

Geometry and Topology Tango in Ordered and Amorphous Chiral Matter

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1 Abstract

2 Systems as diverse as mechanical structures and photonic metamaterials enjoy
3 a common geometrical feature: a sublattice or chiral symmetry first introduced
4 to characterize electronic insulators. We show how a real-space observable, the
5 chiral polarization, distinguishes chiral insulators from one another and resolve
6 long-standing ambiguities in the very concept of their bulk-boundary corre-
7 spondence. We use it to lay out generic geometrical rules to engineer topo-
8 logically distinct phases, and design zero-energy topological boundary modes
9 in both crystalline and amorphous metamaterials.

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47 1 Introduction

48 A century after the foundations of band theory in solids by Félix Bloch [1], physicists have
 49 discovered new states of electronic matter ranging from insulators to superconductors by
 50 exploiting the topological structure of Bloch theory [2–7]. This topological revolution
 51 has built on two cornerstones: an abstract classification based on symmetries [8–15], and
 52 the practical correspondence between bulk topology and the boundary states measurable
 53 in experiments [2–6, 16–19]. During the past decade, these two generic principles spread
 54 frantically across fields as diverse as photonics, acoustics, or mechanics, leading to design
 55 principles and practical realizations of maximally robust waveguides [20, 21].

56 Among the number of symmetries constraining wave topology, chiral symmetry has
 57 a special status. Out of the three fundamental symmetries of the overarching ten-fold
 58 classification [8–10], it is the only one naturally realized with both quantum and classical
 59 waves. It generically takes the form of a sub-lattice symmetry when waves propagate
 60 in frames composed of two connected lattices A and B , with couplings only between, A
 61 and B sites, see e.g. Fig. 1a. In electronic systems, the archetypal example of a chiral
 62 insulator is provided by the polyacetylene molecule described by the Su-Schrieffer-Hegger
 63 (SSH) model [22]. In mechanics, the Hamiltonian description of bead-and-spring networks
 64 is intrinsically chiral [23–26]: the A sites correspond to the beads, and the B sites to
 65 the springs. In topological photonics and cold atoms chiral wave guides are among the
 66 simplest realizations of topological phases. Over the past decade, the modern theory
 67 of electronic polarization based on Zak phases and non-Abelian Wilson loops [27–30] has
 68 illuminated the intimate relation between crystalline symmetries and the topology of band
 69 structures [11–15]. By contrast, the role of chiral symmetry has been overlooked.

70 In this article, by introducing the concept of chiral polarization we determine the zero-
 71 mode content of interfaces between topologically incompatible crystalline and amorphous
 72 chiral meta(materials)

73 In the bulk, the chiral charge, which measures the imbalance between the number

74 of sites on the sub-frames A and B , predicts the number of zero-energy modes of all
 75 Hamiltonians defined on a given chiral frame. To characterize chiral insulators we define
 76 their chiral polarization $\mathbf{\Pi}$ as the spatial imbalance of the bulk waves on the two sub-
 77 frames. This material property does not rely on any crystalline symmetry and can therefore
 78 be defined locally on disordered and amorphous frames. In crystals, although akin to the
 79 time-reversal polarization of \mathbb{Z}_2 insulators [31], we show that $\mathbf{\Pi}$ is not merely set by the
 80 Bloch-Hamiltonian topology but also by the underlying frame geometry. At boundaries,
 81 we show how $\mathbf{\Pi}$ prescribes the surface chiral charge, and therefore the full zero-energy
 82 edge content of crystalline and amorphous chiral matter. Finally, we propose a series of
 83 practical protocols to experimentally measure the chiral polarization of mechanical, and
 84 photonic chiral metamaterials.

85 2 From chiral charge to chiral polarization and Zak phases

86 Introducing the concepts of chiral charge and polarization, we demonstrate that bulk prop-
 87 erties of chiral matter are determined by an intimate interplay between the frame topology,
 88 the frame geometry and the chiral Zak phases of Bloch Hamiltonians.

90 2.1 Chiral charge and chiral polarization.

91 We consider the propagation of waves in chiral material associated to d -dimensional frames
 92 including two sub-frames A and B . The wave dynamics is defined by a Hamiltonian \mathcal{H} .
 93 By definition, the chiral symmetry translates in the anti-commutation of \mathcal{H} with the chiral
 94 unitary operator $\mathbb{C} = \mathbb{P}^A - \mathbb{P}^B$, where \mathbb{P}^A and \mathbb{P}^B are the two orthogonal projectors on
 95 the sub-frames A and B . Simply put, in the chiral basis where \mathbb{C} is diagonal, \mathcal{H} is block
 96 off-diagonal.

97 In order to determine the relative weight of the wave functions of \mathcal{H} on the two sub-
 98 frames, we introduce the chiral charge

$$\mathcal{M} = \langle \mathbb{C} \rangle, \quad (1)$$

99 where the average is taken over the complete Hilbert space. Using the basis of fully
 100 localized states, we readily find that \mathcal{M} is fully prescribed by the frame topology: the chiral
 101 charge counts the imbalance between the number of A and B sites: $\mathcal{M} = N^A - N^B$. We
 102 can however also evaluate Eq. (1) in the eigenbasis of \mathcal{H} . Indexing by n the eigenenergies
 103 of \mathcal{H} , the eigenstates of the chiral Hamiltonian come by pairs of opposite energies related
 104 by $|-n\rangle = \mathbb{C}|n\rangle$. Chirality therefore implies that the chiral charge is solely determined
 105 by the zero modes of \mathcal{H} as $\mathcal{M} = \sum_n \langle n | \mathbb{C} | n \rangle = \langle 0 | \mathbb{C} | 0 \rangle$. Noting that the $|0\rangle$ states are
 106 eigenstates of the chiral operator with eigenvalue $+1$ when localized on the A sites and
 107 -1 when localized on the B sites, it follows that \mathcal{M} also is an algebraic count the zero
 108 modes of \mathcal{H} :

$$\mathcal{M} = N^A - N^B = \nu^A - \nu^B. \quad (2)$$

109 This equality is the classical result established by Maxwell and Calladine in the context of
 110 structural mechanics [33, 34] and independently discussed by Sutherland in the context of
 111 electron localization [35]. Eq. (2) implies that the spectral properties of \mathcal{H} are constrained
 112 by the frame topology. In particular, frames with a non-vanishing chiral charge impose
 113 *all* chiral Hamiltonian to possess flat bands. This simple prediction is illustrated in Fig. 1
 114 where we show the Lieb and the dice lattices, which are both characterized by a unit
 115 chiral charge per unit cell. All Hamiltonians defined on these lattices are therefore bound

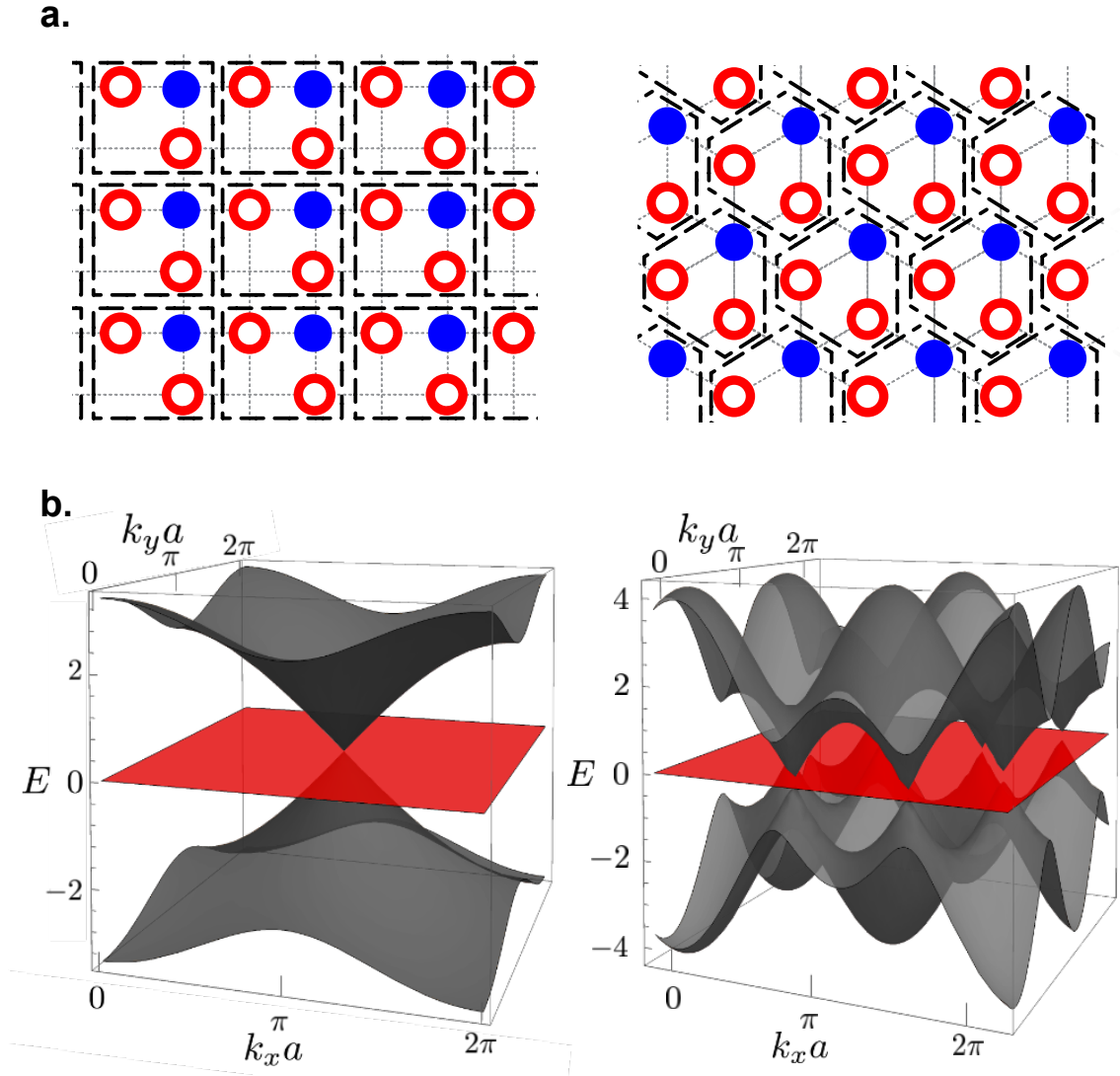


Figure 1: **Lattices with a finite chiral charge.** **a.** The Lieb (left) and dice (right) frames are both characterized by an imbalance between the number N^A and N^B of sites. In both cases the chiral charge per unit cell equals 1. Any Hamiltonian defined on these frames possesses a flat energy band. **b.** Illustration of two band spectra associated to chiral Hamiltonians defined on the Lieb (left) and dice (right) frames. The two band spectra are computed for tight-binding Hamiltonians with nearest neighbour coupling and a hopping parameter set to 1, see e.g. [32].

116 to support at least one flat band, Fig. 1b. No chiral insulators exist on the Lieb and dice
117 lattices.

118 By contrast, in chiral insulators, no zero-energy bulk modes exist and \mathcal{M} must vanish.
119 To probe the relative weight of the wave functions on the two sub-frames, we therefore
120 introduce the chiral polarization vector $\Pi_j = \langle \mathbb{C}x_j \rangle_{E \neq 0}$. As the $|\pm n\rangle$ states contribute
121 equally to Π in chiral systems, we henceforth use the definition

$$\Pi_j = 2 \langle \mathbb{C}x_j \rangle_{E < 0}, \quad (3)$$

122 with $j = 1, \dots, d$ are the indices of the d crystallographic directions and where $E < 0$
123 indicates that the average is taken over the occupied states. Although seemingly identical

124 to the the skew polarization introduced in [36, 37] for topological insulators, and the
 125 mean chiral displacement of quantum walks [38], we emphasize that Π_j does not rely on
 126 any Bloch representation and is therefore defined also in amorphous phases. We stress
 127 that, even in the crystalline case, Π_j includes content beyond the skew polarization, as it
 128 resolves the weighted positions with a sub-unit-cell resolution. These differences are not
 129 mere technicalities, and will prove crucial in the next sections.

130 To gain more physical insight, it may be worth noting that in electronic systems, Π_j
 131 corresponds to the algebraic distance between the charge centers associated to the A and
 132 B atoms. While in mechanical networks, Π_j is the vector connecting the stress-weighted
 133 and displacement-weighted positions. A vanishing polarization indicates that the average
 134 locations of the stress and displacement coincide. Conversely, a finite chiral polarization
 135 reveals an asymmetric mechanical response discussed in [39, 40]. For the sake of clarity,
 136 before revealing topologically protected zero modes in amorphous phases, we first explore
 137 the consequences of a finite chiral polarization in periodic systems such as in the paradigm-
 138 atic example of the SSH model illustrated Fig. 2.

139

140 2.2 Chiral polarization: an interplay between Zak phases and frame 141 geometry.

