Strategically combining four structured domains creates the first ever three-way topological energy-splitter; remarkably, this is only possible using a square, or rectangular, lattice, and not the graphene-like structures more commonly used in valleytronics. To achieve this effect, the two mirror symmetries, present within all fully-symmetric square structures, are broken; this leads to two nondistinct interfaces upon which valley-Hall states reside. These interfaces are related to each other via the time-reversal operator and it is this subtlety that allows us to ignite the third outgoing lead. The geometrical construction of our structured medium allows for the three-way splitter to be adiabatically converted into a wave steerer around sharp bends. Due to the tunability of the energies directionality by geometry, our results have far-reaching implications for applications such as beam-splitters, switches and filters across wave physics.

A fundamental understanding of the manipulation and channeling of wave energy underpins advances in device design in acoustics and optics. For instance, beam-splitters, that split an incident beam of light in two, are extensively used for experiments and devices in quantum computing, astrophysics, relativity theory and other areas of physics. This desire to guide waves, split and redirect them, for broadband frequencies, in a lossless and robust manner, extends well beyond optical devices and into electromagnetism, vibration control and acoustic switches, amongst other fields. Fortunately, the advent of topological insulators in quantum mechanics and their translation into classical systems, has led to waveguides that are more broadband and robust than previous designs and ultimately to robust networks; however, the vast majority of the topological energy-splitters are based upon graphene-like hexagonal structures and hence restricted to a two-way partitioning of energy. Herein we rectify this with an intelligently engineered three-way topological energy-splitter, the geometrical design of which is based upon the square lattice.

Time-reversal symmetric (TRS) topological guides leverage the discrete valley degrees of freedom that arise from degenerate extrema in Fourier space. When constructing topological guides, graphene-like materials are the prime candidate due to their well-defined valleys; these valleys are distinguished by their opposite chirality and related by TRS. The intervalley scattering is heavily suppressed by the large Fourier separation between the two valleys, and each valley becomes an efficient information carrier. These valley modes are attracting growing attention, in part due to their simplicity of construction, leading to the emergent field of valleytronics. The primary benefits of these topologically nontrivial modes over, cavity and topologically trivial interfacial modes, is the additional topological protection afforded by the chiral flux either side of the zero-line modes (ZLMs) and geometrical tunability allowing a bend to be adiabatically converted into a splitter (and vice-versa).

The prevalence of graphene-like structures has primarily limited valleytronic devices to two-way energy-splitters; this is motivated by the conservation of chirality at the valleys' points. A three-way and a four-way partitioning of energy away from a nodal region was shown in and26, respectively, however the latter was dependent upon the tunneling mechanism whilst the former was for a multilayer system. Tunneling would introduce an additional dependency upon the system; namely, the decay length perpendicular to the direction of propagation. Hence, the transmission along the outgoing leads would be heavily contingent upon the location of the mode within the topologically nontrivial band-gap; therefore an alternative method whereby the energy is partitioned away from a well-defined nodal point as opposed to a nodal region is highly desirable. Importantly, this is only possible using a square or rectangular lattice; the three-way energy splitting is dependent upon the equivalence of the interfaces (modulo time-reversal symmetry) that is only achievable using the four-fold symmetric cellular structure. The geometrical tunability, the topological robustness and the three-way partitioning of energy away from a well-defined nodal point are three crucial advantages of the square energy-splitter (see Fig. 1) over competing designs.
We begin in Sec. 2 by explicitly recasting the continuum plate model into the language of quantum mechanics, utilising a Hamiltonian description, while retaining elements of the continuum language to bridge across the quantum and elastic plate communities. Despite us utilising the structured elastic plate equation, our theories are system independent, hence are transposable to other classical systems. We examine a square cellular structure containing only a single mirror symmetry in Sec. 3; we demonstrate how this restricts a medium, comprised of these cells, to solely yield straight valley-Hall guides i.e. the energy cannot be navigated around a bend. Contrastingly, the structure examined in Sec. 4 contains two mirror symmetries which in turn allows for ZLMs to couple around a bend as well as partition three-ways away from a nodal point. A few concluding remarks are drawn together in Sec. 5.

Results

Formulation. The group theoretic and topological concepts foundational to our approach hold irrespective of any specific two-dimensional scalar wave system. We choose to illustrate them here using a structured thin elastic Kirchhoff-Love (K-L) plate\(^4\) for which many results for point scatterers are explicitly available\(^5\); the geometrical ideas themselves carry across to photonics, phononics and plasmonics. Displacement Bloch eigenstates|\(\psi_{n\kappa}\rangle\) satisfy the (non-dimensionalised) K-L equation,

\[
\hat{H}|\psi_{n\kappa}\rangle = \omega_{n\kappa}^2|\psi_{n\kappa}\rangle + F(x)|\psi_{n\kappa}\rangle,
\]

\[
\hat{H} = \int |x\rangle \delta(x-y) \nabla^2_{x\kappa} \langle x\rangle dxdy,
\]

\[
F(x) = \omega_{n\kappa}^2 \sum_{p=1}^{P} \sum_{l=1}^{p} M_{l}^{(p)} \psi_{n\kappa}(x) \delta(x-x_{l}^{(p)}),
\]

for Bloch-wavevector \(\kappa\), \(n\) denoting the eigenmodes and \(\omega_{n\kappa}^2\) the non-dimensionalised frequency; reaction forces at the point constraints, \(F(x)\), introduce dependence upon the direct lattice. The most straightforward constraints, sufficient for our purposes, are point mass-loading \(F(x)\) with the reaction forces proportional to the displacement via an effective impedance coefficient. Here \(l\) labels each elementary cell that repeats periodically to create the infinite physical plate crystal, and each cell contains \(p = 1 \ldots P\) constraints. In an infinite medium the displacements are Bloch eigenfunctions

\[
|\psi_{n\kappa}\rangle = \langle x|\psi_{n\kappa}\rangle = \exp(i\kappa \cdot x) \langle x|u_{n\kappa}\rangle,
\]

where \(|u_{n\kappa}\rangle\) is a periodic eigenstate. The displacements satisfy the following completeness and orthogonality relations:

Figure 1. Three-way topological splitter (\(\omega = 6.5356\)) — (a,b) The canonical splitter geometry of four structured quadrants. Different orientation of scatterers in the orange and blue regions. Source is indicated by circle at left edge. (c) Zoom-in of nodal region cells. (d) Displacement field, illustrating the splitting of energy three ways, panels on right illustrate the opposite chirality at the interfaces.
where $\Delta \kappa' = \kappa' - \kappa$. Due to the completeness of the periodic eigensolution, we can expand $|u_{\kappa_0}|$ in the complete orthogonal basis set $\{|u_{\kappa_0}(x)|\}$ where $\kappa_0$ is fixed,

$$|\psi_{\kappa_0}| = \exp(i\kappa \cdot x)|u_{\kappa_0}|$$

$$= \exp(i\kappa \cdot x) \sum_{m} A_{nm}(\kappa)|u_{\kappa_0}|$$

$$= \exp(i\kappa \cdot x) \sum_{m} A_{nm}(\kappa)|\psi_{\kappa_0}|,$$

where $\Delta \kappa = \kappa - \kappa_0$. After substituting (4), into the governing Eq. (1), we explicitly obtain,

$$\exp(i\Delta \kappa \cdot x) \sum_{m} A_{nm}(\kappa)[(\omega^2_{\kappa_0} - \omega^2_{\kappa_0})^2 + \sum_{N} M_{pq}^{(n)}(\mathbf{x} - \mathbf{x}_N^{(p)})]$$

$$+ 4i\Delta \kappa \cdot \nabla_{\mathbf{x}}^2 + \delta^2(|\Delta \kappa|) |\psi_{\kappa_0}|(\mathbf{x}) = 0.$$  

(5)

up to first-order in $|\Delta \kappa|$. This expansion will be used in the subsequent section, alongside symmetry considerations, to engineer the Dirac cones. A categorisation of planar structures that yield non-symmetry induced Dirac cones was shown in43.

**C$_{2v}$ cellular structure.** In this section we examine the cellular structure shown in Fig. 2(a). Spatially, this structure solely has $\sigma_v$ reflectional symmetry; however, in Fourier space, it has $C_{2v}$ symmetry, due to the presence of time-reversal symmetry. In subsection 3.1, we utilise the expansion (5) and group theoretical considerations to demonstrate how an accidental Dirac cone is engineered. The effects of $\sigma_v$ symmetry breaking, on the bulk bandstructure, are discussed in subsection 3.2. Subsection 3.3 demonstrates how the strategic stacking of geometrically distinct media results in valley-Hall edge states10,15. A ZLM connected to a valley-Hall edge state is shown in subsection 3.4 alongside a justification for why this particular $C_{2v}$ model does not allow for propagation around corners. Since the valley-Hall state is a weak topological state protected solely by symmetry, care must be taken to prohibit backscattering hence knowledge of the long-scale envelope is especially useful for finite length interfaces as it can used to minimise the backscattering. An asymptotic method, more commonly known as high-frequency...
homogenisation (HFH) allows for the characterisation of this long-scale envelope (see Methods); this is applied to a $C_{2v}$ ZLM in subsection 3.4.

**Engineering an accidental Dirac cone.** Band coupling at high-symmetry point for $C_{2v}$ structure. The point group symmetry of the structure, shown in Fig. 2(a), is $\Gamma = C_{2v}$; this is also the point group symmetry at $N, G_N = C_{2v}$ (Table 1). The $C_{2v}$ point group arises from a combination of spatial (reflectional) and time-reversal symmetries; the latter relates $\kappa \rightarrow -\kappa$. The group theoretical arguments used throughout this subsection, are reminiscent of those found in \cite{21} although in our calculations we have applied an actual asymptotic scheme whereby we have judiciously chosen a small parameter with a distinguished limit.

The irreducible representations (IRs) at $N$ are one-dimensional hence there is no symmetry induced degeneracy. Despite this, we shall demonstrate in this subsection how two of the IRs can be tuned such that an accidental degeneracy (that is not symmetry repelled) forms. The four solid bands in Fig. 2(c) (bands numbered 3–6 inclusive) are associated with the eigensolutions, shown in Fig. 3, these match the basis function symmetries of the $C_{2v}$ group (Table 1); hence this indicates that bands 3–6 are symmetry induced and the sequential ordering of them (lowest to highest) is deduced numerically, via the eigensolutions, as:

$$\{A_1, B_1, B_2, A_2\}.$$ 

It is expected, from the dispersion curves (Fig. 2(c)) that the two bands that form the accidental degeneracy, namely $A_1, B_1$, have a strong influence on each other whilst the other two symmetry induced bands, $B_2, A_2$, will have a limited effect on the local curvature or slope of the $A_1, B_1$ bands\cite{44}; the effect, by the bands that lie outside of bands 3–6, on the $A_1, B_1$ bands, is expected to be negligible; to see these points mathematically we initially separate out the eigenket expansion Eq. (4) into two sets of bands; namely, the symmetry set eigensolutions (SSE), bands 3–6, and those that lie outside the SSE,

$$|\psi_{\alpha\beta}\rangle = \exp(i\Delta \kappa \cdot \mathbf{x}) \left( \sum_{j \in \text{SSE}} A_{\alpha j}(\kappa)|\psi_{\beta j}\rangle + \sum_{\alpha \in \text{SSE}} A_{\alpha 0}(\kappa)|\psi_{\alpha 0}\rangle \right).$$

