared to previous studies [28,29], we take a fully continuum modeling approach which provides a general methodology of analysis and allows us to map the topological behavior all the way back to the massless (or massive, when SIS is broken) Dirac equation. An in-depth study of the occurrence of edge states, either at domain walls or at the lattice boundaries, is also presented. The analysis addresses both the topological significance of the different edge states and their lossless behavior. Despite the weak topological nature of the QVHE, we show that selective valley injection can be obtained by a synchronized source approach which is able to target specific eigenstates. In addition, we show tunable and reconfigurable capabilities of the topological medium. Starting from an initial hexagonal D_{6h} lattice symmetry, the



FIG. 1. Schematic illustration of the lattice geometry. (a) The reference crystal (unstrained) consisting of an aluminum trusslike hexagonal lattice having SIS. The dashed black line indicates the primitive Wigner-Seitz cell. (b),(c) The deformed α and β lattices obtained upon application of internal pressure. In both configurations, SIS is broken. (d) The red and blue arrows show the equivalent forces produced by δp . To improve the visualization, the deformation shown in (b),(c) is magnified by a factor of 5 with respect to the actual deformation produced by $\delta p = 20$ MPa.

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symmetry level can be tuned using an external static actuation which allows real-time tuning of the edge states (i.e., of the corresponding topological band gaps), as well as a complete reconfiguration of the lattice from a regular phononic material to a tunable topological medium, and vice versa. Note that, in this paper, we are mostly concerned with the analysis of the physical behavior of the medium and of its tuning capability, and we do not concentrate on the practical details of the implementation of the tuning mechanism. Nevertheless, we note that the local perturbation could be practically implemented by using, as an example, mechanical or thermal loads. In the following, we assume that an isostatic pressure could be applied on the sidewalls to produce the deformation necessary for the topological phase transition.

The individual trusses forming the lattice are assumed made out of aluminum, having the width w = 0.1a and the thickness t = 0.05a, where a is the lattice constant. The fundamental lattice structure clearly exhibits SIS with inversion axes located at the nodal points. The lattice is then deformed by applying an internal pressure δp that lowers the symmetry to D_{3h} . The equivalent force produced by δp is locally normal to the individual truss element, as shown in Fig. 1(d). The application of either a positive or a negative pressure δp results in two different deformed states of the lattice that, for simplicity, we label as α and β states [Figs. 1(b) and 1(c)], respectively. In these states, the SIS is broken and the states α and β are inverted images of each other; that is, they turn into each other as δp switches sign.

We will use a combination of theoretical and numerical tools to show in the following (1) that the evolution from α to β state (and vice versa) is accompanied by a topological phase transition, and (2) that when a phononic system is obtained by assembling the two phases, an edge state is supported along the domain wall [(DW), i.e., the interface between the α and β states]. We note that these edge states can be created and modulated at the boundary of a single lattice by selecting proper boundary conditions and by tuning the internal pressure.

II. TOPOLOGICAL BAND-STRUCTURE ANALYSIS

A. Phononic band structure

For a unit cell in prestrained conditions, the numerical analysis can be divided into two parts: (1) a nonlinear static analysis to calculate the state of displacement \mathbf{u}_0 , stress \mathbf{s}_0 , and strain $\boldsymbol{\epsilon}_0$ induced by the applied pressure load, and (2) a linear wave-propagation analysis around the prestressed equilibrium state.

In our analysis, the prestressed state is calculated using finite-element analysis and used as input for the linear wave analysis. The wave propagation in the prestressed medium can be studied as a linear small-oscillation problem $\mathbf{u}'e^{i\omega t}$ around the new static equilibrium \mathbf{u}_0 . The total displacement

can then be expressed as $\mathbf{u}(t) = \mathbf{u}_0 + \mathbf{u}' e^{i\omega t}$. Under these conditions the linearized wave equation is given by

$$\rho \frac{\partial^2 \mathbf{u}}{\partial t^2} = -\boldsymbol{\nabla} \cdot \mathbf{P},\tag{1}$$

which is readily rewritten as

$$-\rho\omega^2 \mathbf{u}' = -\boldsymbol{\nabla} \cdot \mathbf{P},\tag{2}$$

where ρ is the mass density, $\mathbf{P} = (\nabla \mathbf{u} + \mathbf{I})\mathbf{s}$ is the first Piola-Kirchhoff stress tensor, \mathbf{s} is the second Piola-Kirchhoff stress tensor with $\mathbf{s} = \mathbf{s}_0 + \mathbf{C} : (\mathbf{\epsilon} - \mathbf{\epsilon}_0)$, where \mathbf{C} is a fourth-order elasticity tensor, \mathbf{s}_0 and $\mathbf{\epsilon}_0$ are the initial stress and strain, respectively, and the differential operator ∇ is taken with respect to the material frame. Solving the \mathbf{k} -dependent Bloch eigenvalue problem yields the phononic band structure and the eigenstates. The above described model is assembled and solved using the commercial finite element software COMSOL MULTIPHYSICS.

