

2D topological matter from a boundary Green's functions perspective: Faddeev-LeVerrier algorithm implementation

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1 Abstract

2 Since the breakthrough of twistrionics a plethora of topological phenomena in correlated
3 systems has appeared. These devices can be typically analyzed in terms of lattice mod-
4 els using Green's function techniques. In this work we introduce a general method to
5 obtain the boundary Green's function of such models taking advantage of the numeri-
6 cal Faddeev-LeVerrier algorithm to circumvent some analytical constraints of previous
7 works. We illustrate our formalism analyzing the edge features of a Chern insulator, the
8 Kitaev square lattice model for a topological superconductor and the Checkerboard lat-
9 tice hosting topological flat bands. The efficiency and accuracy of the method is demon-
10 strated by comparison to standard recursive Green's function calculations and direct
11 diagonalizations.

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13 Contents

14	1 Introduction	2
15	2 bGF method for 2D lattice models	4
16	3 Faddeev-LeVerrier algorithm	5
17	4 Tight-binding models	7
18	4.1 Chern insulator	7
19	4.2 2D Kitaev square lattice	9
20	4.3 Flat band Checkerboard lattice	11
21	5 Comparison with recursive approaches	12
22	6 Conclusions and outlook	14
23	A Exact Hamiltonian diagonalization	15
24	B Faddeev-LeVerrier algorithm	15
25	References	16

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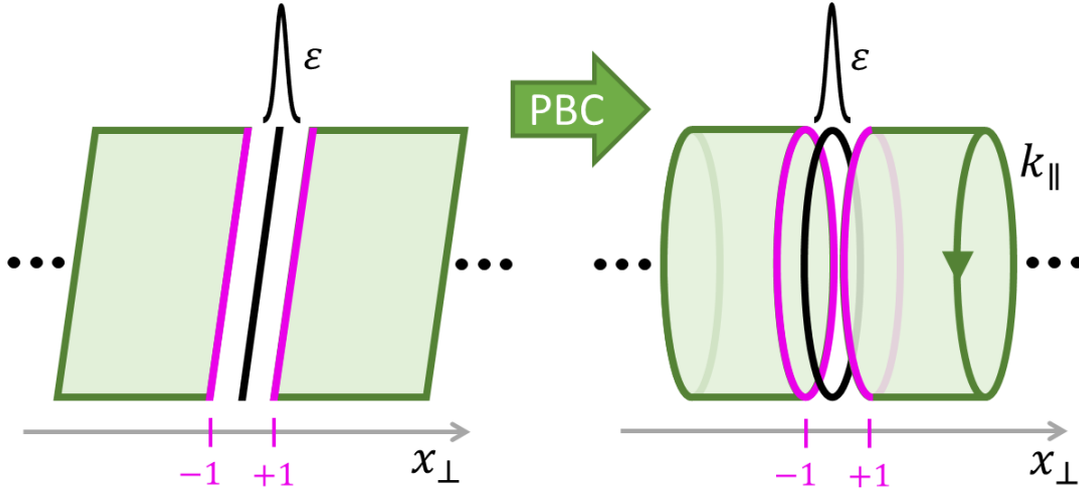


Figure 1: Cylindrical geometry obtained by applying periodic boundary conditions (PBC) along the direction parallel to the boundary in a 2D plane, where x_{\perp} denotes a coordinate in the perpendicular direction measured in units of the lattice constant. In this geometry there is a well defined momentum k_{\parallel} and the open boundaries at $x_{\perp} = \pm 1$ (magenta lines) are obtained by adding a localized impurity line with an amplitude $\varepsilon \rightarrow \infty$ (black line) at $x_{\perp} = 0$. The impurity line breaks the translational symmetry in the x_{\perp} -direction and opens two boundaries in the bulk infinite system.

28 1 Introduction

29 In recent years, due to the appearance of twistrionics [1, 2] and specially since the discovery of
 30 the special properties of twisted bilayer graphene at the magic angle [3, 4], there is a renewed
 31 interest in 2D topological materials exhibiting different phases of matter (e.g. superconductiv-
 32 ity, magnetism, nematicity, etc). In these systems new phenomena arise from the combination
 33 of strong interactions and topology.

34 These circumstances claim for a flexible unified theoretical framework going beyond ide-
 35 alized minimal models to account for interactions, strongly correlated behaviour, spatial inho-
 36 mogeneities or hybrid devices. Several techniques have been developed to analyze open bound-
 37 aries, like exact Hamiltonian diagonalization of finite systems, wave matching in finite scatter-
 38 ing regions [5], some analytical techniques to derive effective boundary Hamiltonians [6] or
 39 the complementary approaches provided by T -matrix and Green's functions formalisms [7–9].
 40 Nevertheless, methods based in exact diagonalization of microscopic Hamiltonians may re-
 41 quire huge computational capabilities with information on several model parameters and gen-
 42 erally, they provide only numerical results with, in some cases, little or no insight in the un-
 43 dergoing physics. For these reasons we are interested in theoretical mesoscopic descriptions
 44 of intermediate complexity which could give us access not only to discrete surface modes but
 45 also to a well defined continuum of excitations.

46 In this work we focus on the boundary Green's function (bGF) method, which is specifi-
 47 cally suited to obtain transport properties in heterostructures based on non-equilibrium Green's
 48 function techniques [10–15]. The bGFs encode information on the local spectral properties
 49 of semi-infinite regions. In addition to transport, such information is of special interest, for
 50 instance, in the case of topological phases where the boundary local density of states (LDOS)s
 51 can reveal the presence of edge states or other type of localized excitations, thus it is possible to
 52 check out the bulk-boundary correspondence of topological phases and computing topological
 53 invariants [16, 17]. Furthermore, Green's function formalism allows to incorporate in a nat-

54 ural way electron-phonon and/or electron-electron interaction effects for example by means
 55 of diagrammatic techniques [18]. Even more, from bGFs it is possible to deduce effective
 56 Hamiltonians including all of these effects and obtain their topological properties [19–21].

57 Here we extend and extrapolate the bGF approach developed in Refs. [13, 22, 23] from 1D
 58 nearest-neighbour (nn) Hamiltonians to generalized d -dimensional systems with an arbitrary
 59 number of degrees of freedom and neighbours. This method performs the Fourier transform
 60 (FT) into real space needed to compute the bGF (see Fig. 1) by the analytic continuation of
 61 the momenta into the complex plane followed by residue integration. This approach exhibits
 62 better convergence performance compared to recursive approaches [24, 25] for which pre-
 63 cision is linked to the number of iterations [23]. However, previous implementations of the
 64 method required analytical expressions for the key building blocks of the formalism such as the
 65 characteristic polynomial. A typical symbolic Laplace expansion to evaluate the characteristic
 66 polynomial is highly inefficient for generalized problems with potentially enormous memory
 67 demand and computational complexity of $O(N!)$ [26] where N is the total Hamiltonian di-
 68 mension. In addition, the other main building block of the method, the adjugate matrix, has
 69 to be obtained with a separate routine.