Motivated by the orthogonality condition (3) and the expansion (6), we multiply Eq. (4), by $\psi_{\alpha 0}(x)$ or $\psi_{\beta 0}(x)$ (where $l \in \text{SSE}, \beta \not\in \text{SSE}$) before integrating over the primitive cell to obtain the following two equations,

$$(\omega_{\alpha 0}^2 - \omega_{\beta 0}^2)A_{\alpha l}A_{\beta l} = \sum_j H_{l j}A_{\alpha j} + \sum_{\alpha} H_{\alpha 0}A_{\alpha 0},$$

$$(\omega_{\alpha 0}^2 - \omega_{\beta 0}^2)A_{\alpha \beta} = \sum_j H'_{\beta 0}A_{\alpha j} + \sum_{\alpha} H'_{\alpha 0}A_{\alpha 0}$$

where the $\kappa$ dependence of the weighting coefficients has been dropped; $\Lambda_1, H_{ab}, H'_{ab}$ are explicitly,

$$\Lambda_1 = 1 + \sum_p M^{(p)}_l|\psi_{\beta 0}(x^{(p)})|^2,$$

$$H_{ab} = \Delta \kappa \cdot \langle \psi_{\alpha 0}|4\nabla^2|\psi_{\beta 0}\rangle + O(|\Delta \kappa|^2).$$

| Classes $\rightarrow$ IR | $E$ | $C_2$ | $\sigma_v$ | $\sigma_h$ | Basis functions |
|---|---|---|---|---|---|
| $A_1$ | $+1$ | $+1$ | $+1$ | $+1$ | $x^2, y^2$ |
| $A_2$ | $+1$ | $+1$ | $-1$ | $-1$ | $xy$ |
| $B_1$ | $+1$ | $-1$ | $+1$ | $-1$ | $x, xy^2$ |
| $B_2$ | $+1$ | $-1$ | $-1$ | $+1$ | $y, x^2y$ |

**Table 1.** $C_{2v}$ character table.

**Figure 3.** Eigensolutions, at $N$, for $C_{2v}$ case with $\sigma_v$ symmetry. Panel (a), IR: $B_2$, Basis: $y$. Panel (b), IR: $B_1$, Basis: $x$. Panel (c), IR: $A_1$, Basis: $x^2, y^2$. Panel (d), IR: $A_2$, Basis: $xy$.
\[ H'_{ab} = H_{ab} + \delta_{ab}(\omega^2_{n_a} - \omega^2_{\alpha_0}) \sum_p M^{(p)}(\psi_{\alpha_0}^d X^{(p)}(\psi_{\beta_0}^d X^{(p)}) + O(|\Delta\kappa|^2). \]

(9)

Rearranging the second equation in (7) to,

\[ A_{n_2} = \frac{\sum_I H_I A_{m_I} + \sum_j H_{j}A_{N_j}}{(\omega^2_{n_2} - \omega^2_{\alpha_0})} \]

(10)

and substituting this into the second summation of the first equation gives,

\[ (\omega^2_{n_2} - \omega^2_{\alpha_0})A_{n_2} = \sum_j H_j A_{n_j} + \sum_j A_{n_j} \sum_\alpha \frac{H_{n_j}H_{\alpha_0}}{(\omega^2_{n_2} - \omega^2_{\alpha_0})}, \]

(11)

where we have neglected terms which couple states outside the SSE to other states outside the SSE. If we let \( n = l \in \text{SSE} \) and \( \kappa = \kappa_0 + \Delta\kappa \) then the frequency term on the left-hand side is expanded to yield,

\[ \omega^2_{n_2} = \omega^2_{\alpha_0} + 2\omega_{\alpha_0}\Delta\kappa \cdot \nabla_{\alpha_0}\omega_{\alpha_0} + O(|\Delta\kappa|^2). \]

(12)

Hence, from this expansion it is easy to see that the second summation in (11), that couples states within the SSE to those outside, falls into second-order hence the effective first-order equation is,

\[ \left(2\omega_{\alpha_0}\Delta\kappa A_{n} \right) A_{n_2} = \sum_{j \in \text{SSE}} H_j A_{n_j}, \]

(13)

where \( \Delta\kappa = \omega_{\kappa} - \omega_{\kappa_0} \) and \( l \in \text{SSE} \). Notably, the higher-order corrections, that encompass the coupling between bands within the SSE to those outside, provide the band curvature details away from a locally linear point. In this instance, Eq. (13) is a 4 x 4 matrix eigenvalue problem, where the Hamiltonian, with components \( H_{\alpha \beta} \) is Hermitian. If, for a particular \( \kappa_0 \), the first-order term is zero we would have to proceed to second-order; here additional terms would come from the fourth-ordered derivative, the \( \omega_{\kappa} \), expansion and band coupling between outside SSE and inside SSE bands.

Compatibility relations and band tunability along NX. Bands tend to vary continuously except possibly at accidental degeneracies where modal inversion may occur which in turn leads to a discontinuity of the intersecting surfaces. Hence, the eigenfunctions continuously transform as you progress along a continuous IBZ path of simple eigenvalues. The associated IRs, that describe the transformation properties of the eigenfunctions, themselves smoothly transition into IRs that belong to the point groups along \( \Gamma N \) or NX.

In physical space the cellular structure only has \( \sigma_\alpha \), spatial symmetry, this is equivalent to \( \sigma_\beta \) symmetry in Fourier space, Fig. 4(b). Recall the definition of a point group symmetry, i.e. any symmetry operator \( R \in G_\Gamma \) that satisfies, \( R \kappa = \kappa \mod G \), where \( G \) is a reciprocal lattice basis vector; this implies that \( \kappa \in \Gamma N \) solely has the mirror symmetry operator, \( \sigma_\alpha \) within its point group. Similarly, for a \( \kappa \in \Gamma N \), only the vertical mirror symmetry operator, \( \sigma_\alpha \in C_{2\kappa} \), satisfies the point group criterion. The symmetries of the eigenfunctions, for a \( \kappa \) belonging to either of the paths, NX and \( \Gamma N \), are shown within the basis functions column of the \( \sigma_{\alpha \beta} \) Table 2. If we solely consider the two strongly coupled bands, represented by the IRs \( A_1 \) and \( B_1 \), then the associated eigenfunctions transitional behaviour, away from \( N \), is described by the \( \sigma_{\alpha \beta} \) character table. Due to the continuity of the bands the \( A_1 \), \( B_1 \) IRs belonging to the \( C_{2\kappa} \) table will transform into the IRs of the \( \sigma_{\alpha \beta} \) table, as we move away from \( N \); the relationships between different IRs are more commonly referred to as compatibility relations. Initially, we consider symmetry \( \sigma_\alpha \), the eigenstates at \( N \) and along NX satisfy the following,

\[ \hat{P}_\alpha |\psi_{A_1, B_1}\rangle = \pm |\psi_{A_1, B_1}\rangle, \quad \hat{P}_\alpha |\psi_{A_1, B_1}\rangle = \pm |\psi_{A_1, B_1}\rangle. \]