142 We begin with a thorough discussion of crystalline materials, defined by periodic frames
 143 and Bloch Hamiltonians. Building on previous works on the electronic polarization [27–
 144 30],

145 we relate the chiral polarization of a crystalline material to the two Zak phases of waves
 146 projected on sub-lattices A and B when transported across the Brillouin zone. To do so, we
 147 first choose a unit cell and consider the basis of Bloch states $|\mathbf{k}, \alpha\rangle = \sum_{\mathbf{R}} e^{i\mathbf{k}\cdot\mathbf{R}} |\mathbf{R} + \mathbf{r}_\alpha\rangle$,
 148 where \mathbf{R} is a Bravais lattice vector, α labels the atoms in the unit cell and \mathbf{k} is the
 149 momentum in the Brillouin Zone (BZ). We henceforth use a convention where the Bloch
 150 Hamiltonian $H(\mathbf{k})$ is periodic in the BZ, see [28, 41] and [Appendix A]. More quantitatively,
 151 considering first Hamiltonians with no band crossing¹, we define the A sub-lattice Zak
 152 phase of the n^{th} energy band along the crystallographic direction j as

$$\gamma_j^A(n) = i \int_{\mathcal{C}_j} d\mathbf{k} \langle u_n | \mathbb{P}^A \partial_{\mathbf{k}} \mathbb{P}^A | u_n \rangle, \quad (4)$$

153 where the $|u_n(\mathbf{k})\rangle$ are the eigenstates of $H(\mathbf{k})$, and \mathcal{C}_j the non-contractible loops over the
 154 Brillouin zone defined along the d crystallographic axes. $\gamma_j^B(n)$ is defined analogously
 155 on the B sublattice. The (intercellular) Zak phase is given by the sum of $\gamma_j^A(n)$ and
 156 $\gamma_j^B(n)$ [43]. In [Appendix B], we show how to decompose the chiral polarization into a
 157 spectral and a frame contribution:

$$\Pi_j = \frac{a}{\pi} (\gamma_j^A - \gamma_j^B) + p_j, \quad (5)$$

158 where a is the lattice spacing (assumed identical in all directions), γ_j^A and γ_j^B are the
 159 sublattice Zak phases defined by

$$\gamma_j^A = \sum_{n < 0} \gamma_j^A(n). \quad (6)$$

¹In the situation where bands cross, our results should be generalized resorting to the Wilson loops of the non-commutative Berry connexion instead of the abelian Zak phase connection [42].

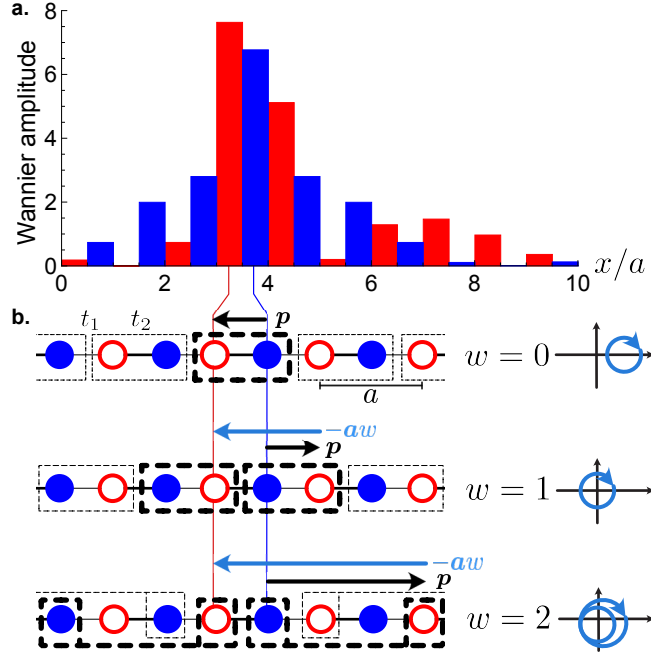


Figure 2: **Chiral polarization and Wannier functions.** **a.** Square of the Wannier amplitude projected into the A (red) and B (blue) sublattices for the ground state configuration of the two-band SSH model with hopping ratio $t_1/t_2 = 0.79$. a denotes the period of the 1D frame. The chiral polarization $\Pi = \langle x^A \rangle - \langle x^B \rangle$ is negative: the chain is left polarized regardless of the choice of unit cell. **b.** The winding number of the Bloch Hamiltonian encodes the chiral polarization *relative* to a given unit cell. The chiral polarization being a material property, the winding number w can therefore take any integer value when redefining the geometry of the unit cell as illustrated in the last column. Whatever the choice of the unit cell, the difference between the geometrical polarization and aw has a constant value given by the chiral polarization Π .

160 In Eq. (5) the p_j are the components of the geometrical-polarization vector connecting the
 161 centers of mass of the A and B sites in the unit-cell:

$$\mathbf{p} = \sum_{\alpha \in A} \mathbf{r}_\alpha - \sum_{\alpha \in B} \mathbf{r}_\alpha. \quad (7)$$

162 In crystals, Eqs. 5 quantifies the difference between the polarity of the ground-state wave
 163 function $\mathbf{\Pi}$ and the geometric polarization of the frame \mathbf{p} . This difference is finite only
 164 when the two sublattice Zak phases differ.

165 3 Topology of chiral insulators

166 We now elucidate the intimate relation between the chiral polarization and the band topol-
 167 ogy of chiral gapped phases defined on periodic lattices. We outline the demonstrations
 168 of our central results below and detail them in [AppendixB].

169

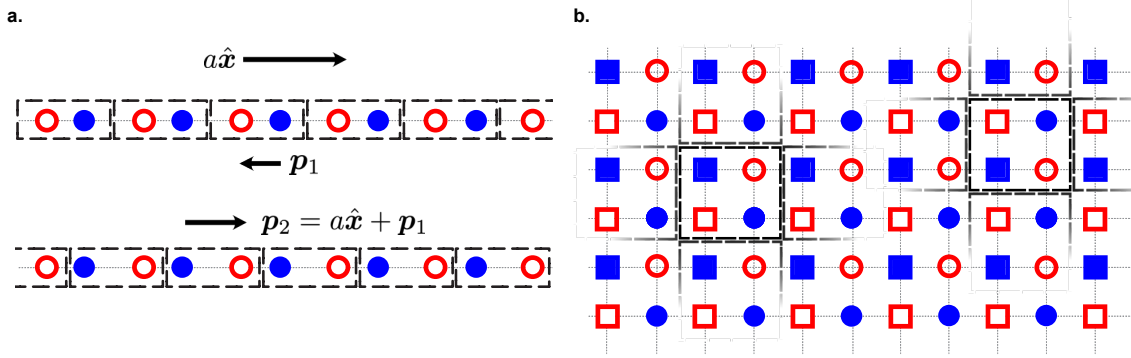


Figure 3: **Inferring the band topology from frame geometry.** **a.** The two-sites Wigner-Seitz cell on a 1D chiral frame have different geometrical polarizations; their difference is given by one Bravais vector. Consequently, we can always define the unit cell so that the Bloch Hamiltonian has a finite winding. **b.** All the Wigner-Seitz unit cells on the checkerboard lattice share the same (vanishing) chiral polarization. Therefore a single winding number w characterizes the Hamiltonians on this frame in virtue of Eq. (11). Evaluating the winding using the Wigner-Seitz cell compatible with the atomic limit of \mathcal{H} yields $w = 0$, by definition.

170 3.1 Sublattice Zak phases and winding numbers.

171 Computing the Wilson loop of the non-Abelian connection $\mathbf{A}_{n,m}(\mathbf{k}) = \langle u_n(\mathbf{k}) | \partial_{\mathbf{k}} | u_m(\mathbf{k}) \rangle$
 172 along \mathcal{C}_j , we show that chirality relates the d Zak phases $\gamma_j^A + \gamma_j^B$ to the windings of the
 173 Bloch Hamiltonian as

$$\gamma_j^A + \gamma_j^B = \pi w_j + 2\pi\mathbb{Z}, \quad (8)$$

174 where $w_j = i/(4\pi) \int_{\mathcal{C}_j} d\mathbf{k} \cdot \text{Tr}[\partial_{\mathbf{k}} H C H^{-1}] \in \mathbb{Z}$. The total Zak phase is quantized but the
 175 arbitrary choice of the origin of space implies that both γ^A and γ^B are only defined up
 176 to an integer. As a matter of fact, a mere $U(1)$ gauge transformation $|u_n\rangle \rightarrow e^{i\alpha_n(k)} |u_n\rangle$
 177 arbitrarily modifies $\gamma_j^A(n)$ and $\gamma_j^B(n)$ by the same quantized value: $\gamma_j^A(n) \rightarrow \gamma_j^A(n) +$
 178 πm , $\gamma_j^B(n) \rightarrow \gamma_j^B(n) + \pi m$, with $m \in \mathbb{Z}$. By contrast, the difference between the two
 179 sublattice Zak phases is left unchanged by the same gauge transformation which echoes
 180 its independence from the space origin. Evaluating the winding of $H(\mathbf{k})$ using the Bloch
 181 eigenstates (see [Appendix A]), we readily establish the essential relation²

$$\gamma_j^B - \gamma_j^A = \pi w_j \in \pi\mathbb{Z}. \quad (9)$$

182 Chirality quantizes the sublattice Zak phases of chiral insulators, even in the absence of
 183 inversion or any other specific crystal symmetry. γ_j^A and γ_j^B are however not independent.
 184 Combining Eqs. (8) and (9) we can always define the origin of space so that $\gamma_j^A = 0$ and
 185 $\gamma_j^B = \pi w_j$.

186 The d winding numbers of Eq. (9) characterize the topology of $H(\mathbf{k})$. In particular,
 187 if for a given Wigner-Seitz cell the corresponding $H(\mathbf{k})$ is associated to a finite winding
 188 ($w_j \neq 0$), then it cannot be smoothly deformed into the atomic limit defined over the
 189 same unit cell. We recall that the atomic limit of a material corresponds to a smooth
 190 deformation of the couplings to separate the energy scales so that the Wannier functions
 191 are exponentially localized, and respect the symmetries of the crystal [45]. In practice, it
 192 consists in choosing a unit cell including the strongest couplings.

²Note that this difference of Zak phases was recently denoted as a chiral phase index in [44].

193 The set of winding numbers is however poorly informative about the spatial distribu-
 194 tion of the charges in electronic systems, or about the stress and displacement distributions
 195 in mechanical structures. The values of w_j are defined only up to the arbitrary choice of
 196 unit cell required to construct the Bloch theory. A well known example of this limitation
 197 is given by the SSH model, where the winding of $H_{\mathbf{k}}$ can either take the values 0 or ± 1
 198 depending on whether the unit cell's leftmost site belongs to the A or B sublattice, see
 199 Fig. 2a and [AppendixB]. We show in the next section, how the chiral polarization allevi-
 200 ates this limitation.

201

202 3.2 Disentangling Hamiltonian topology from frame geometry.

203 Equations (5) and (9) provide a clear geometrical interpretation of the winding number
 204 w_j as the quantized difference between the geometrical and the chiral polarization:

$$\Pi_j = (p_j - a_j w_j). \quad (10)$$

205 We can now use this relation to illuminate the very definition of a chiral topological
 206 insulator. The chiral polarization $\Pi_j = 2\langle \mathbb{C}x_j \rangle_{E < 0}$ is a physical quantity that does not
 207 depend on the specifics of the Bloch representation. Therefore computing Π_j for two unit
 208 cells (1) and (2), we find that the windings of the two corresponding Bloch Hamiltonians
 209 $H^{(1)}(\mathbf{k})$ and $H^{(2)}(\mathbf{k})$ are related via Eq. (10) as

$$w_j^{(2)} - w_j^{(1)} = \frac{1}{a_j} (p_j^{(2)} - p_j^{(1)}). \quad (11)$$

210 This essential relation implies that one can always construct a Bloch representation of \mathcal{H}
 211 where $H(\mathbf{k})$ is topologically trivial, at the expense of a suitable choice of a unit cell. As
 212 a matter of fact, a redefinition of the unit cell can increase, or reduce the geometrical
 213 polarization, and therefore the winding numbers, by an arbitrary large multiple of a_j as
 214 illustrated in Fig. 2b.

215 For instance in the case of Hamiltonians with nearest neighbor couplings, applying
 216 Eq. (11) to Wigner Seitz unit cells ($|w_j| \leq 1$), we find that there exist as many topological
 217 classes of \mathcal{H} , as different geometrical polarizations in the Wigner-Seitz cells. This number
 218 provides a direct count of the chiral 'atomic limits' of \mathcal{H} .

219 Defining the topology of a chiral material therefore requires characterizing both the
 220 winding of its Bloch Hamiltonian, and the frame geometry. Remarkably, this interplay
 221 provides an insight on topological band properties from the sole inspection of the frame
 222 structure.

223

224 3.3 Inferring band topology from frame geometry.

225 There exists no trivial chiral phase in one dimension: one can always choose a Wigner-Seitz
 226 cell such that the Bloch representation of \mathcal{H} has a non-vanishing winding. As a matter of
 227 fact, the geometrical polarization of the Wigner-Seitz cells can only take two finite values
 228 of opposite sign depending on whether the leftmost site in a unit cell is of the A or B type,
 229 see Fig. 3a. Equation (11) therefore implies that, in $1D$, there always exists, at least, two
 230 topologically distinct gapped phases smoothly connected to two atomic limits. The two
 231 gapped phases are characterized by two distinct pairs of winding numbers defined by two
 232 inequivalent choices of unit cells. In other words all SSH Hamiltonians are topological.