70 In the present work we complement the method of Refs. [13, 22, 23] with a straightforward
 71 computational approach using the Faddeev-LeVerrier algorithm (FLA) [27–31] that sorts out
 72 diverse disadvantages of the semi-analytical calculations. The FLA requires a low computa-
 73 tional cost to construct not only the characteristic polynomial but also the adjugate matrix in
 74 the same process and for the same price which are, as mentioned before, the main building
 75 blocks to obtain the bGF using the residue integration method. This algorithm is not only
 76 useful for large dimensions but it is also convenient for smaller problems due to its simple im-
 77 plementation. In addition it does not rely in huge analytical expressions for the characteristic
 78 polynomial which arise for $N \geq 3$, thus avoiding possible algebra errors without relevant time
 79 consuming drawbacks.

80 Furthermore, the semi-analytical approach used in Refs. [13, 22, 23] suffer from rigidity in
 81 the definition of the GF as any new terms that might be inserted in the Hamiltonian impose
 82 a redefinition and consequently, all the analytical coefficients of the characteristic polynomial
 83 have to be re-obtained from scratch by time consuming symbolic algorithms. In contrast, the
 84 only analytical entry for the FLA is the polynomial decomposition of the Hamiltonian in the
 85 analytic continuation variable of the momentum perpendicular to the boundary $z = e^{ik_{\perp}L_{\perp}}$.
 86 This is a simpler and flexible analytical requirement that can be computed without any upper
 87 end limit in the number of degrees of freedom of the system.

88 The rest of the paper is organized as follows: in Sec. 2, we describe the computation
 89 of the Green’s function formalism taking advantage of the residue theorem introduced in
 90 Refs. [23, 32]. We then use Dyson’s equation to open a boundary in the bulk system with
 91 an infinity impurity perturbation. Sec. 3, we describe our method based on FLA to compute
 92 the boundary Green’s functions with barely no analytical demands to operate. In Sec. 4, we use
 93 some relevant model Hamiltonians for 2D topological systems as examples to compute steadily
 94 the FLA, first in a purely analytic problem to then jump into purely computational approaches.
 95 These models include the 2D Chern insulator [33], the 2D Kitaev topological superconduc-
 96 tor [34] and the Checkerboard lattice hosting topological flat bands [35]. Furthermore, we
 97 study the spectral properties at edges of such 2D models exhibiting topological features like
 98 chiral edge states. Sec. 5 includes a study of the convergence of the spectral density of the
 99 semi-infinite translational invariant Checkerboard lattice model comparing the recursive GF
 100 technique with the bGF obtained via FLA.

101 We finally summarize the main results with some conclusions in Sec. 6. Technical details
 102 like the finite system diagonalization or an explicit FLA pseudocode are included in the ap-
 103 pendices. Throughout, we use units with nn hopping amplitude $t = 1$ and lattice parameter

104 $a = 1$.

105 2 bGF method for 2D lattice models

106 To obtain the bGF we start from a d -dimensional bulk infinite system and introduce local
 107 perturbations with the characteristic profile that defines the boundary. As these local pertur-
 108 bations or impurity surface amplitude tends to infinity we are left with two $(d-1)$ -dimensional
 109 open surfaces [7, 8] e.g. two boundary lines in a 2D system induced by an impurity line, see
 110 Fig. 1. The bGF is obtained using the Dyson equations associated to the local surface impurity
 111 potential which breaks translational symmetry albeit the momenta in the direction parallel to
 112 the impurity surface are conserved and thus well defined.

113 The starting GF must be explicitly dependent on the local coordinate associated to the
 114 perpendicular direction to the boundary. In order to get these real space GFs, starting from
 115 $N \times N$ tight-binding Hamiltonians in momentum space $\hat{\mathcal{H}}(\mathbf{k})$, we have to compute the FT of the
 116 bulk GF in the direction perpendicular to the boundary. For this purpose, we decompose the
 117 momenta into parallel and perpendicular components $\mathbf{k} = (k_{\parallel}, k_{\perp})$ relative to the boundary
 118 direction (in higher dimensional models the parallel momentum component would be itself
 119 a vector \mathbf{k}_{\parallel}). The bulk Hamiltonian periodicity in both directions is set by $(L_{\parallel}, L_{\perp})$, such that
 120 $\hat{\mathcal{H}}(\mathbf{k} + 2\pi\mathbf{u}_{\perp}/L_{\perp}) = \hat{\mathcal{H}}(\mathbf{k})$, where \mathbf{u}_{\perp} is the unitary vector in the perpendicular direction. As
 121 to compute the FT we need orthogonal lattice vectors, in some cases like the triangular lattice
 122 we have to double the primitive cell. Using this periodicity, the Hamiltonian can be expanded
 123 in a Fourier series, $\hat{\mathcal{H}}(\mathbf{k}) = \sum_n \hat{\mathcal{V}}_n(k_{\parallel})e^{ink_{\perp}L_{\perp}}$, where n covers the number of neighbours, and
 124 Hermiticity implies $\hat{\mathcal{V}}_{-n} = \hat{\mathcal{V}}_n^{\dagger}$. Then, the advanced bulk GF is defined as

$$\hat{G}^A(\mathbf{k}, \omega) = [(\omega - i\eta)\hat{\mathbb{1}} - \hat{\mathcal{H}}(\mathbf{k})]^{-1}, \quad (1)$$

125 where η is a small broadening parameter that ensures the convergence of its analytic properties
 126 [36] (e.g. to compute the spectral densities and integrated quantities). This parameter is
 127 specially needed in the case of recursive methods where the spectrum is approximated by a
 128 finite set of poles. In this work we set $\eta = 2\Delta\omega/n_{\omega}$, where $\Delta\omega$ is the energy window that we
 129 are studying and n_{ω} is the number of points that we are computing within that window. The
 130 $N \times N$ matrix structure is indicated by the hat notation.