(14)

Hence, the bands \( (A_1, B_1) \) at \( N \) are compatible with \( (A, B) \) (along NX). Physically, this transition is also evident from the eigenvalues; as \( B_1 \to B \) the eigensolution may also satisfy oddness relative to the \( x \)-axis (see Fig. 3). Similarly, at \( N \) and along \( \Gamma N \), the eigenstates transform under \( \sigma_\alpha \) as,

\[ \hat{P}_\alpha |\psi_{A_1, B_1}\rangle = \pm |\psi_{A_1, B_1}\rangle, \quad \hat{P}_\alpha |\psi_{A_1, B_1}\rangle = \pm |\psi_{A_1, B_1}\rangle. \]

(15)

This implies that the bands \( (A_1, B_1) \) at \( N \) are compatible with \( (A, B) \) (along \( \Gamma N \)). These compatibility relations are summarised pictorially in the unfurled IBZ path (Fig. 4(c)). Importantly, that, in deriving Eq. (13) we have only assumed that \( \kappa_0 \) belongs to a particular symmetry set band (surfaces 3–6) (the band at \( \kappa_0 \) must be continuously connected to the same band at \( N \)). Therefore, the compatibility relations allow us to choose any expansion point along the the path \( \Gamma N X \) where the eigensolution basis set, Eq. (4), transforms accordingly i.e. \( |\psi_A\rangle \to |\psi_A\rangle \).
where parity considerations allows for simplification of the Hermitian matrix; the eigensolutions are evaluated at \( \kappa_0 \). Solving the eigenvalue problem yields the following result,

\[
\omega = \pm \Delta \kappa (|\partial + \partial_\gamma|) = 0,
\]

where parity considerations allows for simplification of the Hermitian matrix; the eigensolutions are evaluated at \( \kappa_0 \). Solving the eigenvalue problem yields the following result,

\[
2\omega_1 \Delta \omega_1 \Lambda_A = \Delta \kappa (|\partial + \partial_\gamma|) = 0,
\]

where the \( \pm \) corresponds to the \( A, B \) bands, respectively. This result implies that the \( A, B \) bands have an identical slope, albeit with opposite gradients; hence, if, at \( N \) an instance can be found where \( \omega_A > \omega_B \), then the bands will invariably cross along the path \( NX \). The parametric variation afforded to us, and encompassed in the variable \( \Lambda_{A,B} \), merely increases or decreases the slope thereby increasing or decreasing the distance between \( N \) and the Dirac point. Note that the Dirac cone occurs along the spatial symmetry path, \( \sigma_v \), of the structure due to the opposite parities of the \( A, B \) bands; band repulsion occurs along the \( \Gamma N \) path thereby resulting in a partial band gap along \( NX \). If \( \omega_B > \omega_A \), then the partial gap along \( \Gamma N \) isolates the Dirac cone along a portion of the IBZ path, \( \Gamma NX \).

The distance between the Dirac and high-symmetry point is highly relevant for the transmission properties of the topological guide stated that the transmission is better for short wave envelopes, as opposed to long wave envelopes. This is due to the fact that the short wavelength waves will experience a higher effective mass which results in a higher transmission probability.

### Table 2. \( \sigma_{vh} \) character table and selected portion of \( C_2v \) character table.

| Classes \( \rightarrow \) IR \( \downarrow \) | \( \sigma_h \) | \( \sigma_v \) | Basis functions |
|---|---|---|---|
| \( A_1 \) | +1 | +1 | \( x^2, y^2, xy \) |
| \( B_1 \) | −1 | +1 | \( x, xy^2 \) |
| Classes \( \rightarrow \) IR \( \downarrow \) | \( \sigma_{vh} \) | Basis functions |
| \( A \) | +1 | +1 | \( x^2, y^2, xy \) |
| \( B \) | +1 | −1 | \( x, y, x^3y, xy^2 \) |

### Figure 4. Physical and Fourier space cells — (a) Cellular structure in physical space. (b) IBZ (shaded region) shown within BZ. Presence of \( \sigma_v \) symmetry in physical space translates into \( \sigma_h \) symmetry in Fourier space, this explains the symmetrical placement of the Dirac cones (blue circles) either side of \( \sigma_h \). Symmetries and IRs, along the paths \( \Gamma \rightarrow N \rightarrow X \), are shown. Panel (d) Effect of parametric tuning on \( A, B \) bands — When \( B \) curve lies about the \( A \) curve, the parameters denoted by \( \Lambda_{A,B} \) can be altered to change the intersection location. For our model, the number of masses, their location (\( \sigma_v \) symmetry preserved) and their mass values can all be varied. Panel (e) Dispersion curves \( C_{2v} \) case (when \( \omega_B > \omega_A \) at \( N \)) — Parameter values same as those in Fig. 2; \( \sigma_v \) symmetry present within physical space cell. In this instance, \( B \) curve lies above \( A \) curve at \( N \), hence there is band crossing along \( NX \).
envelopes, hence, for transmission post the nodal region, it is desirable to increase the distance between the Dirac cone and $N$. The latter is true due to the connection between the bulk and projected bandstructures\(^{46}\); the bulk BZ is reduced to a one-dimensional BZ because the only relevant wavevector component for a straight guide is the one parallel to the ZLM. All wavevectors are projected onto the $\Gamma M$ line in Fourier space, hence if the distance between $N$ and the Dirac cone is increased then the Fourier separation between oppositely propagating modes, along the topological guide, would be increased. A mechanism to do this would be by altering the system parameters; Fig. 4(d,e) and Eq. (16) demonstrate that the slopes of the $A$ and $B$ bands can be increased or decreased by the system parameters thereby altering the position of the band intersection.\(^{47,48}\)