The frequency dispersion is normalized by the bulk shear wave speed in aluminum c over the lattice constant a. Note that the current system is effectively a flat waveguide that admits Lamb guided modes [symmetric (S) and antisymmetric (A)], as well as shear horizontal modes. In this paper, we concentrate only on A modes (flexural modes), which are those that are more naturally excited in platelike structures under external excitation. Other modes are filtered out based on the particle-motion polarization of the mode shapes. The value of the pressure perturbation $\delta p = 20$ MPa is chosen in order to achieve large deflection without inducing plastic deformation. Note that, because the lattice is designed based on slender members, it can easily accumulate large deflections while maintaining small strain levels.

Figures 2(a) and 2(b) show the band structure of the flexural modes of the lattice with $\delta p = 0$ (intact SIS) and $\delta p = 20$ MPa (broken SIS), respectively. Note that the two lattices in states α and β have identical band structures [Fig. 2(b)] since they are simply spatially inverted versions of each other and with intact TRS.



FIG. 2. Phononic band structure of (a) the undeformed lattice [shown in Fig. 1(a)], and of (b) the α (or β) lattice [shown in Figs. 1(b) and 1(c)]. An incomplete band gap opens up at the original Dirac point as SIS is broken. The inset in (a) shows a comparison between the numerical dispersion data (the black dots) and the dispersion predicted by the $\mathbf{k} \cdot \mathbf{p}$ method (the red lines) near the Dirac point.



FIG. 3. Effect of the applied perturbation δp on (a) the evolution of the upper and lower bounds of the band gap, and (b) the gap width.

The analysis of the dispersion properties in Fig. 2(a) reveals the existence of a degeneracy at the K point from which locally linear dispersion curves emanate. This dispersion structure, known as the Dirac cone [30], is protected by the lattice configuration. However, when SIS is broken [Fig. 2(b)], the degeneracy is lifted and the initially degenerate modes can separate and give rise to a local band gap. It is the adiabatic evolution of this specific dispersion structure under small perturbations that enables the generation of topological edge modes. We show in this paper that the dispersion near the valleys in either the reference or the perturbed configuration can be mapped to a massless or a massive Dirac equation, respectively. Before proceeding, we should also note that the current configuration produces an incomplete band gap [Fig. 2(b)]. In principle, a complete band gap could be obtained by optimizing the lattice geometric parameters [7]; however, we show later that the valley-dependent design is fairly robust and does not strictly require a full band gap.

As established above, the process of lifting the degeneracy and opening the band gap is connected to the applied pressure perturbation within the cell. Figures 3(a) and 3(b) show the evolution of the gap bounds, as well as of the width of the gap, as a function of the applied pressure δp . An average upward shift of the frequency is observable in the gap bounds as $|\delta p|$ increases. This behavior is directly related to the stress-stiffening effect produced by the geometric nonlinear deformation of the cell. The analysis of the gap width indicates that the gap vanishes and reopens as δp crosses zero, therefore suggesting a topological phase transition in the δp space.

B. Berry curvature and valley Chern number

The evolution of the band gap as a function of δp suggests that a lattice obtained by connecting α and β domains should experience a topological phase transition associated with a vanishing gap occurring exactly at the DW. The topological nature of this transition can be characterized using a topological invariant—that is, the Chern number C_n (module 2π). The parameter C_n is obtained by integrating the Berry curvature $\Omega_n(\mathbf{k}) = \nabla_{\mathbf{k}} \times \langle \mathbf{u}_n(\mathbf{k}) | i \nabla_{\mathbf{k}} | \mathbf{u}_n(\mathbf{k}) \cdot \hat{\mathbf{z}}$ of the *n*th mode throughout the first Brillouin zone. For our system, C_n is expected to be zero due to an odd distribution of the Berry curvature in \mathbf{k} space,



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FIG. 4. (a) The dispersion of the flexural A modes of the α lattice near the **K** valley. (b) The corresponding Berry curvature of the upper mode.

which should be expected given that TRS is preserved [1,26,27]. It follows that these lattices are classified as trivially gapped materials in the context of QHE systems. Nevertheless, for small SIS breaking, the Berry curvature is highly localized at the valleys, and the local integral of the Berry curvature converges quickly to a nonzero quantized value [13,23]. This local integral is often referred to as the valley Chern number C_v of the *n*th band, and it is defined as $2\pi C_v = \int \Omega_n(\mathbf{k}) d^2 \mathbf{k}$, where the integral bounds extend to a local area around the valley. The right-hand side of this equation is also referred to as the topological charge. Previous studies in electronic systems [15] have shown that this quantized value has important implications because it characterizes the bulk-edge correspondence. In fact, the difference between the valley Chern numbers of the upper (or lower) bands of two adjacent lattices indicates the number of gapless edge states expected at the DW.