131 Fourier transforming along the perpendicular direction, the GF components are given by

$$\hat{G}_{jj'}^A(k_{\parallel}, \omega) = \frac{L_{\perp}}{2\pi} \int_{-\pi/L_{\perp}}^{\pi/L_{\perp}} dk_{\perp} e^{i(j-j')k_{\perp}L_{\perp}} \hat{G}^A(k_{\parallel}, k_{\perp}, \omega), \quad (2)$$

132 where j and j' are lattice site indices in the x_{\perp} -direction. By the identification $z = e^{ik_{\perp}L_{\perp}}$, this
 133 integral is converted into a complex contour integral,

$$\hat{G}_{jj'}^A(k_{\parallel}, \omega) = \frac{1}{2\pi i} \oint_{|z|=1} \frac{dz}{z} z^{j-j'} \hat{G}^A(k_{\parallel}, z, \omega). \quad (3)$$

134 Further simplification can be obtained by introducing the roots $z_n(k_{\parallel}, \omega)$ of the character-
 135 istic polynomial in the z -complex plane,

$$P(k_{\parallel}, z, \omega) = \det[\omega\hat{\mathbb{1}} - \hat{\mathcal{H}}(k_{\parallel}, z)] = \frac{c_m}{z^m} \prod_{n=1}^{2m} (z - z_n(k_{\parallel}, \omega)), \quad (4)$$

136 where m is the highest order of the characteristic polynomial and c_m is the highest order
 137 coefficient. In terms of these roots the contour integral in Eq. (3) can be written as a sum over
 138 the residues of all roots inside the unit circle in the complex plane

$$\hat{G}_{jj'}^A(k_{\parallel}, \omega) = \sum_{|z_n| < 1} \frac{z_n^q \hat{M}(k_{\parallel}, z_n, \omega)}{c_m \prod_{l \neq n} (z_n - z_l)}, \quad (5)$$

139 where $q = j - j' + m - m' - 1$ and $z^{-m'} \hat{M}(k_{\parallel}, z, \omega)$ is the adjugate matrix of $[\omega \hat{\mathbb{1}} - \hat{\mathcal{H}}(k_{\parallel}, z)]$
 140 where all the poles at zero were taken out of \hat{M} as a common factor in $z^{-m'}$. Finally, \sum' means
 141 that if $q < 0$ then we include $z_n = 0$ as a pole in the sum of residues (e.g., in the non local GF
 142 components with $j' > j$). Consequently when $q < -1$ higher order poles at zero appear in the
 143 sum of residues. To simplify these situations we can take advantage of the residue theorem to
 144 avoid these poles and compute the integral as

$$\hat{G}_{jj'}^A(k_{\parallel}, \omega) = - \sum_{|z_n| > 1} \frac{z_n^q \hat{M}(k_{\parallel}, z_n, \omega)}{c_m \prod_{l \neq n} (z_n - z_l)}. \quad (6)$$

145 To simplify the notation, we omit the superscript 'A' denoting advanced GFs from now
 146 on. Given the real-space components of the bulk GF in Eq. (5), we next extend the method of
 147 Refs. [11, 13, 37] to derive the bGF characterizing a *semi-infinite* nearest-neighbour 2D systems.
 148 To this effect, we add an impurity potential line ε localized at the frontier region. Taking the
 149 limit $\varepsilon \rightarrow \infty$ the infinite system is cut into two disconnected semi-infinite subsystems with
 150 $j \leq -1$ (left side, L) and $j \geq 1$ (right side, R), see Fig. 1. Using Dyson equation the local GF
 151 components of the cut subsystem follow as [13]

$$\hat{\mathcal{G}}_{jj} = \hat{\mathcal{G}}_{jj}^{(0)} - \hat{\mathcal{G}}_{j0}^{(0)} \left[\hat{\mathcal{G}}_{00}^{(0)} \right]^{-1} \hat{\mathcal{G}}_{0j}^{(0)}, \quad (7)$$

152 where $\hat{\mathcal{G}}^{(0)}$ are the unperturbed bulk GF and $\hat{\mathcal{G}}$ are the semi-infinite perturbed GF. Following
 153 Eq. (7), the bGF for the left and right semi-infinite systems are respectively given by

$$\hat{\mathcal{G}}_L(k_{\parallel}, \omega) = \hat{\mathcal{G}}_{\bar{1}\bar{1}}(k_{\parallel}, \omega), \quad \hat{\mathcal{G}}_R(k_{\parallel}, \omega) = \hat{\mathcal{G}}_{11}(k_{\parallel}, \omega), \quad (8)$$

154 where the over-line in the local indices in the bGF means negative sites. In order to extend the
 155 method to the case of interactions to an arbitrary number of neighbours we need to include
 156 as many impurity surfaces as the number of neighbours in order to completely disconnect
 157 two semi-infinite regions. In an equivalent fashion we could obtain a new Dyson equation
 158 to compute the bGF of the system. Another possibility would be to define a supercell that
 159 transforms the Hamiltonian into a nearest-neighbour one with respect to this supercell and
 160 compute the bGF applying Eq. (8). Using this bGF we can compute the spectral properties of
 161 open (semi-infinite or finite) systems encoded in the spectral densities and the local density of
 162 states respectively

$$\rho_{L,R}(k_{\parallel}, \omega) = \frac{1}{\pi} \Im \text{tr} \{ \hat{\mathcal{G}}_{L,R}(k_{\parallel}, \omega) \}, \quad \langle \rho_{L,R}(\omega) \rangle = \int \frac{dk_{\parallel}}{\Omega_{k_{\parallel}}} \rho_{L,R}(k_{\parallel}, \omega), \quad (9)$$

163 where $\Omega_{k_{\parallel}} = 2\pi/L_{\parallel}$ accounts for the limits of integration.

164 3 Faddeev-LeVerrier algorithm

165 We first summarize FLA for a generic complex matrix. Let \hat{A} be a $N \times N$ matrix with char-
 166 acteristic polynomial $P(\omega) = \det[\omega \hat{\mathbb{1}} - \hat{A}] = \sum_{k=0}^n \bar{C}_k \omega^k$. The trivial coefficients are $\bar{C}_n = 1$

bGF COMPUTATION

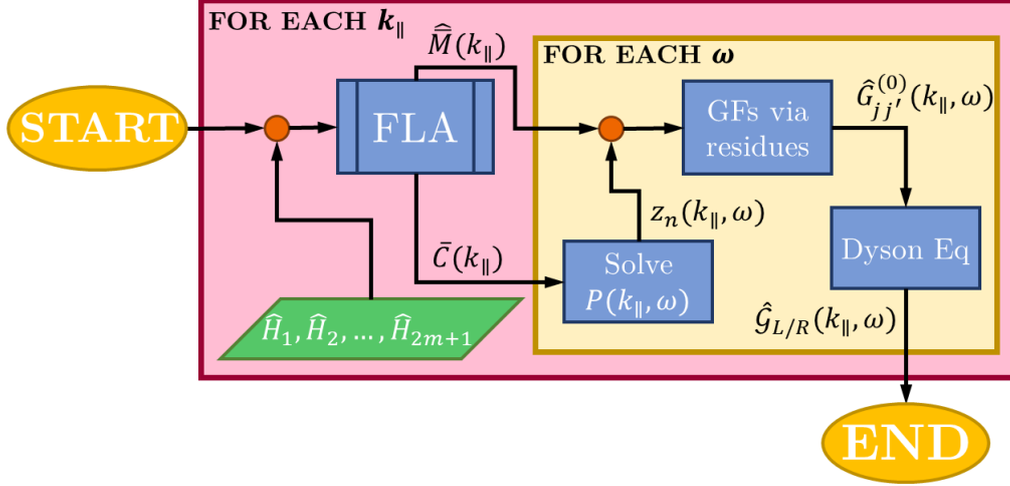


Figure 2: Complete algorithm workflow to compute the bGF using the FLA where the main input is the polynomial decomposition of the Hamiltonian for a given set of momenta k_{\parallel} and frequencies ω .