**Breaking $\sigma_v$ symmetry.** From the previous subsection we know that when the $\sigma_v$ symmetry is preserved an accidental Dirac degeneracy can be created; the bands coalescing along $NX$ in Fig. 5(b) are parametrically engineered to do so. An important nuance is that the Dirac points are solely located along the two high-symmetry lines (HSLs), parallel to $\sigma_v$, and not along the perpendicular HSLs (see Fig. 4(b)); this is critical when it comes to energy-splitting. The $\sigma_v$ symmetry is lost in Fourier space when the internal set of inclusions is rotated and this breaks open the Dirac point to create a band-gap, Fig. 5(c). The locally quadratic curves, in the vicinity of the former Dirac cones, are commonly referred to as “valleys” and they carry nonzero valley Chern numbers (Fig. 5(a)) which in turn leads to the generation of valley-Hall edge modes\(^{47,48}\). The valley Chern number ($C_v$) is formally defined as follows,

\[
C_v = \frac{1}{2\pi} \int_S \nabla \times \left( i \langle \psi_{h\sigma} | \nabla \psi_{h\sigma} \rangle \right) d\kappa
\]

\[
= \frac{i}{2\pi} \oint_{\gamma} \langle \psi_{h\sigma} | \nabla \psi_{h\sigma} \rangle \cdot d\mathbf{l}
\]

\[
= \frac{i}{2\pi} \sum_{\gamma} A_{nj}(\kappa) \nabla \cdot A_{nj}(\kappa) = \frac{\varphi}{2\pi},
\]

where the surface and line integrals are computed in the vicinity of the valleys, $\varphi$ denotes the Berry phase and the Kohn-Luttinger coefficients, $A_{nj}(\kappa)$, are derived from a variation of Eq. (13) that contains the perturbation terms\(^{15}\). Both, $C_v$ and $\varphi$ are dependent upon the perturbation strength and hence are not quantized quantities. Despite this, a topological integer, $\text{sgn}(C_v)$, exists that yields a bulk-boundary correspondence valid for specific interfaces\(^{49}\). These interfaces must ensure that regions of opposite $\text{sgn}(C_v)$ are not projected onto the same point in Fourier space; hence for the $C_{2v}$ case, from Fig. 5(a), it is evident that an interface associated with the $\Gamma M$ direction will lead to well-defined topological integers. In the next subsection, we shall show how, the locations of nonzero $C_v$ dictates how the geometrically distinct media are stacked.

**$C_{2v}$ adjoining ribbons.** Attaching two topological media, with opposite $\text{sgn}(C_v)$ yields broadband chiral edge states\(^{16}\). This is achieved by placing one gapped medium, above its $\sigma_v$, reflected twin; in essence, the stacking in Fourier space results in regions of opposite $\text{sgn}(C_v)$ overlaying each other, this local disparity ensures the presence of valley-Hall edge modes. The two distinct orderings of the media create two distinct interfaces, as seen in Fig. 6(a) one of which supports only the even modes and the other only the odd modes. This evenness and oddness of the edge modes is inherited from the even and odd bulk modes, Fig. 3. The gapless curves are a symptom of the topologically nontrivial nature of the edge states; this is akin to the valley-Hall modes seen for the zigzag interface within hexagonal structures.

The simplicity of this construction, the apriori knowledge of how to tessellate the two media to produce these broadband edge states, and the added robustness\(^{26}\) are the main benefits of these topological valley-Hall modes. The additional functionality of having a three-way topological splitter (Fig. 1) comes with a caveat: The Fourier separation between the valleys controls the intervalley scattering and the smaller separation in the square lattice,
Fig. 6(a), vis-a-vis that for graphene-like structures\textsuperscript{26} leads to increased scattering. This can be mitigated as the Fourier separation can be artificially increased by parametrically increasing the distance between the Dirac cone and \(N\) in Fig. 5(b) thereby acting to increase the robustness of the edge states against shorter-range defects. The property of the \(C_{2v}\) case that prohibits propagation around the bend is the absence of well-defined valleys with nonzero \(C_v\), along the vertical HSLs of the BZ, see Fig. 5(a). Hence, there is no arrangement that can be placed to the right of either stacking in Fig. 6(a) to obtain a ZLM perpendicular to the blue-orange interface, Fig. 6(b). The ZLM, Fig. 6(c), has a long-scale periodic envelope that can be captured using an effective medium theory\textsuperscript{50} (see Methods). Knowledge of the long-scale envelope is especially useful for these finite length interfaces as it can used to minimise the backscattering as one has, in effect, a Fabry-Pérot resonator.

To summarise, for this \(C_{2v}\) case, there are ZLMs along straight interfaces, however the energy cannot navigate around a \(\pi/2\) bend because there is no post-bend mode to couple with.

\(C_{4v}\) cellular structure. We now extend the concepts illustrated in Sec. 3 to a cellular structure that possesses \(C_{4v}\) point group symmetry at \(\Gamma\) (see Fig. 7(a)). Due to this structure possessing two perpendicular mirror symmetries, as opposed to one, there exists regions of nonzero \(C_v\) along both edges of the square BZ, subsection 4.1 (Fig. 7(b)); this will be shown to yield propagation around a corner (subsection 4.2) as well as a three-way splitting of energy (subsection 4.3). An extensive comparison between the square structures, discussed within this article, and the earlier valleytronics models based upon graphene-like structures\textsuperscript{16–20,26–39} will be pictorially shown at the end of subsection 4.3.