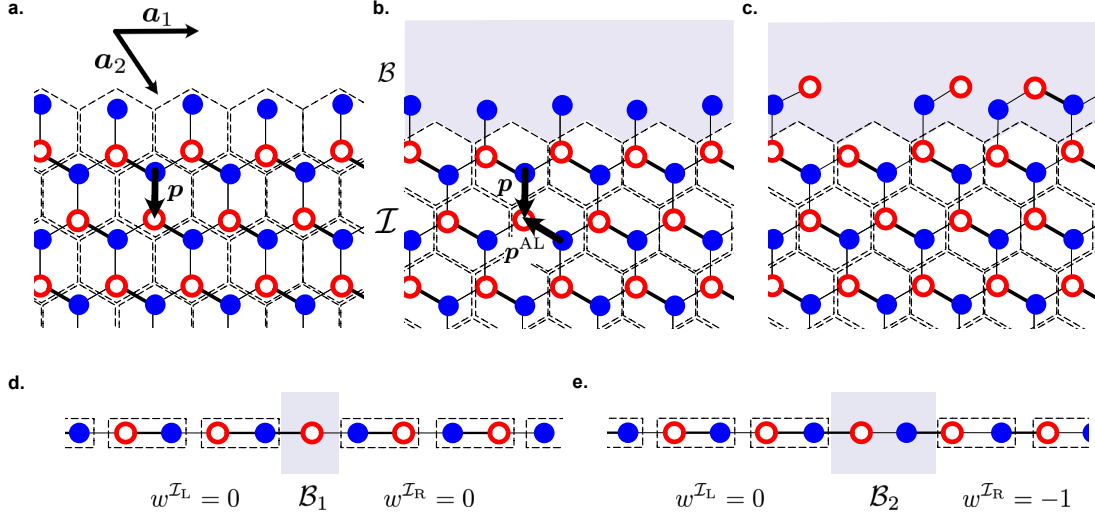


Figure 4: **Bulk-boundary correspondence.** **a.** A chiral crystal defined on a honeycomb frame is terminated by a clean zigzag edge incompatible with the atomic-limit Hamiltonian defined by keeping only the dominant couplings represented by thick solid lines. The dashed rectangles indicate the Wigner-Seitz cells allowing a tessellation compatible with the edge geometry. The arrow indicates the geometrical polarization \mathbf{p} . **b.** Same physical system. The crystalline bulk is now tiled using the unit cell compatible with the atomic limit. This requires a redefinition of the crystal boundary \mathcal{B} (shaded region). The arrows indicate the geometrical polarization of the new unit cell (\mathbf{p}^{AL}). The difference $\mathbf{p} - \mathbf{p}^{\text{AL}}$ is a Bravais lattice vector (\mathbf{a}_2). **c.** Same material as in (a.) and (b.) including a disordered interface \mathcal{B} bearing a non-zero chiral charge $\mathcal{M}^{\mathcal{B}}$. **d.** Two connected SSH chains. The Wigner-Seitz cell in the two materials are compatible with their atomic limits. The interface \mathcal{B}_1 separating the two materials is one-site wide. **e.** Redefining the Wigner-Seitz cell on the right hand side of the interface requires widening the boundary region. This redefinition makes the unit cell incompatible with the atomic limit. The winding of the Bloch Hamiltonian in \mathcal{I}_R takes a finite value and consequently modifies the zero-mode content of the boundary region.

233 Similarly, in $d > 1$ only frames having a geometrical polarization invariant upon re-
 234 definition of the Wigner-Seitz cell can support topologically trivial Hamiltonians. Equa-
 235 tion (11) indeed implies that a topologically trivial Hamiltonian \mathcal{H} constrains the frame
 236 geometry to obey $p_j^{(1)} = p_j^{(2)}$ for all pairs of unit cells and in all directions j . We show a
 237 concrete example of such a frame in Fig. 3b.

238 Before discussing the crucial role of the frame topology and geometry on the bulk-
 239 boundary correspondence of chiral phases, we extend these two notions to chiral insulators
 240 with a flat band.

241

242 3.4 Chiral polarization in the presence of a net chiral charge.

243 It is worth noting that the chiral polarization can also be defined and computed in the
 244 presence of an additional zero-energy flat band in the gap. As detailed in the [\[AppendixC\]](#)
 245 section, it then takes the form

$$\Pi_j = (p_j - p_j^{\text{ZM}}) + a(\gamma_j^A - \gamma_j^B) / \pi. \quad (12)$$

246 In this case, we lose the clear decomposition Π into geometrical and topological con-
 247 tributions. The geometrical polarization is corrected by \mathbf{p}^{ZM} which originates from a
 248 spectral contribution associated to the zero-energy band. Furthermore the second term
 249 on the r.h.s., the difference between two geometrical Zak phases, is not a topological wind-
 250 ing number anymore. Despite the seemingly complex form of Eq. (12), we show in the
 251 next section that the chiral polarization remains an effective tool to relate spectral bulk
 252 properties to the number of zero-energy states localized at boundaries.

253 4 Bulk-boundary correspondence

254 We now establish a bulk-boundary correspondence relating the chiral polarization to the
 255 number of zero modes supported by the free surface of a chiral insulator. For the sake of
 256 clarity, we discuss the two-dimensional case without loss of generality. We consider first a
 257 crystalline insulator \mathcal{I} terminated by a clean edge $\partial\mathcal{I}$ oriented along a Bravais vector, say
 258 \mathbf{a}_1 as illustrated in Fig. 4a.

259 The bulk of the insulator can be described by different types of unit cells. As illustrated
 260 in Fig. 4a, in the presence of a clean edge, it is natural to choose a unit cell which allows
 261 a tessellation of the whole system. However, this unit cell is generically incompatible with
 262 the atomic limit of the Hamiltonian, and therefore does not allow a direct count of the zero
 263 energy boundary states using the simple Maxwell-Calladine count. An obvious strategy
 264 hence consist in redefining the unit cell, as in Fig. 4b to match the constraints of the
 265 atomic limit. This redefinition comes at the expense of leaving sites outside of the bulk
 266 tessellation. We define this ensemble of sites as the boundary region \mathcal{B} . Keeping in mind
 267 that we can smoothly deform the Hamiltonian into its atomic limit without closing the
 268 gap, we use Eq. (2) to count the number of zero energy states hosted by \mathcal{B} . It is given by
 269 $\mathcal{V} = \mathcal{M}^{\mathcal{B}}$. An essential geometrical observation is that the net chiral charge in \mathcal{B} can be
 270 expressed as $\mathcal{N}^{\partial\mathcal{I}}(p_2^{\text{AL}} - p_2)$, where $\mathcal{N}^{\partial\mathcal{I}}$ is the edge length expressed in number of unit cells
 271 and p_2 is the geometrical polarization of the initial unit cell. We can now make use of the
 272 invariance of the chiral polarization formalized by Eq. (11) to relate the geometrical count
 273 of zeromodes to the winding of the Bloch Hamiltonian: $\mathcal{V} = \mathcal{N}^{\partial\mathcal{I}}(p_2^{\text{AL}} - p_2) = \mathcal{N}^{\partial\mathcal{I}}w_2^{\mathcal{I}}$. To
 274 arrive at a bulk boundary correspondence generic to all chiral insulators, we include the
 275 possibility of dealing with irregular interfaces featuring a net chiral charge $\mathcal{M}^{\mathcal{B}}$ as sketched

276 in Fig. 4c. We then find

$$\mathcal{V} = \mathcal{M}^{\mathcal{B}} + \mathcal{N}^{\partial\mathcal{I}} w_2^{\mathcal{I}}. \quad (13)$$

277 Three comments are in order. Firstly, the bulk boundary correspondence defined by
 278 Eq. (13) illuminates the geometrical implication of a nonzero winding: a finite $w_j^{\mathcal{I}}$ echoes
 279 the impossibility to tile a periodic frame with unit cells compatible with the Hamiltonian's
 280 atomic limit. Secondly, Eq. (13) is readily generalized to interfaces separating two chiral
 281 insulators \mathcal{I}_L and \mathcal{I}_R , where we simply have to apply the same reasoning on each side of
 282 the interface: $\mathcal{V} = \mathcal{M}^{\mathcal{B}} + \mathcal{N}^{\partial\mathcal{I}}(w^{\mathcal{I}_L} + w^{\mathcal{I}_R})$, see e.g. Figs. 4d and 4e. Thirdly, the formula
 283 given by Eq. (13) generalizes the Kane-Lubensky index introduced in their seminal work to
 284 count the zero-energy modes localized within isostatic mechanical networks [25]. We show
 285 that this index defines a bulk-boundary correspondence generic to all chiral insulators
 286 and even to flat band insulators such as hyperstatic lattices as further discussed in the
 287 [AppendixC].

288 5 Amorphous Chiral Insulators

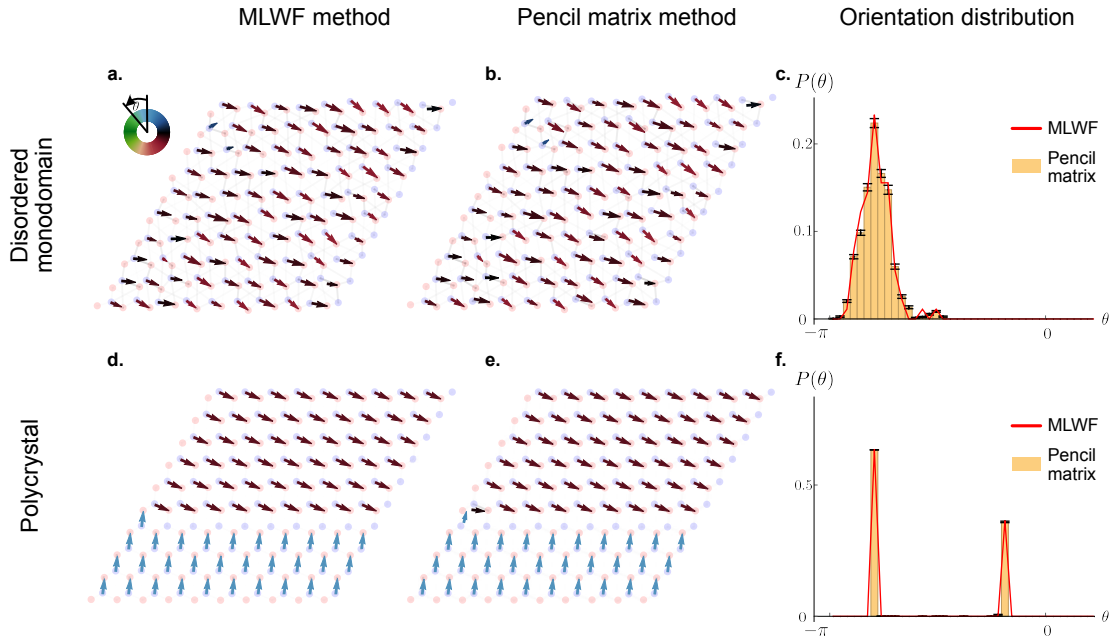


Figure 5: **Pencil matrix versus maximally localized Wannier functions** **a.** Single domain configuration with geometrical and spectral disorder. The chiral polarization field obtained from the maximally localized wannier functions is superposed. **b.** Chiral polarization field obtained from one realization of the pencil matrix procedure. **c.** Orientation distribution obtained from 50 values of α (bar chart), and from the maximally localized wannier functions (red solid line). **d.**, **e.**, **f.** correspond to the same information, this time for two crystalline domains.

289 In condensed matter, chiral symmetry is a low energy feature of electronic Hamilto-
 290 nians, which is unlikely to survive to strong structural disorder. Conversely, in photonic,
 291 acoustic or mechanical metamaterials chirality is built in by design and can therefore be
 292 present both in ordered or amorphous structures [20, 46]. In mechanical metamaterials

chirality is even more robust as it is inherent to any system assembled from elastically coupled degrees of freedoms [21]. In this section, we show how to generalize our physical characterization of zero energy modes to disordered chiral metamaterials.

Over the past two years a number of experimental, numerical and theoretical works showed that crystalline symmetries are not required to define topological insulators, see e.g. [47–50]. Unlike these pioneering studies where topologically inequivalent disordered phases are distinguished by abstract indices defined in real space and related to the quantification of edge currents, our framework solely based on the chiral polarization applies to chiral systems regardless of the presence or not of time reversal symmetry.