167 and $\bar{C}_0 = (-1)^n \det \hat{A}$, also simple is the term $\bar{C}_{n-1} = -\text{tr}\{\hat{A}\}$. The other coefficients can be
 168 calculated using the Faddeev-LeVerrier algorithm [27–31] as

$$\hat{M}_k = \hat{A}\hat{M}_{k-1} + \bar{C}_{n-k+1}\hat{\mathbb{1}}, \quad \bar{C}_{n-k} = -\frac{1}{k}\text{tr}\{\hat{A}\hat{M}_k\}, \quad (10)$$

169 where \hat{M}_k is an auxiliary matrix such that $\hat{M}_0 = 0$. Remarkably the matrices \hat{M}_k allow us to
 170 obtain the adjugate matrix of $[\omega\hat{\mathbb{1}} - \hat{A}]$ as a polynomial

$$\text{adj}[\omega\hat{\mathbb{1}} - \hat{A}] = \sum_{k=0}^n \omega^k \hat{M}_{n-k}, \quad (11)$$

171 which, given that $\hat{M}_0 = \hat{0}$, the adjugate matrix $\hat{M}(\omega)$ has $N - 1$ order in ω .

172 In our case $\hat{A} \equiv \hat{\mathcal{H}}(z)$ is a polynomial complex matrix and it can also be expanded as a
 173 polynomial in z as

$$\hat{\mathcal{H}}(z) = \sum_{i=1}^{2m+1} \hat{H}_i z^{i-(m+1)} = \hat{H}_1 z^{-m} + \dots + \hat{H}_{m+1} + \dots + \hat{H}_{2m+1} z^m. \quad (12)$$

174 In some simple cases where $\text{rg}(\hat{H}_{2m+1}) = N$ we get the highest order polynomial decom-
 175 position for the Hamiltonian and $m = n_n N$ where n_n is equal to the number of neighbours in
 176 the tight-binding model, but in general $m \leq n_n N$.

177 We are interested in expressing \bar{C}_k and \hat{M}_k as two-variable polynomials in ω and z using
 178 two variable FLA [38–40] to compute the complex integral. Still we have $\bar{C}_n = 1$ and $\hat{M}_1 = \hat{\mathbb{1}}$.
 179 Then,

$$\bar{C}_{n-1}(z) = -\text{tr}\{\hat{\mathcal{H}}(z)\} = \sum_{i=1}^{2m+1} \bar{C}_{n-1,i} z^{i-(m+1)}. \quad (13)$$

180 For example, the next coefficients are

$$\hat{M}_2(z) = \hat{\mathcal{H}}(z) + \bar{C}_{n-1}\hat{\mathbb{1}} = \sum_{i=1}^{2m+1} \hat{M}_{2,i} z^{i-(m+1)}, \quad \bar{C}_{n-2}(z) = -\frac{1}{2}\text{tr}\{\hat{\mathcal{H}}(z)\hat{M}_2(z)\} = \sum_{i=1}^{4m+1} \bar{C}_{n-2,i} z^{i-(2m+1)}. \quad (14)$$

181 In this way we could get

$$\hat{M}_k(z) = \sum_{i=1}^{2m(k-1)+1} \hat{M}_{k,i} z^{i-(m(k-1)+1)}, \quad \bar{C}_{n-k}(z) = \sum_{i=i}^{2mk+1} \bar{C}_{n-k,i} z^{i-(mk+1)}, \quad (15)$$

182 and deduce an explicit decomposition of $\text{adj}[\omega \hat{\mathbb{1}} - \hat{\mathcal{H}}(k_{\parallel}, z)]$ in z from which we can extract
 183 the zero poles of the adjugate matrix $z^{-m'}$ as in Eq. (5). In simple cases where $m = n_n N$, it is
 184 straightforward to see that $m' = N - 1$.

185 In Fig. 2 we expose the general structure of the complete algorithm to compute the bGF
 186 given, as an input, the polynomial decomposition of the Hamiltonian particularized at any
 187 k_{\parallel} . Using FLA we obtain the auxiliary matrix to compute the adjugate of the secular equation
 188 $\hat{M}(k_{\parallel})$ and the coefficients of the characteristic polynomial $\bar{C}(k_{\parallel})$, see Appendix B. From $\bar{C}(k_{\parallel})$
 189 we can compute the characteristic polynomial $P(k_{\parallel}, \omega)$ for any desired frequency and solve it
 190 to obtain the roots $z_n(k_{\parallel}, \omega)$.

191 Both $z_n(k_{\parallel}, \omega)$ and $\hat{M}(k_{\parallel})$ are the key ingredients to compute the unperturbed GFs in real
 192 space using Eq. (6) and taking as poles the roots that satisfy that $|z_n(k_{\parallel}, \omega)| > 1$. The order of
 193 the zero poles m and m' are totally determined by the polynomial decomposition in z of $\bar{C}(k_{\parallel})$
 194 and $\hat{M}(k_{\parallel})$ respectively. Finally, we use Dyson equation to compute the bGFs of the system
 195 from the unperturbed ones.

196 4 Tight-binding models

197 In order to illustrate our method in a transparent self-explanatory way we take the example
 198 of common, well-known 2D topological Hamiltonians to compute the bGF explicitly. First, we
 199 start with the fully analytical 2×2 Chern insulator model [33] hosting chiral edge states to
 200 easily follow the FLA step by step. Later we consider more intricate examples where we have
 201 to partially or totally take advantage of the computational power of the FLA. These models
 202 include the 2D Kitaev model [34] for a topological superconductor showing Majorana edge
 203 modes and the 2D Checkerboard model which hosts topological flat bands with chiral edge
 204 states [35]. All these examples are relevant models for the study of topological matter in 2D
 205 and thus we exhibit the spectral density and the LDOS for an open boundary semi-infinite
 206 system to make explicit their topological edge properties. In Fig. 3 a) we show the Brillouin
 207 zone (BZ) for all of these different lattice models.