Breaking \(\sigma_{h\alpha}\) symmetries. The \(C_{4v}\) case, Fig. 7(a), is reminiscent of the \(C_{2v}\) case but now with the addition of \(\sigma_{h\alpha}\) symmetry in physical space. This reflectional symmetry yields additional Dirac cones along, a parallel set of HSLs, perpendicular to those connected with the \(\sigma_v\) symmetry (Fig. 7(b)). This is evident, for the unperturbed \(C_{4v}\) case, in its dispersion curves, Fig. 7(c); note that we have plotted around the \(C_{4v}\) IBZ to clearly illustrate the correspondence between the two sets of dispersion curves, Figs. 4(b) and 7(c). The additional Dirac cone, for the \(C_{4v}\) case, along \(XM\) is due to the additional \(\sigma_{h\alpha}\) reflectional symmetry in physical space. Rotating the inclusion set (Fig. 7(a)) results in the breaking of both \(\sigma_{h\alpha}\) symmetries thereby opening up a band-gap (Fig. 7(d)). Importantly, an identical band-gap is present whether we're plotting along the \(C_{2v}\) or \(C_{4v}\) IBZ's.

**Figure 6.** (a) Interfacial dispersion curves and ZLMs — Top: maroon curve arises when blue medium stacked over orange (left-sided ribbons), whilst navy curves, when orange over blue (right-sided ribbons). Left: even-parity ZLM, \(\omega = 9.00\). Right: odd-parity ZLM, \(\omega = 8.60\). (b) Odd-parity mode along blue over orange interface; importantly, there is no arrangement for the grey cells that ensures a vertical mode. As there are no well-defined valleys of nonzero \(C_v\) along the vertical edges of the BZ in Fig. 5, energy cannot be steered around a \(\pi/2\) bend. (c) Dipolar source placed at leftmost edge, excites odd-parity ZLM. The periodicity of the long-scale envelope is clearly evident; outline around envelope derived from HFH (Methods). Backscattering can be minimised via parametric variation (by decreasing the wavelength of the energy-carrying envelope).
In the subsequent section, we demonstrate how the additional reflectional symmetry enables mode coupling from the pre-bend to post-bend ZLM thereby allowing for energy navigation around a corner.

Propagation around a bend. A crucial property that allows for wave steering for the $C_{4v}$ case is the presence of Dirac cones along both edges of the BZ. Another important property is that, like the $C_{2v}$ case, both, even and odd parity modes exist, however they are now present along the same interface as opposed to different interfaces. The orthogonality of these opposite-parity modes ensures that they do not couple along the same edge. The presence of both parity modes along the same interface (for the $C_{4v}$ case) arises from the relationship between the orange over blue stacking and its reverse (Fig. 8(a,b)). Specifically, it is clearly evident from Fig. 8(a,b) that a right propagating mode for one stacking is a left propagating mode on the other and vice versa. This special property is also what allows for the three-way splitting for the $C_{4v}$ case, Fig. 1, but not for the graphene-like structures.

In the subsequent section, we demonstrate how the additional reflectional symmetry enables mode coupling from the pre-bend to post-bend ZLM thereby allowing for energy navigation around a corner.
conjunction with two-dimensional periodic Bloch conditions. Hence, from Fig. 9, we clearly see that, for a variant of the $C_{4v}$ case, the orange over blue (Fig. 9(a)) or blue over orange (Fig. 9(b)) stacking yields an even-parity decaying mode. More specifically, the orange over blue stacking gives solutions to the right of $\Gamma$, whilst the blue over orange yields solutions to the left of $\Gamma$; this implies that the two stackings host the same mode and are TRS pairs of each other. Note that a parametric variant of the $C_{4v}$ case was used to ensure faster convergence of the Fourier-Hermite spectral method. The local curvature, and thereby the characterisation of the envelope, is obtained for modes in the vicinity of $\Gamma$ (see asymptotics in the bottom panels of Fig. 9). Similar to the earlier $C_{2v}$ structure the edge states that arise are topologically nontrivial and gapless.

Transmission around a bend. The perturbed $C_{4v}$ system has valleys of nonzero $C_v$ along all HSLs of the BZ (Fig. 7(b)). This allows for the strategic arrangement of four structured media such that valleys of opposite $\text{sgn}(C_v)$ overlay each other along, both, horizontal and vertical interfaces, Fig. 10(a). This strategic arrangement necessitates the existence of broadband ZLMs along both of these interfaces simultaneously; therefore, unlike the $C_{2v}$ case, energy is navigable around bends.

The four-cell arrangement shown in Fig. 10(a) encompasses the design of the nodal region (and by extending it outwards, the entire region) for the $\pi/2$ wave steerer and three-way energy-splitter. If the bottom-right inclusion set is rotated clockwise then a wave incident along the leftmost interface will follow the red arrows around the $\pi/2$ bend. The indistinguishable, pre- and post-bend interfaces, ensure that, as the energy traverses the turning point, an even-parity mode will couple into itself. An example of, topological wave steering around a bend, is shown in Fig. 10(b–d). Notably, the $\pi/2$ wave steerers observed within hexagonal structures require coupling between a zigzag mode with an armchair mode. The latter termination hosts topologically trivial edge states due to the overlapping regions of identical $\text{sgn}(C_v)$ resulting in gapped states. Contrastingly, the structure shown in Fig. 10(a) allows for topologically nontrivial $\pi/2$ wave steering.

Similar to the $C_{2v}$ ZLM, the short-scale oscillations are discernible from the long-scale modulation. The importance of this long-scale modulation is numerically elucidated in the subsequent section.

Relevance of envelope to transmission around a bend. The characterisation of the energy-carrying envelope is important, as the tuning of it can lead to higher transmission along finite length interfaces. This principle is elucidated by examining the wave-steering example, Fig. 10(c,f); using finite element integration the intensity of the wave-field in each arm of such a steerer is calculated. The ratio of these intensities is the measure of the transmission of the wave steerser (Fig. 10(f)); this quantity can be seen to oscillate rapidly across the band-gap. This is similar to the behaviour of conventional Fabry-Pérot resonators, where for maximal transmission an integer number of wavelengths must be completely contained in each lead. Thus the length of the interfaces is of importance for optimising the transmission. This effect is clearly seen by the contrast in transmission between Fig. 10(e) and (b–d). Despite the paradigm utilising the valley-Hall topological phase, the robustness and bandwidth of the effect can be further increased, by parametric variation, introducing a TRS-breaking active component, nonlinearity and/or resonators within the nodal region.