Our strategy follows from the fundamental relation: $\Pi_j = p_j - a_j w_j$ of Eq. (10). This relation implies a one-to-one correspondence between the chiral polarization and a topological spectral property quantized by the winding vector. The basic idea hence consists in probing the existence of topologically protected zero modes by *local* discontinuities in the chiral polarization field, even when no winding number or Zak phase can be defined. Relating topologically protected excitations to real-space singularities requires defining a local chiral polarization field $\mathbf{\Pi}(\mathbf{x})$. By definition, $\mathbf{\Pi}(\mathbf{x})$ measures the local imbalance of the wave function carried by the *A* and *B* sites. To express $\mathbf{\Pi}(\mathbf{x})$, it would be natural to consider eigenstates of the position operator $P\mathbf{x}P$ projected onto the occupied states of \mathcal{H} . However, in dimension $d > 1$, the different components of the projected position operator do not commute $[Px_jP, Px_kP] \neq 0$ for $j \neq k$, and do not possess common eigenstates. Instead, we express the polarisation in terms of the maximally localized states \widetilde{W}_m [28], which are centered on the position $\mathbf{x}_m \equiv \langle \widetilde{W}_m | \widehat{X} | \widetilde{W}_m \rangle$. These states generalize the Wannier functions in the absence of translational symmetry, see [AppendixE] for more details. We can then define the *local* chiral polarization as the weighted chiral position evaluated over \widetilde{W}_m :

$$\mathbf{\Pi}(\mathbf{x}_m) = 2 \langle \widetilde{W}_m | \mathbb{C} \widehat{X} | \widetilde{W}_m \rangle. \quad (14)$$

In practice, we can bypass the time consuming numerical determination of the \widetilde{W}_m by taking advantage of the so-called pencil-matrix method [51]. In short, the method consists in replacing in (14) the \widetilde{W}_m by eigenstates of a linear combination of the projected position components $L = \sum_j \alpha_j Px_jP$; $\sum \alpha_j = 1$. The dependence on α_i of the resulting chiral polarization is a measure of the non-commutativity of the Px_j typically associated to a nonvanishing Berry curvature. In practice, as illustrated in Fig. 5, the difference between the actual polarization, computed from the \widetilde{W}_m , and its approximation based on the *R*-matrix eigenstates is smaller than the distance between neighboring sites. Given the excellent agreement found both in mono and polycrystals, we henceforth use the pencil matrix method to locally measure the chiral polarization fields in disordered and amorphous structures out of reach of conventional chiral displacement characterizations [52].

To make the discussion as clear as possible we consider separately the two possible sources of randomness in a disordered chiral insulator: (i) geometrical disorder, which affects the frame geometry leaving the interaction between the *A* and *B* sites unchanged and (ii) Spectral disorder, which alters the interactions while leaving the frame geometry unchanged.

335

5.1 Topological zero modes on amorphous chiral frames.

The reasoning is easily explained starting from a concrete example. Fig. 6 shows the interface between two topologically distinct insulators, \mathcal{I}_T and \mathcal{I}_B , living on a honeycomb

338

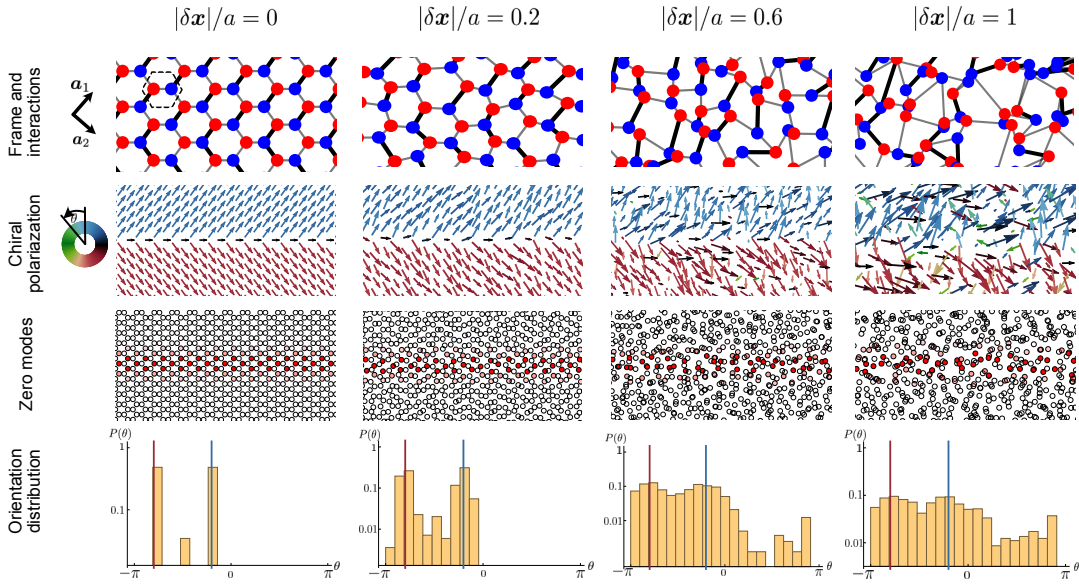


Figure 6: **Topological zero energy states on amorphous frames** First row: Sketch of the frame geometry for increasing positional disorder quantified by the maximal amplitude of the random displacements $|\delta\mathbf{x}|/a$. All panels show the vicinity of a boundary between two different insulators defined on the same frame but with different positions of the stronger couplings. The lines' width indicates the magnitude of the coupling strength. In all panels $t'/t = 20$. In the leftmost panel, we indicate the choice of the unit cell and of the crystallographic axes. Second row: Corresponding chiral polarization fields. The color indicates the orientation of $\mathbf{\Pi}(\mathbf{x})$ Third row: Magnitude of the zero-mode wave function. The zero mode is located at the boundary between topologically inequivalent states even on amorphous frames. Fourth row: Probability density function of the θ , the local orientation of the chiral polarization field. The distributions are peaked on the same two directions (vertical lines) regardless of the magnitude of disorder. This reveals the coexistence of two distinct topological phases robust to positional disorder.

339 frame. They correspond to distinct atomic limits of a nearest-neighbor tight binding
 340 Hamiltonians including two different hopping coefficients, see e.g. [53]. For the choice of
 341 unit cell sketched in Fig. 6, the winding vectors are $\mathbf{w}^{\mathcal{I}_T} = (0, 1)$ and $\mathbf{w}^{\mathcal{I}_B} = (1, 0)$. As
 342 a result the boundary region \mathcal{B} hosts one zero mode per unit cell located on the A sites.
 343 As expected from Eq. (10), on a homogeneous periodic frame, $\mathbf{\Pi}(\mathbf{x})$ takes two distinct
 344 values in the two regions, and is discontinuous across \mathcal{B} . Correspondingly, the distribution
 345 of the chiral polarization in the sample consists of two peaks centered on the two values
 346 associated to two topologically inequivalent phases, see Fig. 6 (left column).

347 We now disorder the frame by shifting all site positions by random displacements of
 348 maximal amplitude $|\delta\mathbf{x}|$ while preserving the magnitude of the interactions in the corre-
 349 sponding Hamiltonian \mathcal{H}_D . For sufficiently large displacements, it is impossible to keep
 350 track of the original periodic lattice, see Fig. 6 (first row). Nonetheless, we clearly see
 351 in the third row of Fig. 6 that the topologically protected zero modes located in \mathcal{B} are
 352 preserved, despite the lack of crystalline symmetry and the impossibility to define a Bloch
 353 Hamiltonian and its topological winding numbers. Note that unlike in [54] both the bulk
 354 and the boundary region are homogeneously disordered. Again, the existence and location
 355 of a line of zero modes is revealed by variations of the chiral polarization field. The vari-

356 ations of the orientation of $\mathbf{\Pi}(\mathbf{x})$ occurs over the penetration length-scale ℓ_G set by the
 357 energy gap. The coexistence of two topologically distinct amorphous phases is signalled
 358 by a (wider) bimodal distribution of $\mathbf{\Pi}(\mathbf{x})$ peaked on the same values as in the pure case,
 359 see Fig. 6 (last row). This robust phenomenology is further illustrated in Supplementary
 360 Video 1, showing the evolution of the polarization field and zero-mode location as the
 361 magnitude of disorder is increased.

362 This observation reflects a generic feature of chiral matter. Randomizing the frame ge-
 363 ometry cannot alter the energy gap provided that the graph defined by the coupling terms
 364 of \mathcal{H} has a fixed chiral connectivity. This observation implies that the concept of topolog-
 365 ical phase naturally applies to amorphous frames that can be continuously deformed into
 366 periodic lattices. In fact, the coexistence of different chiral insulators is effectively probed
 367 by the spatial distribution of the polarization field $\mathbf{\Pi}(\mathbf{x})$. Each peak of the distribution
 368 signals topologically inequivalent regions in amorphous chiral matter. The phase bound-
 369 aries are then readily detected by jumps of the chiral-polarization vector field over ℓ_G .
 370

371 5.2 Topological zero modes of disordered chiral Hamiltonians.

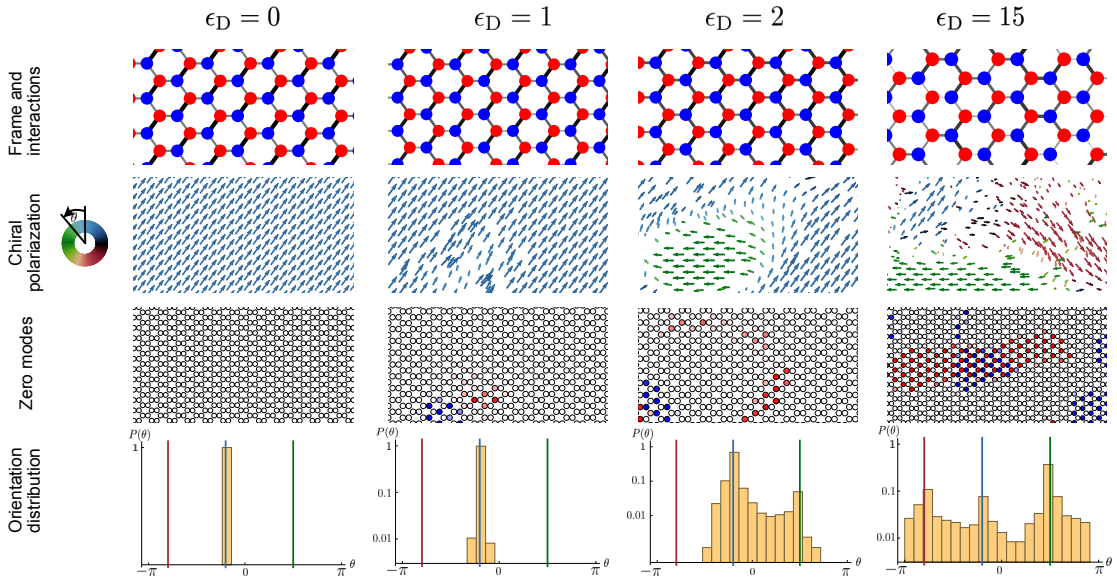


Figure 7: **Topological zero-energy states in the bulk of disordered chiral insulators** First row Sketch of the honeycomb frame and of the coupling strengths for increasing spectral disorder. The strengths of the couplings are represented by the width of the dark lines. Their randomness is quantified by the variance of the Gaussian couplings ϵ_D . The correlation length for all the examples is $\xi = 12a$. Second row Corresponding chiral polarization fields. The color indicates the orientation θ of $\mathbf{\Pi}(\mathbf{x})$. Third Row: Magnitude of the zero-energy modes on the A (red) and B (blue) sites. Fourth row: Probability density function of the orientation θ . Remarkably, even in the disordered cases, the distribution peaks only at values characteristic of the three phases of the homogeneous chiral Hamiltonian.

372 The case of spectral disorder is more subtle as it can trigger topological transitions.
 373 Again, we start with a concrete example. We use the same model of insulator as in the
 374 previous section. Considering the even simpler case of a perfect monocrystal, there is no

375 zero mode in the sample. Keeping the frame unchanged we add disorder to the interactions
 376 in the form of random perturbations to the coupling parameters. We note ϵ_D the width
 377 of the Gaussian disorder distribution, ξ its correlation length and ΔE the energy gap in
 378 the pure case. When $\epsilon_D/\Delta E \ll 1$ no zero mode exists in this finite system see Fig. 7
 379 first column. Consistently, the local chiral polarization hardly fluctuates in space and its
 380 distribution remains peaked around the same constant value.

381 By contrast as $\epsilon_D/\Delta E \sim 1$, zero energy modes emerge in the bulk. Their presence
 382 signals local the emergence of topologically inequivalent regions in the material triggered
 383 by local gap inversions. The distinct phases are revealed by the orientational distribution of
 384 $\mathbf{\Pi}(\mathbf{x})$: as disorder increases additional peaks grow at values of θ characteristic of the other
 385 two homogeneous topological insulators, Fig. 7 (last row). In the limit of strong disorder,
 386 the spatial extent of the coexisting phases is set by the disorder correlation length ξ as
 387 exemplified in Supplementary Movie 2. Gap closings also have a local signature in the
 388 polarization field. As $\mathbf{\Pi}(\mathbf{x}_m)$ is only defined at the generalized Wannier centers (Eq. (14)),
 389 $\mathbf{\Pi}(\mathbf{x}_m)$ cannot be computed at the center of a zero mode, which by definition does not
 390 support any Wannier mode. The proliferation of zero modes in the bulk is therefore
 391 signaled by an increasing number of holes in the polarization field.

392 The above observations do not rely on the specific model we use in Figs. 6 and 7.
 393 Generically, adding spectral disorder to a chiral Hamiltonian results in the nucleation of
 394 additional topological phases decorated by zero modes at their boundaries. Even in the
 395 absence of a Bloch theory, we can distinguish the topological nature of the coexisting
 396 phases by measuring their average chiral polarization. For spatially correlated disorder
 397 the spatial extent of each phase is set by the disorder correlation length ξ .

398

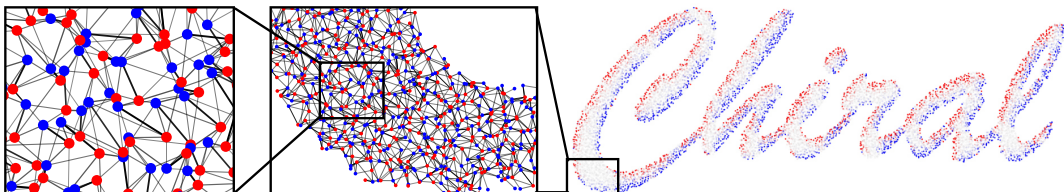


Figure 8: **Disordered chiral metamaterial** Macroscopic view and close ups on an amorphous frame supporting a disordered chiral insulator. The frame is defined adding a strong positional disorder to a Honeycomb lattice $|\delta\mathbf{x}| = a$. Using the same Hamiltonian as in Figs.6 and 7, we add spectral disorder corresponds to $\epsilon_D = 2$. Cutting the sample to form the word "chiral" reveals a continuous distribution of zero modes along the edge.