Numerical integration of the Berry curvature shows that each valley of either the α or the β lattice carries a topological charge of magnitude π . The two lattices clearly exhibit Berry curvatures of opposite signs. It follows that, for the upper mode of the valley **K**, the valley Chern numbers are $-\frac{1}{2}$ for the α lattice and $+\frac{1}{2}$ for the β lattice. The difference $|C_v^{(\alpha)} - C_v^{(\beta)}| = 1$ indicates the existence of a single gapless edge state at the DW between the α and β lattices. Figure 4(a) shows the calculated dispersion surfaces of the flexural modes of the α lattice near the valley **K**. Figure 4(b) shows the Berry curvature corresponding to the upper mode. Equivalently, the lower band carries a Berry curvature of opposite sign.

III. DOMAIN-WALL EDGE STATE

There are two possible configurations of the DWs between the α and β lattices, that is, α above β (+y) or vice versa [see Figs. 5(a) and 5(b)]. These two configurations are not equivalent and are dominated by a mirror



FIG. 5. (a) Lattice structure and (b) schematic illustration of the two possible DW configurations based on the α and β lattices. Each DW allows propagation of an edge state for each valley index. (c) Dispersion curves in a superlattice ribbon containing 37 triangular subcells of α lattices stacked along the *y* direction, with symmetric boundary conditions. The color bar denotes the *y* position of the centroid of the superlattice weighted by the strain energy density: modes in blue are edge states at DW1, while modes in red indicate edge states at DW2. (d),(e) Full-field simulations of the edge states along a straight and an arbitrarily shaped DW. The red stars labeled "src" in the insets indicate the positions of the sources.

symmetry with respect to either DW1 or DW2. As a direct consequence of the mirror symmetry, the edge states propagating along the DWs can be either symmetric or antisymmetric (with respect to the DW interface).

This symmetry is exploited in order to reduce the size of the finite-element model. Specifically, we build an α -state superlattice ribbon made of 37 triangular subcells stacked along the y direction [see the red part of Fig. 5(a)] and having a length of about 64*a*. Then, either symmetric or antisymmetric boundary conditions along DW1 or DW2 are applied to simulate the presence of the β -state ribbon. The model is solved to get the Bloch eigenvalues and eigenvectors. Results show that the edge state at DW1 is antisymmetric, while at DW2 it is symmetric. Figure 5(c) shows the dispersion relation of such a ribbon structure.

The color bar denotes the y position of the centroid of the superlattice weighted by the strain energy density. Blue indicates edge states at DW1, while red indicates edge states at DW2. The two modes have equal and opposite group velocities but, since they are supported by the two different edges of the lattice, they cannot couple. In addition, if intervalley mixing is neglected (due to the large separation in \mathbf{k} space between the \mathbf{K} and \mathbf{K}' points), these edge states are immune to backscattering. Note that this assumption is well verified in the absence of short-range disorder. We highlight that this dynamic behavior is effectively equivalent to the quantum spin Hall effect in topological insulators [1] if the spin index is replaced by the valley index.

In order to further characterize the edge modes, we perform full-field numerical simulations. Specifically, we simulate the steady-state response of two different DW shapes (i.e., straight and arbitrary) under a point excitation. Low-reflecting boundaries are used all around the model to suppress reflections. In both cases, results show that the edge states are well concentrated near the DWs and are guided along the wall itself [Figs. 5(d) and 5(e)].

IV. SEMIANALYTICAL APPROACH

Although the band structure and the valley Chern number are accurately calculated via a numerical approach, it is still useful to introduce a semianalytical model to further analyze the effects of SIS breaking on the phononic waveguide. Starting from the (undeformed) reference lattice configuration which still preserves SIS, the $\mathbf{k} \cdot \mathbf{p}$ perturbation approach [31] can be used to show that the linear dispersions at the valleys can be mapped to a massless Dirac equation. The method allows us to obtain approximate solutions to the governing equations by using the degenerate modes as the fundamental basis of an expansion process. In this method, the origin of the k axes is shifted to the degeneracy by letting $\mathbf{q} \equiv \mathbf{k} - \mathbf{K}$ (or $\mathbf{k} - \mathbf{K}'$) and $\nu \equiv \omega - \omega_0$. Hence, the problem of determining the dispersion relations ν -q is cast in the form of a 2×2 eigenvalue problem associated with the following Hamiltonian:

$$\mathcal{H}(\mathbf{q}) = v_q \mathbf{q} \cdot \vec{\sigma},\tag{3}$$

where v_g is the group velocity and $\vec{\sigma} \equiv (\sigma_1, \sigma_2, \sigma_3)$ are Pauli matrices. The Hamiltonian maps to the massless Dirac equation associated with locally linear dispersion. By extracting the eigenstates from the previous simulations, we can numerically confirm that the group velocity $v_g = \pm 0.138c$ obtained from the $\mathbf{k} \cdot \mathbf{p}$ method matches well with the tangent slopes of the exact dispersion data obtained from numerical calculations [inset Fig. 2(a)].

When SIS is broken by the application of a pressure perturbation δp , the degeneracy at **K** (or **K'**) is lifted, the modes become nondegenerate, and a gap opens up. It is well known that breaking SIS introduces a σ_3 component into the Hamiltonian that can be expressed as [1,26]

$$\mathcal{H}(\mathbf{q}) = v_q \mathbf{q} \cdot \vec{\sigma} + m\sigma_3. \tag{4}$$

It represents the Hamiltonian of a massive Dirac equation, which we show applies also to our specific lattice structure.

Since the SIS-breaking perturbation is small, it can be expressed as a perturbation of the massless Hamiltonian,

$$\mathcal{H}(\mathbf{q}) = v_a \mathbf{q} \cdot \vec{\sigma} + \vec{m} \cdot \vec{\sigma},\tag{5}$$

where we assume a general perturbation that potentially contains all four of the Pauli-matrix components

 $(\vec{\sigma} = [\sigma_1, \sigma_2, \sigma_3])$ as well as the identity matrix σ_0 . At the **K** (or **K**') point (**q** = **0**), the Hamiltonian reduces to

$$\mathcal{H}(\mathbf{0}) = \vec{m} \cdot \vec{\sigma}.$$
 (6)

Let $\mathbf{u}_{1,2}^d$ be the two degenerate modes of the undeformed lattice and $\mathbf{u}_{1,2}^n$ be the two nondegenerate modes of the deformed lattice at $\mathbf{q} = \mathbf{0}$. These modes are available from the numerical calculation of the band structure. By performing the inner product

$$I_{ij} \equiv \int_{\text{unit cell}} \mathbf{u}_i^d \cdot \mathbf{u}_j^n d^3 \mathbf{r}, \qquad (7)$$

it is seen that matrix I is diagonal (or antidiagonal, depending on the numbering of the modes), which implies that $\mathcal{H}(\mathbf{0}) = \vec{m} \cdot \vec{\sigma}$ is also diagonal. This means that the perturbation term can contain only σ_0 and σ_3 components, at most. Specifically, the splitting of the eigenfrequencies is connected to the σ_3 component, while the average upward shift of the bands (as a function of δp) is connected to the σ_0 component. Note that the σ_0 term does not affect either the eigenvectors or the topological properties; hence, it can be omitted in the Hamiltonian. The above arguments confirm that the Bloch eigenvalue problem of the SIS-broken lattice can be mapped to the massive Dirac equation [Eq. (4)].

The bulk dispersion is then given by $\nu(\mathbf{q}) = \pm \sqrt{|v_g \mathbf{q}|^2 + m^2}$, with a gap equal to 2|m|. From the numerical results in Fig. 3(b), we find that |m| = 0.000795 MPa⁻¹ δp , where δp is expressed in megapascals.

The corresponding Berry curvature, associated with the evolution of the eigenvectors in \mathbf{k} space, has a distribution sharply peaked at the two Dirac points,

$$\Omega(\mathbf{q}) = \frac{1}{2} \tau m v_g^2 (|\mathbf{q}|^2 v_g^2 + m^2)^{-(3/2)}, \qquad (8)$$

where $\tau = \pm$ allows simple labeling of the two valleys **K** and **K'**. Integrating Eq. (8) gives the valley Chern number $C_v = \frac{1}{2}\tau \text{sgn}(m)$. Under TRS, $m_{\mathbf{K}} = m_{\mathbf{K}'}$ for either a lattice α or β , and $m^{(\alpha)} = m^{(\beta)}$ at the same valley. Thus, $|C_v^{(\alpha)} - C_v^{(\beta)}| = 1$ is always true. Hence, we can conclude that there is one gapless edge state living on the DW connecting the two lattices α and β .