**Topological 3-way splitter.** We now move onto the construction of the three-way energy-splitter; rotating the bottom-right inclusion set anti-clockwise, in Fig. 10(a); results in four partitions of geometrically distinct media. A wave incoming, from the leftmost interface, will now follow, both, the red and green arrows thereby splitting
the energy three-ways. The resulting scattering solution, for a monopolar source, is shown in Fig. 1; the topological nature of the modes is demonstrated by the chiral fluxes. The three-way splitter can be tuned to a wave steerer by rotating the cellular structures in lower-right quadrant. For a mode to couple, from one lead to another, the chirality of the modes must match. For the square case this condition is satisfied due to the relationship between the interfaces Fig. 8(a,b); an incident even mode couples to itself along the three exit leads, Figs. 1 and 11(a). The excited for the upper and lower leads (Fig. 1) matches the left lead interface (orange over blue), hence an incident even mode at valley will couple to itself along the upper and lower leads. Importantly, the right-sided interface (blue over orange) is the reverse of the left-sided interface, hence a right propagating mode on the right-sided interface is identical to a left propagating mode on the left-sided interface. Mathematically, the latter implies that \( \kappa_{LR} = -\kappa_L \) where \( \kappa_L \) and \( \kappa_R \) are the wavevectors along the left and right-sided interfaces respectively. Due to the chiral relationship between the interfaces the \( \kappa_L \) and \( \kappa_R \) modes conveniently have matching chirality. This allows for the seamless coupling between the left-sided and right-sided modes however there remains a phase difference to account for. In order for the left-sided mode to couple with the right-sided mode a phase must be acquired as the incident wave passes through the nodal point; this is similar to how an incident wave acquires a phase when it passes through a gratings coupler. This phase difference between the modes implies that \( \kappa_L + \kappa_R = 0 \) thereby resulting in a zero imaginary component along the rightward lead, Fig. 11(b). Despite this phase difference, the chiral relationship between the two interfaces, Fig. 8(a,b), ensures that there is conservation of chirality (a necessary condition for topological mode coupling) throughout this four-region structured domain. The relationship between the interfaces is crucial in allowing the third lead to be triggered. This provides further evidence for why only two-way splitting has thus far been obtained for TRS-breaking topological systems.

Comparing our design with that of a similar hexagonal network, see and Fig. 11(c), we note that the chirality and/or phase velocity mismatch results in energy being redirected solely along the two vertical partitions.
Additionally there is no such relationship between the blue over orange and orange over blue zigzag interfaces, see Fig. 8(c,d). This conservation of chirality and phase velocity, as well as the two distinct interfaces, restricts the hexagonal structures to two-way energy-splitting. A comprehensive pictorial comparison between the $C_{v2}$, $C_{v4}$ cases described herein and the, more common, topologically nontrivial and trivial hexagonal examples described in C26 is shown in Fig. 12.

**Discussion**

We have demonstrated how to geometrically engineer the first-ever broadband three-way energy-splitter. This novel paradigm adds a degree of freedom unavailable to all current designs; namely, the hexagonal valley-Hall energy-splitters and the two-way cavity guide beam-splitters. This design is reliant upon the time-reversal relationship between the interfaces and hence serves as a paradigm for all scalar wave systems: plasmonics, photonics, acoustics, as well as, for vectorial systems such as plane-strain elasticity, surface acoustic waves and Maxwell equation systems. The additional degree of freedom afforded by this three-way energy-splitter, along with latest advancements in topological physics, will inevitably lead to a myriad of highly tunable, broadband and efficient crystalline networks.

**Methods**

**Dispersion curves and scattering solutions.** The dispersion curves throughout the article were obtained using a combination of standard spectral methods as well as the Galerkin method. The standard scheme utilised an adaptation of the plane-wave expansion method to determine the eigenstates. Specifically, a doubly periodic Fourier series expansion is employed by applying Floquet-Bloch conditions on opposite sides of the unit cell, resulting in a generalised eigenvalue problem, that is solved for the non-dimensionalised eigenstates presented throughout.

The dispersive behaviour of the edge states is obtained in a similar manner whereby we consider a stretched unit cell centred on the interface between the two media. Floquet-Bloch conditions are applied to both edges of the ribbon and the height of the ribbon is taken such that it is much greater than the decay length of the localised edge states. The relatively slower decay of the $C_{v4}$ edge mode (Fig. 9) required an alternative method; namely, a Galerkin method where the rate of the decay is built into the expansion of the wavefield through scaled orthonormal Hermite functions.

The scattering solutions for point forcings are obtained through the solution of a system of linear equations by standard methods. When a forcing is applied we utilise a Green’s function approach where the total wavefield is given for $P$ scatterers by

$$\psi_{inc}(x) = \psi(x) + \sum_{p=1}^{P} F_g(\omega_{inc}, |x - x_p|).$$

Using the well-known Green’s function $g(\omega_{inc}, \rho) = (i/2\omega_{inc})[H_0(\omega_{inc}) - H_0(\omega_{inc}, \rho)]$, the unknown reaction terms $F_m (m = 1 \ldots P)$ come from the linear system

$$F_m = M_m\omega_{inc}^2 \psi(\omega_{inc}, |x_m - x_p|).$$

(Fig. 11(c)). Additionally there is no such relationship between the blue over orange and orange over blue zigzag interfaces, see Fig. 8(c,d). This conservation of chirality and phase velocity, as well as the two distinct interfaces, restricts the hexagonal structures to two-way energy-splitting. A comprehensive pictorial comparison between the $C_{v2}$, $C_{v4}$ cases described herein and the, more common, topologically nontrivial and trivial hexagonal examples described in C26 is shown in Fig. 12.

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$$F_m = M_m\omega_{inc}^2 \psi(\omega_{inc}, |x_m - x_p|).$$
### Characterising energy-carrying envelope and relevance to robustness.