399 5.3 Designing topologically protected zero modes in amorphous chiral 400 matter.

401 It is worth stressing that disordered chiral insulators generically support topologically
 402 protected zero-energy modes at their boundaries. Unlike crystalline topological insulators,
 403 the lower the bulk and edge symmetries the more robust the edge states.

404 Cutting an amorphous sample into two parts without inducing the proliferation of
 405 boundary zero modes is virtually impossible. It would require cutting bonds while pre-

406 serving the connectivity between all pairs of A and B site connected by the local polar-
 407 ization vectors $\mathbf{\Pi}(\mathbf{x}_m)$; only this type of configurations can be continuously deformed into
 408 crystals having edges matching that of tilings generated by the unit cell of an atomic limit.
 409 These cuts require extreme fine tuning in macroscopic samples and are therefore virtually
 410 impossible to achieve. This property makes the design of zero energy wave guides very
 411 robust in amorphous chiral matter. As illustrated in Fig. 8.

412

413 5.4 Measuring the chiral polarization.

414 In this section we show that the chiral polarization is not only a powerful theoretical
 415 concept, but an actual material property readily accessible to experiments. Two scenarios
 416 are possible: when the (low energy) eigenfunctions can be measured, the chiral polarization
 417 can be directly evaluated using its definition, Eq. (3). This technique is straightforward
 418 e.g. in mechanical metamaterials [55], where the vibrational eigenmodes can be imaged in
 419 real space in response to mechanical actuation.

420 Alternatively, when spectral properties are out of reach of quantitative measurements,
 421 we can infer the value of the chiral polarization from the dynamic spreading of localized
 422 chiral excitations. This approach builds and generalizes the technique pioneered in the
 423 context of periodically driven photonic quantum walk [56, 57]. For the sake of clarity we
 424 henceforth limit our discussion to 1D, two-band insulators although the reasoning applies
 425 in higher dimensions.

426 We introduce the dynamical chiral polarization $\Pi_{\Psi}(t) = \langle \Psi(t) | \mathcal{C}\hat{X} | \Psi(t) \rangle$ defined over
 427 the time-evolved states $\Psi(t) = \exp(-iHt)\Psi(0)$, where $\Psi(0)$ is a localized chiral state.
 428 Should one be able to initialize an experiment in a Wannier State $\Psi(0) = W_{n,\mathbf{R}}$, the wave
 429 function would spread as in Fig. 9a, but remarkably the dynamical chiral polarization
 430 $\Pi_{\Psi}(t)$ would be stationary and equal to Π in a homogeneous system as illustrated in
 431 Fig. 9a, and demonstrated in the Method section. In practice, it would be always eas-
 432 ier to approximate the Wannier state by excitations Ψ_{AB} (resp. Ψ_{BA}) localized on two
 433 neighboring A and B sites (resp. B and A). The result of this procedure is shown in
 434 Fig. 9b and reveals that the long-time dynamics of $\Pi_{\Psi}(t)$ converges towards the chiral
 435 polarization Π . However, we stress that the essential information about the orientation
 436 of Π is already accessible at very short times and would not suffer from possible damping
 437 issues. When $\Pi_{\Psi}(t=0)$ and Π have opposite signs, we observe very large amplitude oscil-
 438 lations reflecting the dynamic reversal of the chirality of the wave packet at short times.
 439 Conversely when $\Pi_{\Psi}(t=0)$ and Π are parallel the convergence is very fast and devoid of
 440 large amplitude fluctuations.

441 It is worth noting that the chiral initial state $\Psi(t=0) = \Psi_{AB}$ is an atomic-limit
 442 eigenstate. The dynamics can then be seen as the result of a quench at $t=0$ starting
 443 from the atomic-limit Hamiltonian. The amplitude of the fluctuations in Fig. 9b then
 444 reveals the topological nature of the quench. As a last comment we stress that although
 445 our protocol is close to the chiral displacement method introduced and used in [56–59], it
 446 is not tight to a Bloch Hamiltonian model, but characterizes an intrinsic (meta)material
 447 property.

448 6 Conclusion

449 We have established a generic framework to characterize, elucidate and design the topo-
 450 logical phases of chiral insulators. In crystals, we show that the frame topology and the
 451 frame geometry conspire with Bloch Hamiltonian topology to determine the zero-mode

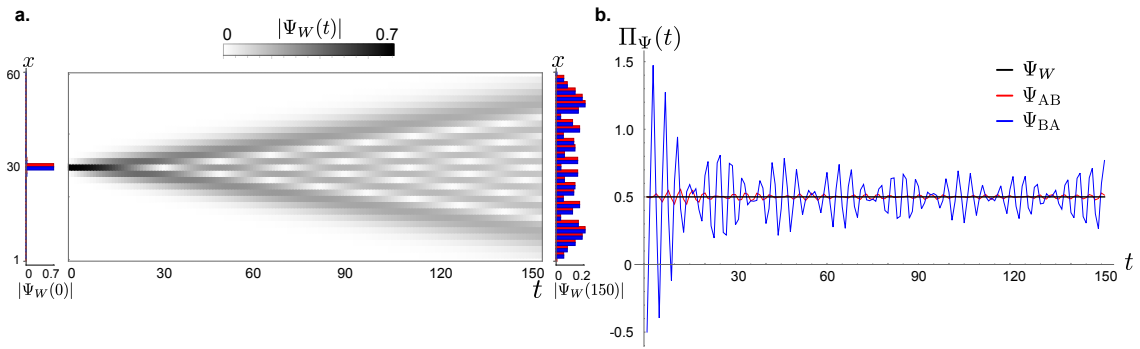


Figure 9: **Measuring the chiral polarization in time.** **a.** Left: Dynamical evolution of a Wannier state in the ground state of a two-band SSH model, with hopping ratio $t_1/t_2 = 0.1$. The state is localized in the middle of a finite system of 60 unit cells. Center: Time evolution of the wave-function amplitude. Right: The amplitude of the final state at time $t = 250$ is represented on the A (red) and B (blue) sites of the lattice. **b.** The dynamical chiral polarization $\Pi_\psi(t) = \langle \Psi(t) | \mathcal{C}\hat{X} | \Psi(t) \rangle$ corresponding to the protocol described in **a** is constant in time (black solid line). By comparison, the dynamical chiral polarization starting from a state $\Psi_{AB}(t = 0)$ (resp. $\Psi_{BA}(t = 0)$), localized on two neighboring sites A and B (resp. B and A) shows fluctuations around the static chiral polarization whose amplitude depends on the initial state. The sign $\Pi_\psi(t)$ is reversed at short time when the chiral polarization of the initial state is opposite to the static chiral polarization of the SSH chain. This results in large amplitude oscillations. The short time dynamics of Π_ψ therefore provides a direct access to the orientation of the material chiral polarization.

452 content of the bulk and interfaces. In the bulk, the frame topology fully determines the
 453 algebraic number of zero-energy modes counted by the chiral charge \mathcal{M} . Chiral insula-
 454 tors, however, are distinguished one another via their chiral polarization $\mathbf{\Pi}$ set both by
 455 the frame geometry and Bloch-Hamiltonian topology. At their surface, the number of
 456 zero-energy states is prescribed by the interplay between the Bloch Hamiltonian topology
 457 and the frame geometry in the bulk on one hand, and by the frame topology of the bound-
 458 ary on the other hand. This subtle tango goes beyond the bulk-boundary-correspondence
 459 principles solely based on Hamiltonian topology.

460 We have shown that chiral symmetry alone translates real-space properties into spectral
 461 phases without relying on any crystalline symmetry and translational invariance when
 462 expressed as a sublattice symmetry. Chiral symmetry does not merely complement the
 463 classification of topological quantum chemistry [45, 60–62] but also makes it possible to
 464 distinguish topological phases in amorphous matter. In disordered system, introducing the
 465 concept of chiral polarization field, we provide a practical platform to detect topological
 466 phases coexisting in disordered samples, an to design robust zero-mode wave guides at
 467 their boundaries.

468 We expect our framework to extend beyond Hamiltonian dynamics when dissipative
 469 processes obey the chiral symmetry [63]. We therefore conjecture that real-space topol-
 470 ogy, geometry and non-Hermitian operator topology should cooperate in chiral dissipative
 471 materials as diverse as cold atoms to photonics, robotic devices and active matter.

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 474 P. Massignan, A. Po, A. Schnyder and A. Vishwanath for insightful discussions.

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 476 breakthrough programs.

477 A Bloch theory convention and Wannier states.

478 A.1 Conventions for the Bloch decomposition.

479 For the sake of clarity, we first introduce the main quantities used throughout all the
 480 manuscript to describe waves in periodic lattices. We note $|\Psi_{n,\mathbf{k}}\rangle$ the Bloch eigenstates.
 481 They correspond to wavefunctions $\langle \mathbf{x} | \Psi_{n,\mathbf{k}} \rangle = \varphi_{n,\mathbf{k}}(\mathbf{x}) e^{i\mathbf{k} \cdot \mathbf{x}}$, where \mathbf{k} is the momentum in
 482 the Brillouin Zone (BZ), and where the normalized function $\varphi_{n,\mathbf{k}}$ has a periodicity of one
 483 unit cell [28]. In this article, we use the following convention to express the Bloch states
 484 as a superposition of plane waves:

$$|\Psi_{n,\mathbf{k}}\rangle = \sum_{\alpha} u_{n,\alpha}(\mathbf{k}) |\mathbf{k}, \alpha\rangle, \quad (15)$$

485 where α labels the different atoms in the crystal, and $|\mathbf{k}, \alpha\rangle$ represents the Fourier trans-
 486 form of the real-space position basis: $|\mathbf{k}, \alpha\rangle = \sum_{\mathbf{R}} \exp(i\mathbf{k} \cdot \mathbf{R}) |\mathbf{R} + \mathbf{r}_{\alpha}\rangle$, \mathbf{R} being a Bravais
 487 lattice vector and \mathbf{r}_{α} a site position within the unit cell. We stress that here the compo-
 488 nents $u_{n,\alpha}(\mathbf{k})$ are periodic functions of \mathbf{k} over the BZ. It is worth noting, however, that
 489 there exists multiple conventions to decompose the Bloch states as discussed e.g in the
 490 context of graphene-like systems in [64–66]. A common alternative uses nonperiodic com-
 491 ponents over the BZ which carry an additional phase encoding the position of each atom

492 within the unit cell: $|\Psi_{n,\mathbf{k}}\rangle = \sum_{\alpha} \tilde{u}_{n,\mathbf{k},\alpha} e^{i\mathbf{k}\cdot\mathbf{r}_{\alpha}} |\mathbf{k}, \alpha\rangle$. We will comment on the translation
 493 of our results from one convention to the other in the following.

494 A.2 Wannier functions.

495 By definition the Wannier function associated to a Bloch eigenstate is given by the inverse
 496 Fourier transform (up to a phase):

$$|W_{n,\mathbf{R}}\rangle = \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} e^{-i\mathbf{k}\cdot\mathbf{R}} |\Psi_{n,\mathbf{k}}\rangle. \quad (16)$$

497 where Ω is the volume of the BZ. Note that for sake of clarity, we here and henceforth
 498 assume that the spectrum does not include band crossings. The technical generalization
 499 of our results to degenerated spectra is straightforward but involves some rather heavy
 500 algebra, see e.g. [28]. In addition, to ease the notation and calculations we work with or-
 501 thogonal coordinates such that $\int_{\text{BZ}} dk_j = \Omega^{1/d}$, $\forall j$. The generalization to non-orthogonal
 502 lattices is straightforward and amounts to considering different geometrical factors along
 503 each reciprocal direction

504

505 A.3 Projected position operator and sublattice Zak phases.

506 Ignoring the distinction between the A and B sites, we can first compute the action of the
 507 position operator on the Wannier states following [28]:

$$\begin{aligned} \langle \mathbf{x} | \hat{X} | W_{n,\mathbf{R}} \rangle &= \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} \mathbf{x} e^{i\mathbf{k}\cdot(\mathbf{x}-\mathbf{R})} \varphi_{n,\mathbf{k}}(\mathbf{x}) \\ &= \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} \left(-i\partial_{\mathbf{k}} e^{i\mathbf{k}\cdot(\mathbf{x}-\mathbf{R})} + \mathbf{R} e^{i\mathbf{k}\cdot(\mathbf{x}-\mathbf{R})} \right) \varphi_{n,\mathbf{k}}(\mathbf{x}) \\ &= \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} e^{-i\mathbf{k}\cdot\mathbf{R}} \left[e^{i\mathbf{k}\cdot\mathbf{x}} (\mathbf{R} + i\partial_{\mathbf{k}}) \right] \varphi_{n,\mathbf{k}}(\mathbf{x}), \end{aligned} \quad (17)$$