208 4.1 Chern insulator

209 We first illustrate the FLA with the well-known 2×2 Chern insulator Hamiltonian [33] in a
 210 square lattice described by

$$\hat{\mathcal{H}}(\mathbf{k}) = (M - \cos k_y - \cos k_x) \sigma_z + \sin k_x \sigma_x + \sin k_y \sigma_y, \quad (16)$$

211 where σ_{μ} with $\mu = x, y, z$ are the Pauli matrices and M is the mass term.

212 We FT along $k_x = k_{\perp}$ thereby we made the analytic continuation $z = e^{ik_x}$. We can now
 213 obtain the polynomial expansion of the Hamiltonian in z following Eq. (12) where

$$\hat{H}_1 = \hat{H}_3^{\dagger} = (i\sigma_x - \sigma_z)/2, \quad \hat{H}_2 = (M - \cos k_y) \sigma_z + \sin k_y \sigma_y. \quad (17)$$

214 We then compute the trivial \bar{C}_n coefficients that define the characteristic polynomial in
 215 frequencies (ω)

$$\bar{C}_2 = 1, \quad \bar{C}_1 = 0, \quad \bar{C}_0 = (M - \cos k_y)(z + z^{-1}) - [M^2 + 2(1 - M \cos k_y)], \quad (18)$$

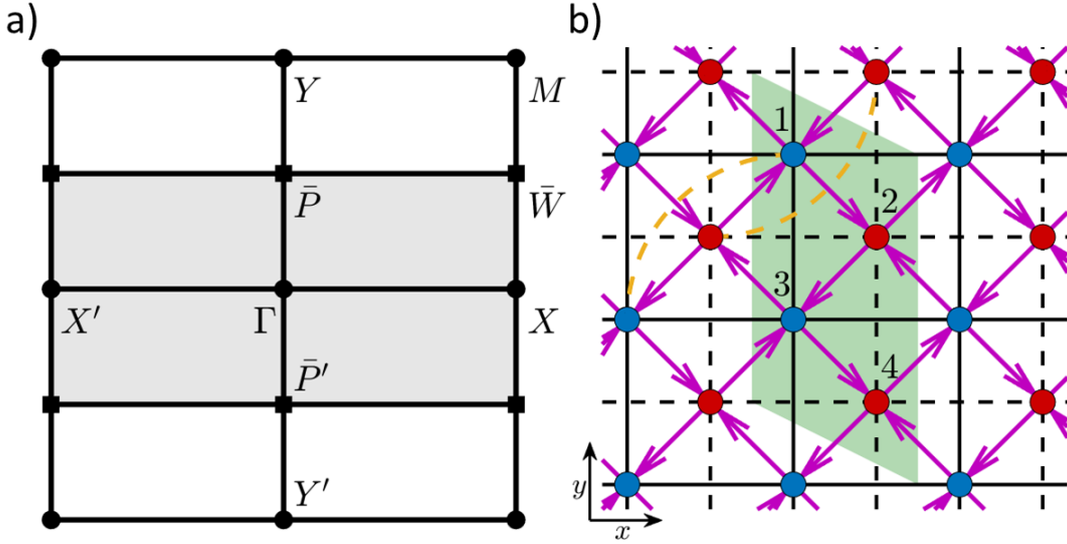


Figure 3: Brillouin zone for the square lattice models and real space representation of the Checkerboard lattice model. a) Square (white) and rectangular (grey shaded) BZ showing the high symmetry points in each one. The over-line in the high symmetry points denotes that they belong to the folded rectangular BZ in the k_y -direction. b) Checkerboard lattice. Red and blue dots indicate the sublattice sites. The magenta arrow, black dashed (solid) line and yellow dashed line accounts for the nn hopping t , the nnn hopping t'_1 (t'_2) and the $nnnn$ hopping t'' respectively. The arrow direction shows the sign of the accumulated phase ϕ in the nn hopping terms. The shaded green region in b) corresponds to the doubling of the original primitive cell which produces the folding of the square BZ into a rectangular one as indicated in panel a).

216 consequently, their explicit decomposition in the z polynomial

$$\bar{C}_{02} = \bar{C}_{04}^* = (M - \cos k_y), \quad \bar{C}_{03} = -[M^2 + 2(1 - M \cos k_y)]. \quad (19)$$

217 Due to the aforementioned relation, as $\text{rg}(\hat{H}_{2m+1}) < N$, then $m < n_n N$ and for that we
 218 have a reduced degree of the characteristic polynomial obeying $\bar{C}_{01} = \bar{C}_{05} = 0$. Nevertheless,
 219 we have used the indexation of the polynomial in z as the maximum degree polynomial for
 220 the sake of generalization of the method, similarly to the criteria taken in the pseudocode
 221 formulation in Appendix B.

222 Then, the characteristic polynomial takes the form

$$P(\omega) = (M - \cos k_y)(z + z^{-1}) + \omega^2 - M^2 - 2(1 - M \cos k_y), \quad (20)$$

223 where $c_m = \bar{C}_{04}$ and the non-trivial contributions to \hat{M} matrix are defined by

$$\hat{M}_{21} = \hat{H}_1 + c_{11} \hat{\mathbb{I}} = \hat{H}_1, \quad \hat{M}_{22} = \hat{H}_2 + c_{12} \hat{\mathbb{I}} = \hat{H}_2, \quad \hat{M}_{23} = \hat{H}_3 + c_{13} \hat{\mathbb{I}} = \hat{H}_3. \quad (21)$$

224 Finally, the integral by residues for the bulk GF takes the form

$$\hat{G}_{jj'}(k_y, \omega) = -\frac{z_-^{j-j'}}{z_-} \frac{\begin{pmatrix} -(1+z_-^2) + \alpha z_- & i(1-z_-^2) - \beta z_- \\ i(1-z_-^2) + \beta z_- & (1+z_-^2) - \alpha z_- \end{pmatrix}}{2(M - \cos k_y)(z_- - z_+)}, \quad (22)$$

225 where $\alpha = 2(M + \omega - \cos k_y)$, $\beta = i2 \sin k_y$ and we have regularized the zeros of the ad-
 226 jugate matrix $\text{adj}[\omega \hat{\mathbb{I}} - \hat{H}(k_{\parallel}, z)]$ with the zeros of $P(\omega)$ knowing that $m = m' = 1$. Fur-
 227 thermore, we solve the trivial roots for $P(\omega)$ in Eq. (20), $z_{\pm} = (-b \pm \sqrt{b^2 - 4})/2$ where