V. EDGE STATES AT EXTERNAL BOUNDARIES

The topological phase transition illustrated above can be obtained also at the boundary of an individual lattice when proper boundary conditions are selected. The two specific boundary conditions considered in this paper are traction-free and fixed. Each boundary condition can be applied on the top (+y) and bottom (-y) boundaries of the ribbon (in either the α or the β state), hence resulting in a total of eight



FIG. 6. (a)–(d) Schematic of the four different boundary configurations involving fixed and traction-free conditions on the top (+y) or bottom (-y) boundaries of the α or β ribbons. The configurations (a) and (c) yield two edge states associated with the dispersion shown in (e). The two boundary conditions in (b) and (d) do not yield topological edge states, as shown in (f).

possible configurations [Figs. 6(a)-6(d)). However, since the α and β lattices are inverted images of each other, a given boundary condition on the edge of an α ribbon corresponds to the inverted image of the same boundary condition on the opposite edge of a β ribbon, therefore yielding only four different edge dispersions [see, for example, the pairs displayed in Figs. 6(a) and 6(c) and Figs. 6(b) and 6(d)]. Note that, although they have the same edge dispersion, the valley indexes interchange.

Not all four groups of the boundary conditions yield edge modes. Only the two edges of the ribbon β in Fig. 6(a) [or the edges of ribbon α in Fig. 6(c)] have propagating edge states, as shown in Fig. 6(e). The other two groups [Figs. 6(b) and 6(d)] do not support edge states [Fig. 6(f)]. Note that, in both cases [Figs. 6(e) and 6(f)], there are localized modes near the free edge whose dispersion curves run along the bulk bands corresponding to the faster flexural branch. These modes have no topological significance and they are effectively surfacelike waves with depth of the same order of the wavelength, which will be discussed in detail in a later section. These results can be explained by identifying the similarity between the dynamic behavior imposed by the boundary conditions and the DWs, respectively.

Recall that DW1 supports an antisymmetric edge state. Antisymmetry requires $\mathbf{u} \cdot \mathbf{t} = \mathbf{0}$ on the interface, where \mathbf{t} is any tangent vector to the interface, therefore having the direction of either $\hat{\mathbf{x}}$ or $\hat{\mathbf{z}}$. To satisfy the antisymmetry condition, u_x and u_z must be zero, while the u_y component is nonzero. The *A* modes under consideration in this paper exhibit particle displacement mostly dominated by the u_z component. When a fixed boundary condition is imposed along the edge, the particle displacement \mathbf{u} is set to zero, therefore resembling the behavior of the antisymmetric

FIG. 7. Full-field numerical simulations showing that (a) edge states can be excited on the fixed edge on the top of an α ribbon, while (b) they cannot be excited at the same position in a β ribbon.

interface. On the other hand, DW2 supports symmetric edge states. At the interface, we have $\mathbf{u} \cdot \hat{\mathbf{y}} = \mathbf{0}$, with no restrictions on u_z . Similarly to the discussion above, the traction-free boundary condition will allow u_z , therefore behaving more closely to a symmetric interface.

As a result, each of the fixed boundaries in Figs. 6(a) and 6(c) (note that this figure is equivalent to Fig. 5, reported here for convenience) is dynamically equivalent to half of DW1. This boundary configuration supports edge states similar to those observed along DW1, but without connecting the two continuous bulk bands [i.e., the red modes in Fig. 6(e)]. Similarly, each of the free boundaries in Figs. 6(a) and 6(c) supports an edge state that does not cross the band gap. On the contrary, the other two groups of boundary conditions [Figs. 6(b) and 6(d)] break the similarity with the DWs; therefore, the edge states are strongly suppressed.

Full-field numerical simulations confirm this behavior and show that edge states can be excited on the fixed edge on the top of an α ribbon, but they cannot be excited under equivalent conditions in a β ribbon [see Figs. 7(a) and 7(b)]. These two conditions correspond to the fixed edges in Figs. 6(c) and 6(b), respectively.