508 where in the last step we applied an integration by parts, using that $|\Psi_{n,\mathbf{k}}\rangle = |\Psi_{n,\mathbf{k}+\mathbf{G}}\rangle$ with
 509 \mathbf{G} a primitive reciprocal vector. The generalization of Eq. (17) to the position operator
 510 projected on the sublattice $a = A, B$ is straightforward:

$$\langle \mathbf{x} | \hat{X} \mathbb{P}^a | W_{n,\mathbf{R}} \rangle = \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} e^{-i\mathbf{k}\cdot\mathbf{R}} \left[e^{i\mathbf{k}\cdot\mathbf{x}} (\mathbf{R} + i\partial_{\mathbf{k}}) \right] \mathbb{P}^a \varphi_{n,\mathbf{k}}(\mathbf{x}), \quad (18)$$

511 which allows us to define the average positions $\langle \mathbf{x}^a \rangle_{n,\mathbf{R}}$ restricted to the site $a = A, B$ and
 512 to the n^{th} band excitations:

$$\begin{aligned} \langle \mathbf{x}^a \rangle_{n,\mathbf{R}} &\equiv \langle W_{n,\mathbf{R}} | \mathbb{P}^a \hat{X} \mathbb{P}^a | W_{n,\mathbf{R}} \rangle \\ &= \frac{\mathbf{R}}{\Omega} \int_{\text{BZ}} d^d \mathbf{k} \langle \varphi_{n,\mathbf{k}} | \mathbb{P}^a | \varphi_{n,\mathbf{k}} \rangle + \frac{1}{\Omega} \mathbf{\Gamma}_{\text{Zak}}^a(n), \end{aligned} \quad (19)$$

513 where $|\varphi_{n,\mathbf{k}}\rangle = e^{-i\mathbf{k}\cdot\hat{X}} |\Psi_{n,\mathbf{k}}\rangle$, and $\mathbf{\Gamma}^a(n)$ is the vector composed of the d generalized
 514 sublattice Zak phases associated to the n -th band:

$$\Gamma_j^a(n) = i \int_{\text{BZ}} d^d \mathbf{k} \langle \varphi_{n,\mathbf{k}} | \mathbb{P}^a \partial_{k_j} \mathbb{P}^a | \varphi_{n,\mathbf{k}} \rangle \quad (20)$$

515 We can further simplify Eq. (19) noting that the orthonormality of the $|\varphi_{n,\mathbf{k}}\rangle$ implies
 516 $\langle \varphi_{n,\mathbf{k}} | \mathbb{P}^A + \mathbb{P}^B | \varphi_{n,\mathbf{k}} \rangle = 1$ and $\langle \varphi_{n,\mathbf{k}} | \mathbb{P}^A - \mathbb{P}^B | \varphi_{n,\mathbf{k}} \rangle = 0$, which yields $\langle \varphi_{n,\mathbf{k}} | \mathbb{P}^a | \varphi_{n,\mathbf{k}} \rangle =$
 517 $1/2$. All in all, we find a simple relation between the average of the position operator and
 518 the Zak phase of the Bloch eigenstates over the BZ:

$$\langle \mathbf{x}^a \rangle_{n,\mathbf{R}} = \frac{\mathbf{R}}{2} + \frac{1}{\Omega} \mathbf{\Gamma}^a(n). \quad (21)$$

519 B Chiral polarization, Zak phases and winding.

520 B.1 Chiral polarization and sublattice Zak phases.

521 We are now equipped to compute the chiral polarization, defined as the difference between
 522 the expected value of the projected position operators over the occupied eigenstates ($n <$
 523 0). It readily follows from Eq. (21) that $\mathbf{\Pi}$ corresponds to the difference of the sublattice
 524 Zak phases:

$$\begin{aligned} \mathbf{\Pi} &\equiv 2 \sum_{n < 0} \langle \mathbf{x}^A \rangle_{n, \mathbf{R}} - \langle \mathbf{x}^B \rangle_{n, \mathbf{R}} \\ &= \frac{2}{\Omega} \sum_{n < 0} \mathbf{\Gamma}^A(n) - \mathbf{\Gamma}^B(n). \end{aligned} \quad (22)$$

525 Two comments are in order. Firstly, the sum could have been also taken over the unoc-
 526 cupied states ($n > 0$). As $\mathbb{C}^2 = \mathbb{I}$, the sublattice phase picked up by $|\varphi_{n, \mathbf{k}}\rangle$ is indeed the
 527 same as that of its chiral partner $|\varphi_{-n, \mathbf{k}}\rangle = \mathbb{C} |\varphi_{n, \mathbf{k}}\rangle$. Secondly, we stress that Eq. (22)
 528 does not depend on the specific convention of the Bloch representation. This relation,
 529 however does not disentangle the respective contributions of the frame geometry and of
 530 the Hamiltonian on the chiral polarization. To single out the two contributions, we now
 531 use the specific Bloch representation (15). Given this choice, the sublattice Zak phase is
 532 naturally divided into two contributions leading to

$$\mathbf{\Gamma}^a(n) = \int_{\text{BZ}} d^d \mathbf{k} \sum_{\alpha \in a} (u_{n, \alpha}^* u_{n, \alpha} \mathbf{r}_\alpha + i u_{n, \alpha}^* \partial_{\mathbf{k}} u_{n, \alpha}). \quad (23)$$

533 The first term on the r.h.s. is the intracellular contribution to the Zak phase while the
 534 second is proportional to the sublattice intercellular Zak phase following to the definitions
 535 of [43]

$$\gamma_j^a(n) \equiv i \int_{\mathcal{C}_j} d\mathbf{k} \sum_{\alpha \in a} u_{n, \alpha}^*(\mathbf{k}) \partial_{\mathbf{k}} u_{n, \alpha}(\mathbf{k}). \quad (24)$$

536 Summing Eq.(23) over all occupied bands, and using the orthogonality of the chiral com-
 537 ponent $u_{n, \alpha}$ we then recover our central result:

$$\mathbf{\Pi} = \mathbf{p} + \frac{2}{\Omega^{1/d}} (\gamma^A - \gamma^B), \quad (25)$$

538 where $\mathbf{p} = \sum_{\alpha \in A} \mathbf{r}_\alpha - \sum_{\alpha \in B} \mathbf{r}_\alpha$ is the geometrical polarization of the corresponding unit-
 539 cell and $\gamma^a = \sum_{n < 0} \gamma^a(n)$. The chiral polarization is the sum of one contribution coming
 540 only from the frame geometry and one contribution characterizing the geometrical phase
 541 of the Bloch eigenstates.

542

543 B.2 Chiral polarization in different Bloch conventions.

544 Although the physical content of the chiral polarization does not depend on the choice of
 545 the Bloch convention, it is worth explaining how to derive its functional form for the other
 546 usual representation where $|\Psi_{n, \mathbf{k}}\rangle = \sum_{\alpha} \tilde{u}_{n, \alpha}(\mathbf{k}) e^{i\mathbf{k} \cdot \mathbf{r}_\alpha} |\mathbf{k}, \alpha\rangle$. Within this convention the
 547 vector of Zak phases take the form

$$\mathbf{\Gamma}^a(n) = i \int_{\text{BZ}} d^d \mathbf{k} \sum_{\alpha \in a} \tilde{u}_{n, \alpha}^* \partial_{\mathbf{k}} \tilde{u}_{n, \alpha}, \quad (26)$$

548 which does not allow the distinction between the geometrical and the Hamiltonian con-
 549 tributions to Π when performing the sum over the occupied band in Eq. (22). This
 550 observation further justifies our choice for the Bloch representation.

551

552 B.3 Quantization of the intercellular Zak-phase in chiral insulators.

553 To demonstrate the quantization of $\gamma_j = \gamma_j^A + \gamma_j^B$, we resort to the Wilson loop formalism
 554 reviewed e.g. in Ref. [42].

555 Let us first recall the definition of the non-Abelian Berry-Wilczek-Zee connection along
 556 the Brillouin zone for a set of smooth vectors $|u_n(\mathbf{k})\rangle, n = 1, \dots, M$:

$$\mathbf{A}_{nm}(\mathbf{k}) = \langle u_n(\mathbf{k}) | \partial_{\mathbf{k}} | u_m(\mathbf{k}) \rangle. \quad (27)$$

557 The associated Wilson loop operator defined along the path \mathcal{C}_j through the Brillouin zone
 558 is given by the ordered exponential

$$W_j = \overline{\exp} \left(- \int_{\mathcal{C}_j} d\mathbf{k} \cdot \mathbf{A}(\mathbf{k}). \right) \quad (28)$$

559 The topological properties of a generic gapped chiral Hamiltonian are conveniently cap-
 560 tured by smooth deformations yielding a flat spectrum $E = \pm 1$. The corresponding Bloch
 561 Hamiltonian is then given by

$$H = \begin{pmatrix} 0 & Q(\mathbf{k}) \\ Q^\dagger(\mathbf{k}) & 0 \end{pmatrix} \quad (29)$$

562 where $Q(\mathbf{k})$ is a nonsingular unitary matrix. Without loss of generality, we write the
 563 corresponding eigenstates as

$$|u_{\pm n}(\mathbf{k})\rangle = \frac{1}{\sqrt{2}} \begin{pmatrix} \pm Q(\mathbf{k}) |e_n^B\rangle \\ |e_n^B\rangle \end{pmatrix} \quad (30)$$

564 where the sign \pm identifies the sign of the eigenvalue $E = \pm 1$ and the normalized vectors
 565 $|e_n^B\rangle$ form a basis of the Hilbert space of Q^\dagger . The non-Abelian connection (27) for the
 566 negative (resp. positive) energy states then takes the simple form

$$\mathbf{A}_{nm}^-(\mathbf{k}) = \frac{1}{2} \langle e_n^B | Q^\dagger(\mathbf{k}) \partial_{\mathbf{k}} Q(\mathbf{k}) | e_m^B \rangle = \mathbf{A}_{nm}^+(\mathbf{k}) \quad (31)$$

567 It follows from the definition of the Wilson-loop operator (Eq. (28)) that the intercellular
 568 Zak phase for the negative energy bands $\gamma = \gamma^A + \gamma^B$ is defined in terms of the Wilson
 569 loops for the non-Abelian connection $\mathbf{A}^-(\mathbf{k})$ as

$$\gamma_j = -i \ln \det W_j^- \quad (32)$$

570 The quantization of all d intercellular Zakk phases then follows from Eqs (28) and (31):

$$\gamma_j = -i \operatorname{tr} \ln \left[\overline{\exp} \left(- \frac{1}{2} \int_{\mathcal{C}_j} d\mathbf{k} \cdot \partial_{\mathbf{k}} \ln Q(\mathbf{k}) \right) \right] \quad (33)$$

$$= \pi w_j \operatorname{mod} (2\pi) \quad (34)$$

571 where the $\text{mod}(2\pi)$ indetermination stems from the choice of the branch cut of the
572 complex \ln function, and where w_j is the standard winding of the chiral Hamiltonian (29):

$$w_j = \frac{i}{4\pi} \int_{\mathcal{C}_j} d\mathbf{k} \cdot \text{tr} [\partial_{\mathbf{k}} H \mathbb{C} H^{-1}] \in \mathbb{Z}, \quad (35)$$

$$= \frac{1}{2\pi i} \int_{\mathcal{C}_j} d\mathbf{k} \cdot \text{tr} [Q^{-1} \partial_{\mathbf{k}} Q]. \quad (36)$$

573 We therefore conclude that the d Zak phases are topological phases defined modulo 2π .

574

575 **B.4 Relating the sublattice Zak phases to the winding of the Bloch** 576 **Hamiltonian.**

577 We here demonstrate the essential relation given by Eq. (9). To do so, we relate the
578 winding w_j to the sublattice Zak phases by evaluating the trace in Eq. (35) using the
579 eigenstate basis. Noting that $\langle u_n | \partial_{\mathbf{k}} H(\mathbf{k}) \mathbb{C} H^{-1}(\mathbf{k}) | u_n \rangle = -2 \langle u_n | \mathbb{C} \partial_{\mathbf{k}} | u_n \rangle$, the winding
580 takes the simple form

$$w_j = -\frac{i}{2\pi} \int_{\mathcal{C}_j} d\mathbf{k} \sum_n \langle u_n | \mathbb{C} \partial_{\mathbf{k}} | u_n \rangle. \quad (37)$$

581 Decomposing the chiral operator on the two sublattice projectors $\mathbb{C} = \mathbb{P}^A - \mathbb{P}^B$, yields

$$\pi w_j = (\gamma_j^B - \gamma_j^A) \in \pi \mathbb{Z}. \quad (38)$$

582 **B.5 Quantization of the sublattice Zak phases.**

583 Eqs. (34) and (38) shows that both the sum and the difference of the sublattice Zak
584 phases are quantized:

$$\begin{aligned} \gamma_j^A + \gamma_j^B &= \pi w_j + 2\pi m, & m \in \mathbb{Z}, \\ \gamma_j^B - \gamma_j^A &= \pi w_j. \end{aligned} \quad (39)$$

585 It then follows that both sublattice phases γ_j^A and γ_j^B are integer multiples of π .

586

587 **B.6 How does the winding number of a chiral Bloch Hamiltonian change** 588 **upon unit cell redefinition?**

589 Starting from a chiral Hamiltonian \mathcal{H} , we demonstrate below the relation between the
590 winding numbers associated to the Bloch Hamiltonians constructed from different choices
591 of unit cells, Eq. (11).