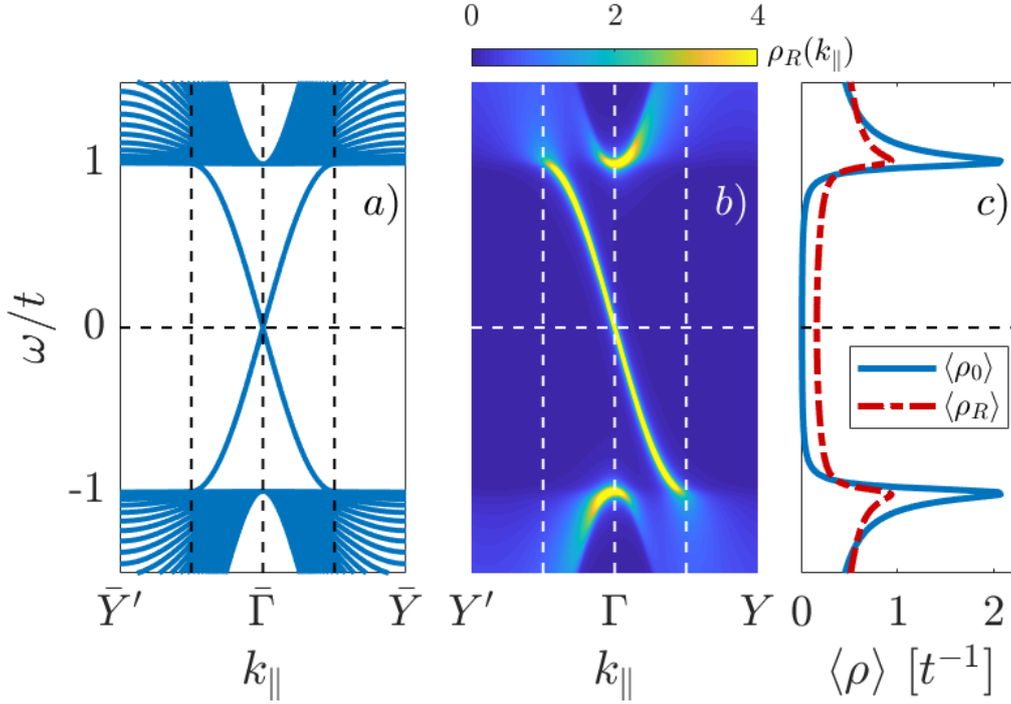


Figure 4: Open boundary characterization for the Chern Insulator model showing chiral edge states under the effect of the mass term $M \rightarrow 1$. a) Electronic bands obtained by exact diagonalization of a finite size system with $N_{sites} = 40$ sites. The spectrum shows 2 chiral edge states each one associated to a different boundary. b) Spectral density for a right boundary in the semi-infinite limit obtained from the bGF calculation. c) Integrated LDOS where straight (dot-dashed) line represents bulk (right boundary) LDOS.

228 $b = [\omega^2 - M^2 - 2(1 - M \cos k_y)] / (M - \cos k_y)$ defining $|z_-| > 1$ and $|z_+| < 1$. Once the
 229 bulk GF in real space has been constructed we use Eq. (8) to obtain the corresponding bGFs.
 230 In Fig. 4 we illustrate the open boundary spectral density for the topological phase of the Chern
 231 insulator exhibiting chiral edge states obtained using FLA. For comparison we also show the
 232 bands obtained using exact finite size Hamiltonian diagonalization, see Appendix A. As can be
 233 observed, while two chiral edge states are present in the finite system calculation, only one
 234 appears in the bGF calculation as expected for a semi-infinite system.

235 4.2 2D Kitaev square lattice

236 Now we apply FLA to obtain the characteristic polynomial of the 2×2 Kitaev square lattice
 237 model [34] and solve it computationally, in this way we can then obtain the bGF in a semi-
 238 analytic manner. The model Hamiltonian is given by

$$\hat{H}(\mathbf{k}) = (\mu - \cos k_y - \cos k_x) \sigma_z - \Delta (\sin k_x + \sin k_y) \sigma_y, \quad (23)$$

239 where μ is the chemical potential and Δ is the pairing potential.

240 Again, the FT along $k_x = k_{\perp}$ is obtained using the analytic continuation $z = e^{ik_x}$. The
 241 polynomial expansion of the Hamiltonian in z takes the expression

$$\hat{H}_1 = \hat{H}_3^{\dagger} = (-\sigma_z - i\Delta\sigma_y)/2, \quad \hat{H}_2 = (\mu - \cos k_y) \sigma_z + -\Delta \sin k_y \sigma_y. \quad (24)$$

242 We next compute the \bar{C}_n coefficients that define the characteristic polynomial in powers of

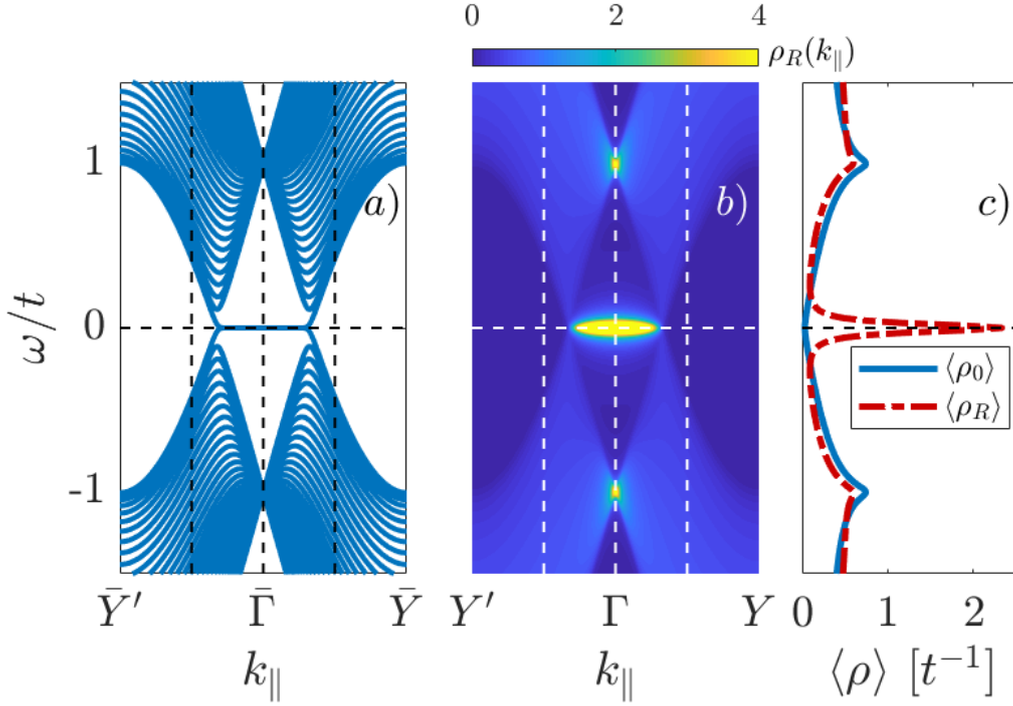


Figure 5: Open boundary characterization for the 2D Kitaev model showing Majorana flat band edge modes in the topological phase $\Delta = 1$ and $\mu = 1$. a) Electronic bands obtained by exact diagonalization of a finite size system with $N_{sites} = 40$ sites. The spectrum shows flat bands at both ends of the system. b) Spectral density for a right boundary in the semi-infinite limit obtained from the bGF calculation. c) Integrated LDOS where straight (dot-dashed) line represents bulk (right boundary) LDOS.