VI. LOSSLESS EDGE STATES

The results above deserve some further discussion to clarify why the edge states can be excited independently from the bulk states existing at the same frequency. Because of the lack of a full band gap, when a point excitation is placed at the domain wall, both bulk and edge states can potentially be excited. However, two factors contribute to make the edge states predominant compared to the bulk states. First, the bulk states amplitude decays as 1/r from the source, while the edge-state amplitude remains constant. Second, since the lattice is inhomogeneous, the mode shapes of either the bulk or the edge states have nonuniform distribution over the unit cell. If a point source is located at the point where the edge state has the largest displacement amplitude, most of the input energy is injected into the edge rather than the bulk state. In Fig. 7(b), if the edge state is suppressed, the longwavelength ripples of the bulk state is visible. In summary, both states exist, but their large separation in k space allows



FIG. 8. The equifrequency contour of the bulk state having the *x* axis aligned with the edge (DW or boundary) at the edge state's frequency. Imposing the conservation of the wave-vector component parallel (k_x) to the interface and providing a bulk state that is faster than the edge state results in scattered waves towards the bulk that are evanescent in the normal (k_y) direction, and the edge state is therefore nonleaky.

selective excitation of the edge states. In a similar way, if a distributed source other than a point source is used, the valley injection can still be achieved by targeting a selected wave number that supports strong edge modes. This result can be achieved by using excitation methods such as comb transducers.

In addition, it can be shown that the edge state propagates without loss. At the frequency of the edge state, we calculate the equifrequency contour (where the x axis is aligned with the edge, the DW, or the boundary) of the faster bulk state which is indicated by the solid curve in Fig. 8. The curve shows an almost circular profile for the propagating bulk state, while the dashed line indicates the imaginary $k_{\text{bulk},v}a$ component corresponding to an arbitrary large $k_x a$ beyond the circle. This is equivalent to a conventional slowness diagram, but different by a constant factor a/ω . When the edge state propagates and induces bulk-wave scattering, the $k_x a$ component must be conserved, that is, $\|\mathbf{k}_{edge}\| = k_{bulk,x}$. Clearly, since the bulk state is faster, there exists no \mathbf{k}_{bulk} on the solid curve satisfying this condition. The scattered bulk state then has a complex wave vector with a purely imaginary k_v component, as illustrated graphically in Fig. 8. This means that the scattered bulk wave is evanescent along the y direction and travels with the same speed as the edge state along the xdirection. As a result, the edge state is lossless and does not necessarily require a bulk band gap, as long as the bulk state is faster (in terms of phase velocity) than the edge state at the same frequency.

VII. TUNING THE BOUNDARY EDGE STATES

In addition to the effect of the boundary conditions, adjusting the pressure of the outermost cells (those close to the boundary) can significantly tune the edge-state dispersion. Yao *et al.* [15] showed that, on a graphene system with staggered sublattice potential, the edge-state



FIG. 9. (a),(b) Edge states at the fixed and free edges when the outer cell pressure is set to $\delta p_o = +2.5|\delta p|$ and $-2.5|\delta p|$, respectively. The edge states can be tuned to bend towards either the top or the bottom bulk bands. (c),(d) Edge states at the fixed and free edges when the outer cell pressure is set to $\delta p_o = -2|\delta p|$ and $+2|\delta p|$, respectively. The edge states become flatbands.

dispersion can be controlled by the on-site energy on the boundary cells. Similarly, in our system, the phononic edge-state dispersion can be manipulated by tuning the pressure of the outermost row of lattice. From the dispersion curve [Fig. 6(e)], we observe that the edge state develops from the bulk band near the valley and, as k_r increases, the edge state localizes more sharply on the edge. At $k_x a = \pi$, the energy is mostly confined to the outermost row of the lattice; therefore, controlling the edge state requires tuning the outermost cell properties. For our phononic waveguide, a general trend can be established by observing that a positive pressure on the outer cells tends to increase the edge-state frequency at $k_x a = \pi$, while a negative pressure tends to decrease it. This result suggests that the edge-state dispersion can be tailored by controlling the local pressure in the edge cells.

Numerical results in Fig. 9 show that a combination of boundary conditions and local pressure allows us to tune the edge states to a partially gapped band, a gapless band, or a flat mode. As an example, by setting the outer cell pressure to $\delta p_o = +2.5 |\delta p|$ near the fixed edge [see Fig. 6(c)], the dispersion curve of the edge state bends up towards the top bulk band [Fig. 9(a)]. Similarly, by setting the outer cell pressure to $\delta p_o = -2.5 |\delta p|$ near the free edge [see Fig. 6(a)], the dispersion curve bends down towards the bottom bulk band [Fig. 9(b)]. Flatbands can also be obtained [Figs. 9(c) and 9(d)] by applying $\delta p_o = 2|\delta p|$ and $-2|\delta p|$, respectively, in the previous two cases. For the two cases supporting the gapless edge states crossing the gap, the pressure applied on the outer cells has a different sign than the bulk lattice. This situation is somewhat equivalent to introducing a DW on the boundary, and it results again in the occurrence of gapless edge states due to the contrast between bulk topological charges.



FIG. 10. Cross-section view of the mode shapes of a finlike elastic waveguide with thickness *t* and width *b*, at kb = 0 and kb = 157. The fundamental flexural mode develops into a surface wave as $kb \gg 1$.