592 The definition of Bloch waves and Bloch Hamiltonians require prescribing a unit cell.
593 Starting with a first choice of a unit cell geometry, say unit cell (1), we can write $H^{(1)}(\mathbf{k})$
594 in the chiral basis as

$$H^{(1)}(\mathbf{k}) = \begin{pmatrix} 0 & Q^{(1)} \\ Q^{\dagger(1)} & 0 \end{pmatrix}, \quad (40)$$

595 Let us now opt for a second choice of unit cell, say choice (2). The Bloch Hamiltonians
596 $H^{(1)}$ and $H^{(2)}$ are then related by a unitary transformation

$$H^{(2)} = U^\dagger H^{(1)} U, \quad (41)$$

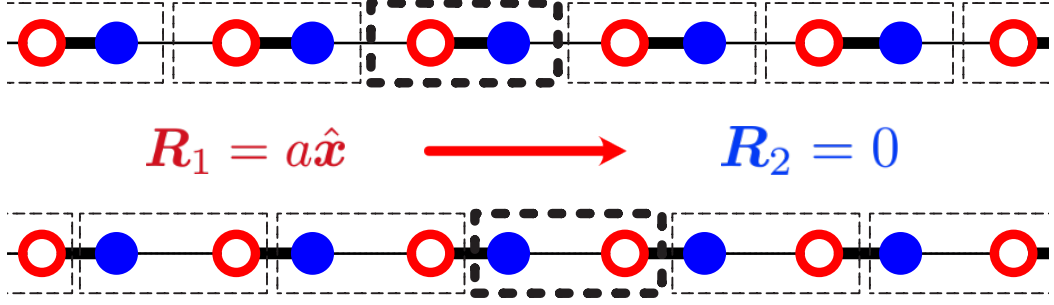


Figure 10: **Unit cell transformation.** We illustrate the definition of the \mathbf{R}_α vectors using the simple example of a SSH chain. For the first atom (empty symbol) $\mathbf{R}_1 = a\hat{x}$ while $\mathbf{R}_2 = 0$ for the second atom (solid symbol).

597 where the components of the unitary matrix are given by

$$U_{\alpha\beta} = \exp\left(i\mathbf{k} \cdot \mathbf{R}_\alpha^{(12)}\right)\delta_{\alpha\beta}, \quad (42)$$

598 where the \mathbf{R}_α^{12} are the Bravais vectors connecting the position of the atoms in the two unit-
 599 cell conventions, see Fig. 10 for a simple illustration. We note that, we have implicitly
 600 ignored the trivial redefinitions of the unit cell that reduce to permutations of the site
 601 indices. We can then express the winding of $H^{(2)}$ using Eq. (41) in the definition of
 602 Eq. (35), which yields

$$w_j^{(2)} = \frac{i}{4\pi} \int_{\mathcal{C}_j} d\mathbf{k} \operatorname{tr} \left[\partial_{\mathbf{k}} (UH^{(1)}U^\dagger) \mathcal{C} (UH^{(1)}U^\dagger)^{-1} \right]. \quad (43)$$

603 Expanding the gradient, using the trace cyclic property and noting that $[\mathcal{C}, U] = 0$, we
 604 find

$$w_j^{(2)} = w_j^{(1)} - \frac{i}{2\pi} \int_{\mathcal{C}_j} d\mathbf{k} \operatorname{tr} [\partial_{\mathbf{k}} U \mathcal{C} U^{-1}]. \quad (44)$$

605 This equation relates the winding numbers of the two Bloch Hamiltonians to the winding
 606 number of the transformation matrix U , which is by definition a geometrical quantity
 607 independent of \mathcal{H} . Using Eq. (42) leads to the remarkable relation which relates the
 608 spectral properties of the Hamiltonian to the unit-cell geometry

$$w_j^{(1)} - w_j^{(2)} = \frac{i}{2\pi} \int_{\mathcal{C}_j} d\mathbf{k} \operatorname{tr} [\partial_{\mathbf{k}} U \mathcal{C} U^{-1}] = \frac{1}{a_j} \left(\sum_{\alpha \in A} R_\alpha - \sum_{\alpha \in B} R_\alpha \right). \quad (45)$$

609 C Zero energy flat-band insulators.

610 We consider a flat-band chiral insulator, defined on a lattice with an non-vanishing chiral
 611 charge. In mechanics this situation is readily achieved adding extra bonds to further
 612 rigidify an otherwise isostatic lattice. It is characterized by a finite gap separating positive

613 and negative energy states and by an additional flat band at $E = 0$. In such a phase, there
 614 may exist additional zero energy edge states in addition to the bulk zero-energy modes.
 615 These edge states are analogous to the topological edge modes of insulators. Our goal is
 616 here to derive a bulk-boundary correspondence for these materials and provide a count of
 617 their zero-energy edge states. We will show that this correspondence involves the specific
 618 geometry of the eigenstates as opposed to their topology in the case of genuine insulators.

619 To show this we first derive the expression of the chiral polarization in the presence
 620 of a finite bulk chiral charge. Our starting point is Eq. (22), which relates to the chiral
 621 polarization of a crystal to the sublattice Zak phases given by Eq. (20):

$$\mathbf{\Pi} \equiv 2 \sum_{n < 0} \langle \mathbf{x}^A \rangle_{n, \mathbf{R}} - \langle \mathbf{x}^B \rangle_{n, \mathbf{R}} = \frac{2}{\Omega} \sum_{n < 0} \mathbf{\Gamma}^A(n) - \mathbf{\Gamma}^B(n). \quad (46)$$

622 The sum over all the negative energy bands $n < 0$ is half the sum over the non-zero energy
 623 states $n \neq 0$ given by

$$\begin{aligned} \sum_{n \neq 0} \mathbf{\Gamma}^a(n) &= \int_{\text{BZ}} d^d \mathbf{k} \sum_{\alpha \in a} \sum_n u_{n, \alpha}^* u_{n, \alpha} \mathbf{r}_\alpha + \frac{2}{\Omega^{1/d}} \gamma^a \\ &= \int_{\text{BZ}} d^d \mathbf{k} \sum_{\alpha \in a} \left(1 - \sum_{n_0} u_{n_0, \alpha}^* u_{n_0, \alpha} \right) \mathbf{r}_\alpha + \frac{2}{\Omega^{1/d}} \gamma^a. \end{aligned} \quad (47)$$

624 In the last line, we single out the role of the bulk zero-energy modes indexed by n_0 . Using
 625 the above expression to evaluate the r.h.s. of Eq. (46), we find an expression similar to
 626 Eq. (25) in the main text:

$$\mathbf{\Pi} = (\mathbf{p} - \mathbf{p}_{\text{ZM}}) + \frac{2}{\Omega^{1/d}} (\gamma^A - \gamma^B). \quad (48)$$

627 A first noticeable difference with Eq. (25) is a spectral correction to the geometrical polar-
 628 ization stemming from the localized zero-energy bulk modes. This zero-mode polarization
 629 is given by

$$\mathbf{p}_{\text{ZM}} = - \int_{\text{BZ}} d^d \mathbf{k} \sum_{n_0} \left(\sum_{\alpha \in A} - \sum_{\alpha \in B} \right) u_{n_0, \alpha}^* u_{n_0, \alpha} \mathbf{r}_\alpha. \quad (49)$$

630 Three comments are in order. Firstly, we stress that while the geometrical polarization \mathbf{p}
 631 depends on the choice of origin in the presence of an excess of chiral charge, the difference
 632 $\mathbf{p} - \mathbf{p}_{\text{ZM}}$, and $\mathbf{\Pi}$, are both independent of the frame's origin. Secondly, unlike in insulators,
 633 the difference between the intercellular sublattice Zak phases, $\gamma^A - \gamma^B$ does not identify
 634 with the winding number of the Bloch Hamiltonian. In fact it is not a topological quantity:
 635 it continuously depends on the specific couplings of the Hamiltonian. Finally, we point
 636 that, by definition, the chiral polarization does not depend on the Bloch convention.
 637 A change in the Bloch convention changes the geometrical polarization, the zero-mode
 638 polarization, and the intercellular zak phases in such a way that all corrections cancel one
 639 another.

640 Equipped with Eq. (48), we now turn to the generalization of the bulk boundary
 641 correspondence for flat-band insulators. We consider a crystalline material \mathcal{S} terminated
 642 by a clean edge $\partial \mathcal{S}$ oriented along the Bravais vector \mathbf{a}_1 . This edge may host \mathcal{V}^{NT} non-
 643 trivial zero-energy modes, in addition to the (trivial) bulk zero modes associated to the
 644 flat band. The edge defines a unit cell that may not be compatible with that of the
 645 atomic limit. We can nonetheless extend the edge region such that it matches the unit-cell
 646 compatible with the atomic limit (AL). The idea being that \mathcal{V}^{NT} is fully determined by
 647 the additional chiral charge of the edge with respect to that provided by the bulk chiral

648 charge density. Following the same reasoning as in the main text, this extra chiral charge
 649 is given by the difference of geometrical polarization and zero-mode polarization:

$$\mathcal{V}^{\text{NT}} = \mathcal{N}^{\mathcal{B}} [(p_2 - p_{\text{ZM}_2})_{\text{AL}} - (p_2 - p_{\text{ZM}_2})], \quad (50)$$

650 where $\mathcal{N}^{\mathcal{B}}$ is the boundary length expressed in units of unit-cell length. The first term
 651 is computed in the unit cell compatible with the atomic limit, and the second term is
 652 computed in the original unit cell defined by the edge $\partial\mathcal{S}$.

653 The invariance of the chiral polarization with respect to unit cell transformations allows
 654 the connection with the intercellular sublattice Zak phase:

$$\left(p_2 - p_{\text{ZM}_2} + \frac{2}{\Omega^{1/d}} (\gamma_2^A - \gamma_2^B) \right)_{\text{AL}} = p_2 - p_{\text{ZM}_2} + \frac{2}{\Omega^{1/d}} (\gamma_2^A - \gamma_2^B), \quad (51)$$

655 where AL denotes the terms evaluated in the unit-cell compatible with the atomic limit.
 656 All in all, the non-trivial zero-energy content of flat band insulators is given by a formula
 657 which generalizes Eq. (13):

$$\mathcal{V}^{\text{NT}} = \mathcal{N}^{\mathcal{B}} \frac{2}{\Omega^{1/d}} [(\gamma_2^A - \gamma_2^B) - (\gamma_2^A - \gamma_2^B)_{\text{AL}}]. \quad (52)$$

658 It is worth noting that in the case of genuine insulator, $(\gamma_2^A - \gamma_2^B)_{\text{AL}} = -w_{\text{AL}} = 0$ since it
 659 corresponds to the winding number in the unit cell compatible with the AL. Once again
 660 the chiral polarization field and its relation with the geometric phases allow us to predict
 661 the existence of non-trivial zero-energy modes by observing the local discontinuities of the
 662 chiral polarization field at any interface.

663 D Basis of localised states: matrix pencil

664 Finding a localized basis of the space of negative energy states poses several challenges
 665 when working in high-dimensional systems. In one dimension, this is an easy task that
 666 can be directly solved by finding the eigenstates of the projected position operator, $P\hat{X}P$,
 667 where

$$P = \sum_{E < 0} |\Psi_E(\mathbf{r})\rangle \langle \Psi_E(\mathbf{r})|, \quad (53)$$

668 is the projector onto the occupied energy states and $|\Psi_E\rangle$ are the eigenstates of the real
 669 space hamiltonian \mathcal{H} .

670 It would be tempting to generalize this approach to two and three dimensional sys-
 671 tems to find a common basis for the independent components of the projected position:
 672 $P\hat{\mathbf{R}}P \equiv (P\hat{X}P, P\hat{Y}P, P\hat{Z}P)$. However, in general these components do not commute.
 673 As proposed in the seminal work of Marzari and Vanderbilt [67], a workaround consists in
 674 computing the set of maximally localized Wannier functions $\{W_n\}$, which minimizes the
 675 quadratic spread:

$$\Delta r^2 = \frac{1}{Na^2} \sum_n^N \left[\langle W_n | (P\hat{\mathbf{R}}P)^2 | W_n \rangle - |\langle W_n | P\hat{\mathbf{R}}P | W_n \rangle|^2 \right], \quad (54)$$

676 Other minimization functions can be used to define localized states such Foster-Boys or
 677 the Edmiston-Ruedenberg criteria [68].