The efficacy of transmitting energy around a bend, coupling modes between different leads within a network or even transmission through a straight ZLM is contingent upon the displacement of the mode at the turning, nodal or end point. Knowledge of the long-scale envelope is especially useful for these finite length interfaces as it can used to minimise the backscattering as one has, in effect, a Fabry-Pérot resonator. Examples of the characterisation of the energy-carrying envelope, using high-frequency homogenisation (HFH)\(^5\)\(^0\), for the \(C_\text{v}_2\) and \(C_\text{v}_4\) cases are shown. In addition to this, the interfacial dispersion curves for a variation of the \(C_\text{v}_4\) case are derived using a Fourier-Hermite spectral method\(^5\)\(^1\).

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#### Table

| Broken Symmetries | Hexagon | Square |
|-------------------|---------|--------|
| \(\{\sigma_v, \sigma_h\}\) | \(\sigma_v\), \(\sigma_h\) parity (\(\pi\)) | \(\sigma_v\), \(\sigma_h\) |

#### Berry Curvature Sign (+) (-)

| Edge config. | Topologically nontrivial | Topologically trivial |
|--------------|--------------------------|-----------------------|
| Distinct Interfaces | ![Image](image1.png) | ![Image](image2.png) |

2 distinct interfaces 2 similar interfaces

#### Edge States:

| Band shape | Mode parity |
|------------|-------------|
| \(K'\) \(K\) \(K'\) \(K\) | Even |
| \(K'\) \(K\) \(K'\) \(K\) | Odd |

#### Coupling

**Pros (+)** | **Cons (-)**
--- | ---
+ Topologically nontrivial | + Dirac cones at BZ vertices, hence larger Fourier separation (\(d\)) between modes of opposite chirality: enhanced robustness
+ Large Fourier separation between \(\sigma_v < 0\) and \(\sigma_v > 0\) modes | - Only two-way splitting possible without using tunnelling
+ More than two way splitting possible due to matching phase and group velocity of pre/post-nodal region leads | + Topologically nontrivial
+ Topologically trivial, hence no added chiral flux protection | + Dirac cones along both sets of HSLs hence wave-steering possible
+TRS related interfaces yield a 3-way topologically nontrivial energy-splitter | - Small Fourier separation (\(d'\)) between modes of opposite chirality

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**Figure 12.** Comparison table between different geometrically engineered states. A topologically nontrivial and trivial example are given for the hexagonal structure\(^2\)\(^6\); the two square cases are those included within this article: \(C_\text{v}_2\), \(C_\text{v}_4\).
To fully characterise the long-scale periodic behaviour of topological edge states along a crystal interface we utilise HFH, applying the methodology directly in reciprocal space to further bolster the plane wave expansion (PWE) method that was used to obtain the dispersion curves. This technique is a multiple scale asymptotic method, that (for non-degenerate curves with locally quadratic curvature) results in the following homogenised PDE,

\[ T_0 f_0, x_k x_l - \omega_0^2 f_0 = 0, \]  

(20)

where \( f_0 \) is the long-scale envelope defined on the coordinate system \( (X_k, X_l) \); whilst the \( T_0 \) coefficients fully encapsulate the short-scale behaviour (similar analysis can be carried over to any scalar and vectorial system\(^{51,69-73}\). The tensor coefficients \( T_0 \) are geometrically dependant and, from the simple solution of the homogenised PDE, determine the envelope wavelength for a given frequency. These coefficients are determined entirely from integrated quantities of the wave-field in physical space. To avoid the need for regularisation (higher order corrections) we work in reciprocal space and calculate the \( T_0 \)’s directly, using the PWE method. Our eigenvalue problem is recast into matrix form,

\[ [A(\kappa) - \omega^2 B(\kappa)] W = 0, \]  

(21)

with the matrices \( A, B \) encoding the geometry and forcing of the mass loading.

Expanding in the vicinity of a high symmetry point leads to the following ansatz;

\[ \omega^2 = \omega_0^2 + \varepsilon \omega_1^2 + \varepsilon^2 \omega_2^2 + \mathcal{O}(\varepsilon^3), \]

\[ W = W_0 + \varepsilon W_1 + \varepsilon^2 W_2 + \mathcal{O}(\varepsilon^3), \]

\[ A = A_0 + \varepsilon \sum_i \kappa_i \left( \frac{\partial A}{\partial \kappa_i} \right)_{\kappa_i} + \varepsilon^2 \sum_{i,j} \kappa_i \kappa_j \left( \frac{\partial^2 A}{\partial \kappa_i \partial \kappa_j} \right)_{\kappa_i,\kappa_j} + \mathcal{O}(\varepsilon^3), \]

with a similar expansion for \( B(\kappa) \). Applying suitable solvability conditions and imposing Bloch conditions on the microscale results in the following tensor coefficients \( T_0 \)

\[ \tilde{W}(j) \equiv \frac{\langle W(j) \rangle}{\kappa_j}, \]

\[ = - \frac{[A_0 - \omega_0^2 B_0]}{\omega_0^2 B_0} (A^{(1)} - \omega_0^2 B^{(1)}) W_0 T_0 \]

\[ = \frac{W_2(A^{(2)} - \omega_0^2 B^{(2)}) (W_0 + \frac{1}{2} W_1 (A^{(2)} - \omega_0^2 B^{(2)}) W_0)}{W_2 B_0 W_0}, \]

(22)

where \( \omega_0 \) and \( W_0 \) are the solutions obtained from the PWE method, and \([\ldots]^\dagger\) denotes the pseudoinverse. The explicit characterisation of the envelope is shown in Fig. 6(c).

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Author contributions
M.P.M. conceived the idea, did the majority of the theory, numerical simulations and write-up. G.C. did the numerical simulations shown in Figures 1(c) and 6(c) (envelope component), 9, 10(f). The underlying theory for these components, as well as the illustrative table Fig. 12 were also done by G.C.

Competing interests
The authors declare no competing interests.

Additional information
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