A. Discussion of the nontopological edge state

These results deserve some additional discussion concerning the dispersion of the traction-free edge [Figs. 6(e), 6(f), 9(b), and 9(d)]. Note that the dispersion curves highlight the existence of a localized mode near the free edge, which develops along the bulk band corresponding to the faster flexural branch. These parts have no topological significance, and they are effectively surfacelike waves with a depth of the same order of wavelength. To clarify this aspect further, consider a finlike elastic waveguide with thickness t and width b. The fin can be thought as a continuum version of our phononic ribbon without the lattice structure. It is well known that, as $kb \gg 1$, the fundamental flexural mode develops into a surface wave (Fig. 10). This evolution is similar to the S_0 and A_0 Lamb modes evolving into Rayleigh surface modes as $kt \to \infty$. Therefore, this edge mode has no topological significance, as confirmed also by the zero Berry curvature associated with the bulk bands. Nevertheless, there is strong coupling between this mode and the topological edge states propagating along the same traction-free edge. To clarify the interaction between these two edge modes, we can plot again the results of Fig. 9(b) with an emphasized nonlinear color scheme in Fig. 11. The gapless topological edge state and the nontopological edge state meet at the upper left of the cone, and there is a strong level repulsion indicating hybridization. When they enter the continuous bulk band of the other, they become strongly leaky due to the coupling with bulk modes.



FIG. 11. Results of Fig. 9(b) plotted using a nonlinear color scheme based on the particle displacement. This plot clarifies the interaction between the two edge modes. The gapless topological edge state and the nontopological edge state meet at the upper left of the **K** point and experience a strong repulsion, which indicates mode hybridization. When both modes enter the continuous bulk band, they become leaky waves due to the coupling with the bulk modes.

VIII. SELECTIVE VALLEY INJECTION AND EXCITATION OF ONE-WAY PROPAGATING EDGE STATES

Finally, we show that, despite the TRS remaining intact, the two valley-dependent edge states can be excited individually, thereby giving rise to one-way propagation. Since the two valley edge modes are largely separated in momentum space, the coupling between them is vanishingly small. This means that accurate *valley injection* can be achieved by using a source having a stronger coupling with one of the two valleys. This selective coupling can be obtained using a set of two point sources having a carefully controlled phase difference. The procedure is explained below.

As discussed and shown in Figs. 5(c) and 9, edge states can have a positive or a negative group velocity near a valley, depending on the DW configuration (or, equivalently, on the boundary conditions). Each of those modes has its time-reversed counterpart defined around the opposite valley (Fig. 12). The edge state associated with the valley $\mathbf{K} = [(4\pi/3a), 0]$ (in reciprocal space) maps to $k_{\rm r}a = -2\pi/3$ in the first Brillouin zone of the superlattice. For simplicity, we denote this state as the **K**-polarized edge mode. Similarly, we designate its TR counterpart at $k_x a =$ $2\pi/3$ as the **K**'-polarized edge mode, as shown in Fig. 12. Note that the term *polarization* is used here not to classify the particle velocity but instead to indicate the coupling with a specific valley. This situation suggests that, although both the polarized modes exist at the same frequency, they can still be selectively excited by a source that is polarized in favor of a specific valley.

Consider a **K**-polarized edge mode ($k_x a = -2\pi/3$). The associated displacement field is a Bloch wave function $\mathbf{u}_K(x) = \hat{\mathbf{u}}_K(x)e^{-2i\pi x/3a}$, where $\hat{\mathbf{u}}_K$ a periodic function of



FIG. 12. Edge states can have a positive or a negative group velocity near a valley depending on either the DW configuration or the external boundary conditions, as the red and the blue curves shown in (a) and (b). Each state has its own time-reversed counterpart defined around the opposite valley. The edge state around $k_x a = -2\pi/3$ is associated with valley **K** and is denoted as the **K**-polarized edge mode. Its TR counterpart around $k_x a = 2\pi/3$ is denoted as the **K**'-polarized edge mode.

period *a*. When selecting two points in physical space on the DW (or on the boundary) such that their positions differ by a distance *a* [for example, *x* and *x* + *a*; see Fig. 13(a)], the Bloch function must satisfy $\mathbf{u}_{\mathbf{K}}(x + a) = \mathbf{u}_{\mathbf{K}}(x)e^{-2i\pi/3}$. In other words, the phase of the particle displacement at *x* + *a* differs from that at *x* by $-\frac{2}{3}\pi$. Similarly, a phase difference of $+\frac{2}{3}\pi$ is obtained for the **K**'-polarized edge mode.