243 ω and z

$$\begin{aligned} \bar{C}_2 = 1, \quad \bar{C}_1 = 0, \quad \bar{C}_{01} = \bar{C}_{05}^* = (\Delta^2 - 1)/4, \quad \bar{C}_{02} = \bar{C}_{04}^* = (\mu - \cos k_y) - i\Delta^2 \sin k_y, \\ \bar{C}_{03} = \frac{(\Delta^2 - 1)}{2} \cos 2k_y + 2\mu \cos k_y - (1 + \Delta^2 + \mu^2), \end{aligned} \quad (25)$$

244 where $c_m = \bar{C}_{05}$ and finally the non-trivial contributions to the \hat{M} matrix are defined as

$$\hat{M}_{21} = \hat{H}_1, \quad \hat{M}_{22} = \hat{H}_2, \quad \hat{M}_{23} = \hat{H}_3. \quad (26)$$

245 We regularize the zeros of the $\text{adj}[\omega \hat{\mathbb{L}} - \hat{\mathcal{H}}(k_{\parallel}, z)]$ with the zeros of $P(\omega)$ knowing that
246 $m = 2$ and $m' = 1$. The integral by residues for the bulk GF takes the form

$$\hat{G}_{jj'}(k_y, \omega) = -2z_4^{j-j'} \frac{\begin{pmatrix} -(1+z_4^2) + \alpha z_4 & -\Delta[(1-z_4^2) - \beta z_4] \\ \Delta[(1-z_4^2) - \beta z_4] & (1+z_4^2) - \alpha z_4 \end{pmatrix}}{(\Delta^2 - 1)(z_4 - z_1)(z_4 - z_2)(z_4 - z_3)} + (z_4 \longleftrightarrow z_3), \quad (27)$$

247 where $\alpha = 2(\mu + \omega - \cos k_y)$, $\beta = i2 \sin k_y$ and $|z_4|, |z_3| > 1$, thus $|z_2|, |z_1| < 1$. We omit the
248 explicit analytical expression of the roots of the characteristic 4th degree polynomial due to
249 their extension. As mentioned before, for this example it is convenient to obtain the roots com-
250 putationally. In Fig. 5 we show typical results for the open boundary LDOS in the topological
251 phase of the 2D Kitaev model showing Majorana flat band edge modes. Again, the comparison
252 with the finite size diagonalization shows good agreement.

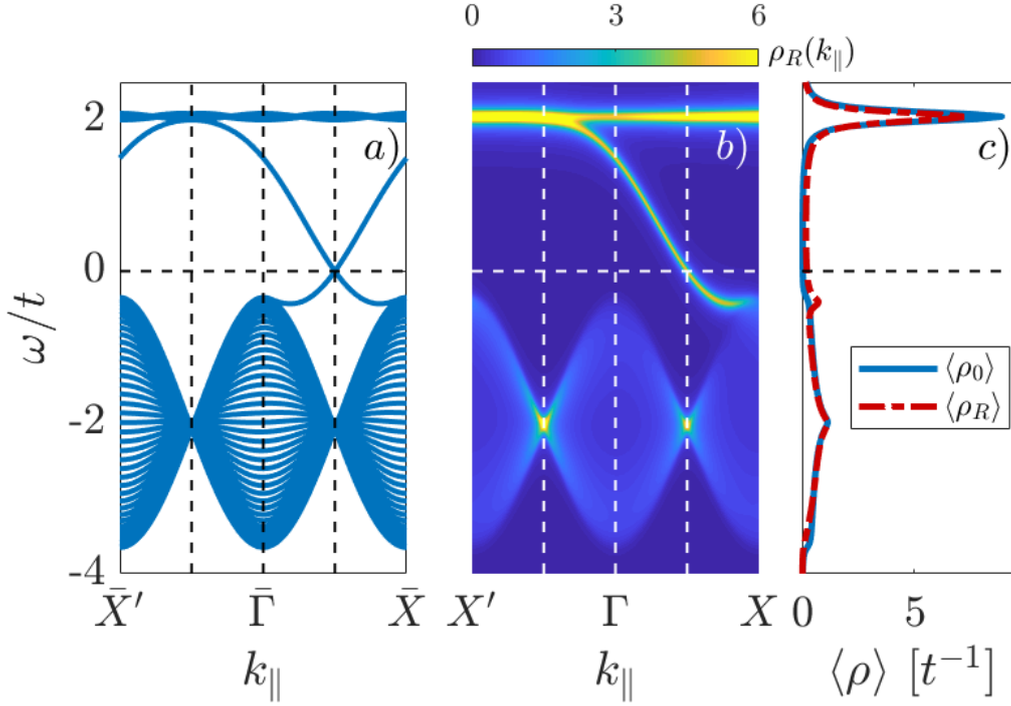


Figure 6: Open boundary characterization for the Checkerboard lattice model showing topological flat band at $\omega/t = 2$ with a chiral edge mode in the topological phase $\phi = -\pi/4$, $t'_1 = -t'_2 = t/(2 + \sqrt{2})$ and $t'' = -t/(2 + 2\sqrt{2})$. a) Electronic bands obtained by exact diagonalization of a finite size system with $N_{sites} = 20$ sites with the unit cell doubled. The spectrum shows 2 chiral edge states each one associated to a different boundary. b) Spectral density for a right boundary in the semi-infinite limit obtained from the bGF calculation. c) Integrated LDOS where straight (dot-dashed) line represents bulk (right boundary) LDOS.