678 Here, we introduce an alternative method based on the matrix pencil of the projected
 679 positions. The matrix pencil of two matrices PXP and PYP corresponds to their linear

680 combination $L(\alpha_1, \alpha_2) = \alpha_1 PXP + \alpha_2 PYP$, where α_i are two non-zero real coefficients [51].
 681 When $[PXP, PYP] \neq 0$, the eigenvectors of the matrix pencil leads to a localised basis,
 682 whose spreading is comparable to the standard Maximally Localized Wannier Functions
 683 (MLWF) at a much reduced computational cost. To illustrate this result, we study in
 684 detail the case of a chiral Halmiltonian on a 10×10 honeycomb lattice, see Fig. 6. We
 685 compare the MLWF and the matrix pencil methods in Fig. 11. We implement MLWF
 686 method using a gradient descent protocol [68]. The minimization of Ω converges slowly
 687 as shown in Fig. 11b. We now discuss the case of the matrix pencil method and first note
 688 that the diagonalization of L for the two limiting cases $\alpha_1 = 0$ and $\alpha_2 = 0$ corresponds
 689 to finding a basis of completely localized states along the \hat{y} and \hat{x} direction, respectively.
 690 In the general case where $\alpha_{1,2} \neq 0$, L corresponds to the position operator along the
 691 direction $\alpha_1 \hat{x} + \alpha_2 \hat{y}$. It is therefore convenient to parametrize this axis according to its
 692 polar angle: $L(\theta) = \cos \theta PXP - \sin \theta PYP$. Fig. 11c shows the spreading functional
 693 evaluated at the local basis obtained from $L(\theta)$ for different polar angles. In practice,
 694 we achieve a comparable and even better localisation with respect to the MLWF method
 695 after 200 iterations. The only values of θ for which the matrix pencil method is not
 696 effective corresponds to the crystallographic directions of the lattice, see Fig. 11d. As these
 697 directions are known a priori, we can safely and effectively use the matrix pencil method
 698 to compute a set of localized states. The gain in terms of computing time is obvious.
 699 Both the diagonalization of L and each minimization step of Δr^2 have a computational
 700 complexity of order N^2 , where N is the system size. Choosing a value of θ avoiding the
 701 Bravais directions allows us to find a set of localized states in one step using the matrix
 702 pencil method. It is also worth noting that this method is unrelated to the chiral symmetry
 703 of the Hamiltonians considered in the main text and applies broadly.

704 We now switch to disordered systems and illustrate the performance of the matrix
 705 pencil method in Figs. 11d, e and f. The minimization of Δr^2 for the MLWF method is
 706 more time consuming than in crystals, Fig. 11e. Conversely, the diagonalization time of
 707 L remains unchanged. The difference with the ordered case is visible when plotting the
 708 spreading function Ω as a function of θ . The peaks along the crystallographic directions
 709 widen, as expected, when disorder increases.

710 In practice, we compute the θ average of the chiral polarization associated to a set of
 711 localized eigenstates along the θ directions which yields excellent approximations of the
 712 Wannier states, see Fig. 5.

713 E Chiral polarization in amorphous materials.

714 We have seen that the chiral polarization does not depend on the specifics of the unit cell:
 715 it is an intrinsic property of the material. In fact, as we show below, this framework is far
 716 more general and we can define the chiral polarization in amorphous solids.

717 We start by revisiting the definition of the chiral polarization in a crystal given by
 718 eq. (22):

$$\mathbf{\Pi} \equiv 2 \sum_{n < 0} \langle \mathbf{x}^A \rangle_{n, \mathbf{R}} - \langle \mathbf{x}^B \rangle_{n, \mathbf{R}}. \quad (55)$$

719 Strictly speaking this polarization is defined at the position \mathbf{R} . However, the discrete
 720 translational invariance of the crystal and by consequence, of the Wannier functions, makes
 721 the polarization field homogeneous. We can thus we drop the \mathbf{R} indices.

722 The definition of the Wannier function as the inverse Fourier transform of the Bloch
 723 eigenstate cannot be used when dealing with a disordered configuration. Instead, we work

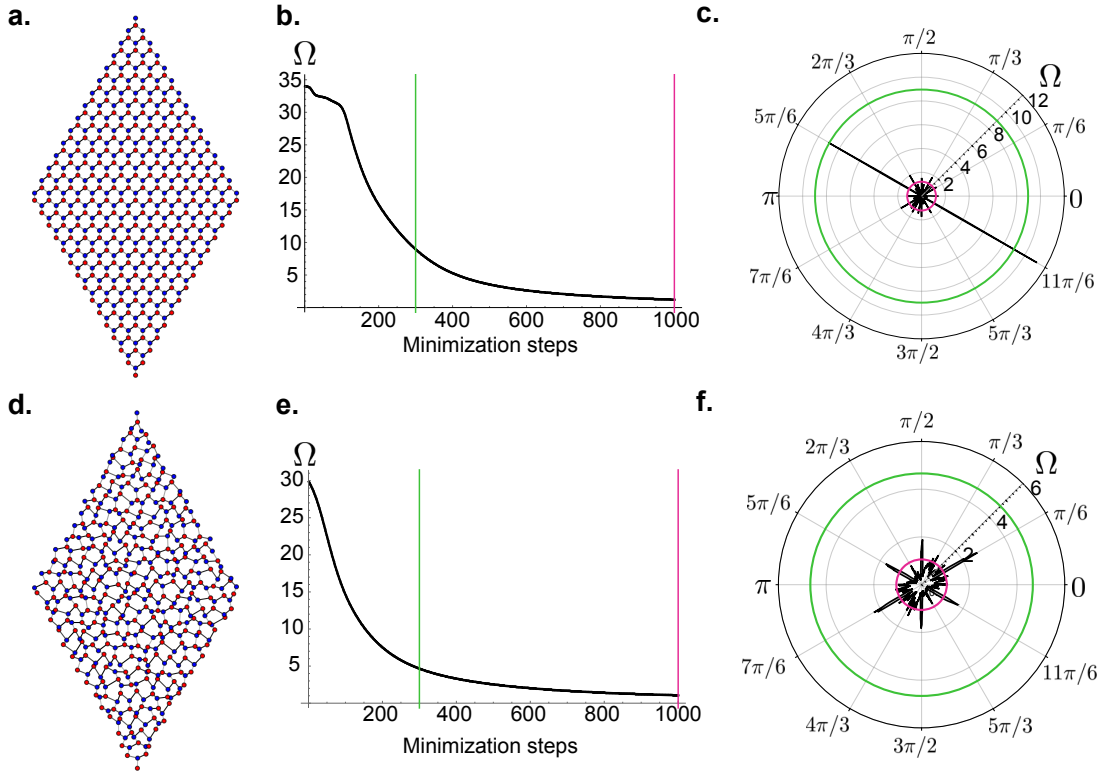


Figure 11: **Comparison between the matrix pencil and the MLWF method a.** Sketch of the frame geometry for a crystalline chiral honeycomb lattice made of 15×15 unit cells. **b.** Spreading functional as a function of the amount of minimization steps for the determination of maximally localised Wannier functions. After 300 (green) and 1000 (pink) minimization steps, the spreading corresponds to $\Omega_{\text{Wannier}_{300}} \approx 8.9$ and $\Omega_{\text{Wannier}_{1000}} \approx 1.2$, respectively. **c.** Spreading functional as a function of the angle θ for the localised basis determined from the matrix pencil $L(\theta)$ (black line). In green and pink we show the spreading obtained from the wannier states after 300 and 1000 minimization steps, respectively. Except for a few given directions, notably $\theta = 11\pi/6$ and $\theta = 5\pi/6$, the matrix pencil method gives a more localised basis at a much lower computational cost. **d., e., f.** Same as before applied for a disordered system with $|\delta\mathbf{x}|/a = 0.2$, $\epsilon_D = 0.4$, and $\xi = 10a$. (see figs. 6 and 7)

724 with a another set of fully localized functions: the eigenstates of the projected position
 725 operator onto the occupied bands [42]. The projected position operator is given by $P\hat{X}P$,
 726 where

$$P = \sum_{E<0} |\Psi_E(\mathbf{r})\rangle \langle \Psi_E(\mathbf{r})|, \quad (56)$$

727 is the projector onto the occupied energy states (not to be confused with the projectors
 728 \mathbb{P}^α), and the $|\Psi_E\rangle$ are the eigenstates of the real space hamiltonian \mathcal{H} . Let us denote
 729 the m^{th} eigenstate of the projected position operator as \widetilde{W}_m (notice that there are as
 730 many eigenstates as occupied energy states of the Hamiltonian). This is a localized func-
 731 tion around the center given by $\mathbf{x}_m = \langle \widetilde{W}_m | \hat{X} | \widetilde{W}_m \rangle$, similarly to the Wannier centers.
 732 Moreover, using each localized function, we can compute the difference of the weighted
 733 positions on both sublattices, in other words, the local chiral polarization:

$$\mathbf{\Pi}(\mathbf{x}_m) = 2 \langle \widetilde{W}_m | \mathbb{C}\hat{X} | \widetilde{W}_m \rangle \quad (57)$$

734 In a periodic frame, the eigenstates of the projected position operator reduce to a linear
 735 combination of the Wannier functions W_n : $|\widetilde{W}_m\rangle = \sum_n V_{mn} |W_n\rangle$, with $n < 0$, indicating
 736 the occupied energy bands, V a unitary matrix in the energy space, and V_{mn} a diagonal
 737 matrix in the position space. We can then rewrite the chiral polarization in eq. (57) as

$$\begin{aligned} \mathbf{\Pi}(\mathbf{x}_m) &= 2 \sum_{n,l} \langle W_n | V_{mn}^\dagger \mathbb{C} V_{ml} | W_l \rangle \\ &= 2 \sum_{n<0} \langle W_n | \mathbb{C}\hat{X} | W_n \rangle, \end{aligned} \quad (58)$$

738 where in the last line we used the fact that the V_{ml} commutes with $\mathbb{C}\hat{X}$ and the unitarity
 739 of V . As a result, we recover the first expression defined in crystals using the Bloch
 740 formalism as given by Eq. (22).

741 F Chiral polarization of time evolved Wannier states.

742 In Ref. [57], the mean chiral displacement under Hamiltonian dynamics was introduced as a
 743 measure of the Zak phase of periodic Hamiltonians in $d = 1$. This quantity characterizes a
 744 representation of a Hamiltonian associated to a given unit cell definition, and corresponds
 745 to the long-time displacement of an initially fully localized state, measured in units of
 746 the unit-cell size. As a consequence, several choices of unit cells were necessary to fully
 747 characterize the dynamics of a given (meta)material [56]. Although seemingly similar in
 748 its formal definition, the chiral polarization which we extensively use in this article is an
 749 intrinsic (meta)material property, defined in real space, and which does not rely on any
 750 underlying frame periodicity, Eq. (5). In the specific case of periodic frames $\mathbf{\Pi}$ crucially
 751 resolves the chiral imbalance of wave packets with a sub-unit-cell resolution.

752 In this method section, we show how $\mathbf{\Pi}$ relates to the dynamics of a maximally localized
 753 Wannier state spreading in the bulk of a chiral crystal. To do so we consider the time
 754 evolution of a wave function $|\psi_n(t)\rangle = U(t) |W_{n,\mathbf{R}}\rangle$ starting from a of a Wannier state in
 755 band n , initially localized at \mathbf{R} , with an evolution operator $U(t) = \exp(-iHt)$. Using the

756 notations introduced in Eq. (16), the position at time t is given by

$$\begin{aligned} & \langle \mathbf{x} | \hat{X} | \psi_n(t) \rangle \\ &= \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} \mathbf{x} e^{i\mathbf{k} \cdot (\mathbf{x} - \mathbf{R})} e^{-iE_n(\mathbf{k})t} \varphi_{n,\mathbf{k}}(\mathbf{x}) \end{aligned} \quad (59)$$

$$= \Omega^{-1} \int_{\text{BZ}} d^d \mathbf{k} e^{-i\mathbf{k} \cdot \mathbf{R}} \left[e^{i\mathbf{k} \cdot \mathbf{x}} (\mathbf{R} + \mathbf{v}_n(\mathbf{k})t + i\partial_{\mathbf{k}}) \right] \varphi_{n,\mathbf{k}}(\mathbf{x}), \quad (60)$$

757 where $\mathbf{v}_n(\mathbf{k}) = \partial_{\mathbf{k}} E_n(\mathbf{k})$ is the group velocity in the energy band n . We can also generalize
758 Eq. (21) to define the instantaneous average positions restricted to the $a = A, B$ sublattices
759 which read

$$\langle \mathbf{x}^a(t) \rangle_{n,\mathbf{R}} \equiv \langle \psi_n(t) | \mathbb{P}^a \hat{X} \mathbb{P}^a | \psi_n(t) \rangle \quad (61)$$

$$= \oint_{\mathbf{k}} \frac{1}{2} (\mathbf{R} + \mathbf{v}_n(\mathbf{k})t) + \frac{1}{\Omega} \Gamma_{\text{Zak}}^a(n) \quad (62)$$

$$= \frac{1}{2} \mathbf{R} + \frac{1}{\Omega} \Gamma_{\text{Zak}}^a(n) \quad (63)$$

$$= \langle \mathbf{x}^a(t=0) \rangle_{n,\mathbf{R}}. \quad (64)$$

760 This result indicates that the chiral polarization of each Wannier state is a stationary
761 quantity although they all evolve in time. When summed over (half of) the spectrum, we
762 recover the static definition of the chiral polarization

$$\mathbf{\Pi}(t) = \text{Tr}(U^{-1}(t) \mathbb{C} \hat{X} U(t)) \quad (65)$$

$$= \sum_n \langle \psi_n(t) | \mathbb{C} \hat{X} | \psi_n(t) \rangle \quad (66)$$

$$= 2 \sum_{n < 0} \langle \mathbf{x}^A(t) \rangle_{n,\mathbf{R}} - \langle \mathbf{x}^B(t) \rangle_{n,\mathbf{R}}$$

$$= \frac{2}{\Omega} \sum_{n < 0} \Gamma_{\text{Zak}}^A(n) - \Gamma_{\text{Zak}}^B(n). \quad (67)$$

763 We note that the trace operation in Eq. (65) can be evaluated using any basis of the
764 Hilbert space, such as the ensemble of states fully localized on the A and B sites.

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