Figure 13 shows a schematic view of both the point forces and the displacement vectors in a complex plane, where the single arrows represent the forces and the double arrows represent the particle displacements.

Assume a harmonic point force applied at position x. Without loss of generality, we can use this point as reference for the phase calculation; that is, we assume that the force f(x) is in phase with the displacement u(x) at position x [the red arrows in Fig. 13(a)]. In this case, the force $\mathbf{f}(x)$ contributes an equal amount of energy per period $\int_{T} \mathbf{f}(x) \cdot \mathbf{u}_{\mathbf{K}}(x) dt$ [or, equivalently, $\int_{T} \mathbf{f}(x) \cdot \mathbf{u}_{\mathbf{K}'}(x) dt$] to both the K- and K'-polarized modes. Next, we apply another harmonic point force at position x + a with a phase difference designed to target a specific valley injection. As an example, if the phase difference is selected as $-\pi/3$ [the single blue arrow $\mathbf{f}_{\mathbf{K}}(x+a)$ in Fig. 13(c)], the **K**-polarized mode is excited because the force produces an additional 0.5 units of energy on the K-polarized mode. However, this same force produces a negative unit of energy on the \mathbf{K}' -polarized mode, which, ultimately, eliminates the



FIG. 13. Schematic illustration showing the selective valleyinjection procedure. (a) Two point forces are applied on the DW edge at two locations separated by a distance a (equal to the lattice constant). The phase between the force and the displacement at the two points can be schematically visualized in a complex plane. (b) Choosing location x as the reference, we can assume that the force f(x) (the single red arrow) and the corresponding particle displacements $\mathbf{u}_{\mathbf{K}}(x)$ and $\mathbf{u}_{\mathbf{K}'}(x)$ (the double red arrows) are initially in phase. In this case, the force contributes one unit of energy to each mode. (c) The phase of the K- and K'-polarized modes (the blue and yellow double arrows, respectively) and of the harmonic forces at x + a necessary to create pure K- and K'-polarized modes (the blue and yellow single arrows, respectively). The forces produce negative work on the modes with different subscripts, therefore canceling the work done by the force at x and realizing single-mode excitation.



FIG. 14. Full-field numerical simulations showing the ability to trigger a specific unidirectional edge state (around either the **K** or the **K**' point) along either (a),(b) a DW or (c),(d) a fixed edge. The insets show that, by selecting a proper two-point excitation with an appropriate phase difference (matched to the targeted Bloch mode), accurate valley injection can be achieved.

energy produced at x by the first point source. Similarly, a phase difference of $+\pi/3$ [the single yellow arrow $\mathbf{f}_{\mathbf{K}'}(x+a)$ in Fig. 13(c)] creates a pure \mathbf{K}' -polarized mode. In general, we observe that a phase difference of $-2\pi/3$ ($2\pi/3$) is the most efficient for generating a \mathbf{K} -polarized (\mathbf{K}' -polarized) mode since the source is perfectly aligned with the mode polarization, therefore applying two units of energy to the target mode. However, in this case, 0.5 units of energy would be done on the \mathbf{K}' -polarized (\mathbf{K} -polarized) mode as well, hence resulting in a polarization ratio of 0.8. In general, either of the two approaches described above would be effective in triggering a specific edge state; therefore, the preferred excitation strategy could be selected based on the capability of driving a specific mode in a given material configuration.

Full-field numerical results are shown in Figs. 14(a) and 14(b) for the propagation along the DW and in Figs. 14(c) and 14(d) for the fixed boundary. Both cases are obtained using a phase difference of $\pm 2\pi/3$. The point-force sources are applied in the direction normal to the plate and low reflecting boundaries are used all around the model to suppress reflection.

IX. CONCLUSIONS

In this paper, we present the design of a tunable topological elastic phononic waveguide based on the acoustic analog of the QVHE. The lattice structure exploits SIS breaking while preserving TRS, hence resulting in a weak topological acoustic material. SIS breaking is induced by producing elastic deformations of the original lattice structure, while the topological phase transition is achieved by contrasting ribbons having different topological charges. This configuration could produce clear edge states at the domain-wall interface. In a similar way, the proposed design is capable of achieving topological edge states even in a single lattice configuration where boundary conditions are properly selected to mimic the second topological phase. We show that, in both configurations, the edge states can be tailored efficiently by simply tuning the pressure within the cells. Despite TRS being preserved in the proposed design, selective valley injection can be effectively achieved by using a two-point source excitation strategy, thereby giving rise to well-defined unidirectional edge states. We note that, in the presence of smooth disorder, the intervalley mixing proves to be sufficiently weak to result (almost completely) in the backscattering of immune edge states.

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