253 4.3 Flat band Checkerboard lattice

254 Finally we consider the 2×2 Checkerboard lattice model [35] which hosts topological flat
 255 bands and is defined by the Hamiltonian

$$\hat{\mathcal{H}}(\mathbf{k}) = \Omega_0(\mathbf{k})\hat{1} + \Omega_1(\mathbf{k})\sigma_x + \Omega_2(\mathbf{k})\sigma_y + \Omega_3(\mathbf{k})\sigma_z, \quad (28)$$

256 where

$$\begin{aligned} \Omega_0(\mathbf{k}) &= (t'_1 + t'_2)(\cos k_x + \cos k_y) + 4t'' \cos k_x \cos k_y, & \Omega_1(\mathbf{k}) &= 4t \cos \phi \cos \frac{k_x}{2} \cos \frac{k_y}{2}, \\ \Omega_2(\mathbf{k}) &= 4t \sin \phi \sin \frac{k_x}{2} \sin \frac{k_y}{2}, & \Omega_3(\mathbf{k}) &= (t'_1 - t'_2)(\cos k_x - \cos k_y). \end{aligned} \quad (29)$$

257 The system is thus characterized by nn hopping t , nnn hopping t'_1, t'_2 and $nnnn$ hopping
 258 t'' terms, also the nn terms accumulate a phase ϕ pointed out in Fig. 3 b).

259 This model is an exemplification of a typical obstacle to tackle with our algorithm due to the
 260 sublattice degree of freedom. Due to that, the Hamiltonian includes lattice spacing fractions,
 261 hence if we try to FT with the analytic continuation $z = e^{ik_{\perp}L_{\perp}/2}$ instead of having a complex
 262 integral over the closed unit circle we arrive to an open arc integral in the complex plane, so
 263 we cannot apply the residue theorem to solve it. This kind of problems may also appear in
 264 Bravais lattices with non-orthogonal lattice vectors (e.g. the triangular lattice).

265 To circumvent this kind of obstacles we proceed to double the unit cell to obtain a new
 266 lattice with orthogonal lattice vectors and integer powers of $z = e^{ik_\perp L_\perp}$. The drawbacks of
 267 doubling the unit cell are that we are now working in a folded BZ and we have doubled the
 268 Hamiltonian degrees of freedom. Consequently the Hamiltonian in the new unit cell expressed
 269 in the basis $\Psi_{\mathbf{k}} = (\psi_{A1,\mathbf{k}}, \psi_{B2,\mathbf{k}}, \psi_{A3,\mathbf{k}}, \psi_{B4,\mathbf{k}})^T$ takes the form

$$\hat{\mathcal{H}}(\mathbf{k}) = \begin{pmatrix} \hat{A} & \hat{B} \\ \hat{B}^\dagger & \hat{A} \end{pmatrix}, \quad (30)$$

270 with

$$\hat{A} = \begin{pmatrix} \delta_2 & \beta_- \\ \beta_-^* & \delta_1 \end{pmatrix}, \quad \hat{B} = \begin{pmatrix} \alpha_1(1 + e^{ik_y}) & e^{ik_y} \beta_+^* \\ \beta_+ & \alpha_2(1 + e^{ik_y}) \end{pmatrix}, \quad (31)$$

271 where $\beta_\pm = e^{\pm i(k_x \pm \phi)} + e^{-i\phi}$, $\alpha_\mu = (t'_\mu + 2t'' \cos k_x)$ and $\delta_\mu = 2t'_\mu \cos k_x$ with $\mu = 1, 2$.

272 In Fig. 3 b) we show the unit cell doubling in the y -direction for the Checkerboard lattice
 273 problem leading to a folded BZ along the k_y -direction. To avoid foldings in the spectral
 274 densities we have made the analytic continuation in $z = e^{ik_y}$ with $k_y = k_\perp$, in this way we
 275 have the explicit momenta dependence of the Hamiltonian in the unfolded BZ coordinate
 276 $k_x = k_\parallel$. The polynomial expansion of the Hamiltonian in z adopts the expression

$$\hat{H}_1 = \hat{H}_3^\dagger = \begin{pmatrix} 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ \alpha_1 & 0 & 0 & 0 \\ \beta_+ & \alpha_2 & 0 & 0 \end{pmatrix}, \quad \hat{H}_2 = \begin{pmatrix} \delta_2 & \beta_- & \alpha_1 & 0 \\ \beta_-^* & \delta_1 & \beta_+ & \alpha_2 \\ \alpha_1 & \beta_+^* & \delta_2 & \beta_- \\ 0 & \alpha_2 & \beta_-^* & \delta_1 \end{pmatrix}. \quad (32)$$

277 Due to the cell doubling we have a characteristic off-diagonal representation of the z -
 278 dependent terms of the Hamiltonian which induces that $\text{rg}(\hat{H}_{2m+1}) < N$, then again we have a
 279 degree reduction of the characteristic polynomial. We now could obtain analytically the \bar{C}_n co-
 280 efficients that define the characteristic polynomial but we omitted them due to their extension.
 281 These coefficients along with the adjugate matrix $\hat{M}(k_\parallel, z, \omega)$ can be obtained computationally
 282 in a straightforward way using Eq. (14), see Appendix B.

283 In Fig. 6 we show results for the open boundary LDOS for the topological phase of the
 284 Checkerboard lattice model exhibiting topological flat bands and chiral edge states. Again,
 285 the comparison with the bands obtained by direct diagonalization gives excellent agreement,
 286 except for the doubling of the edge states.

287 5 Comparison with recursive approaches

288 As mentioned before, the recursive GF method is a well established tool to compute bGFs.
 289 Below we briefly describe the recursive method taking advantage of the Hamiltonian decom-
 290 position into two perpendicular directions already introduced for FLA. We define the recursive
 291 method to compute the bGF at a dimensionless n -site as

$$[\hat{\mathcal{G}}_R^{rc}(n)]^{-1} = \omega \hat{\mathbb{1}} - \hat{\mathcal{H}}_0(k_\parallel) - \Sigma_R(n), \quad [\hat{\mathcal{G}}_L^{rc}(n)]^{-1} = \omega \hat{\mathbb{1}} - \hat{\mathcal{H}}_0(k_\parallel) - \Sigma_L(n), \quad (33)$$

292 where $\hat{\mathcal{H}}_0(k_\parallel)$ is the local contribution defined in each iteration step and the recursive expres-
 293 sion of the self-energy takes the form

$$\Sigma_R(n) = \hat{T}_{LR} [\hat{\mathcal{G}}_R^{rc}(n-1)]^{-1} \hat{T}_{LR}^\dagger, \quad \Sigma_L(n) = \hat{T}_{LR}^\dagger [\hat{\mathcal{G}}_L^{rc}(n-1)]^{-1} \hat{T}_{LR}. \quad (34)$$

294 As can be observed, the self-energy at a given n -site couples this site with the previous one
 295 where n goes from $n = 1$ to $n = N_{it}$ with N_{it} is the number of recursive steps. The self-energy

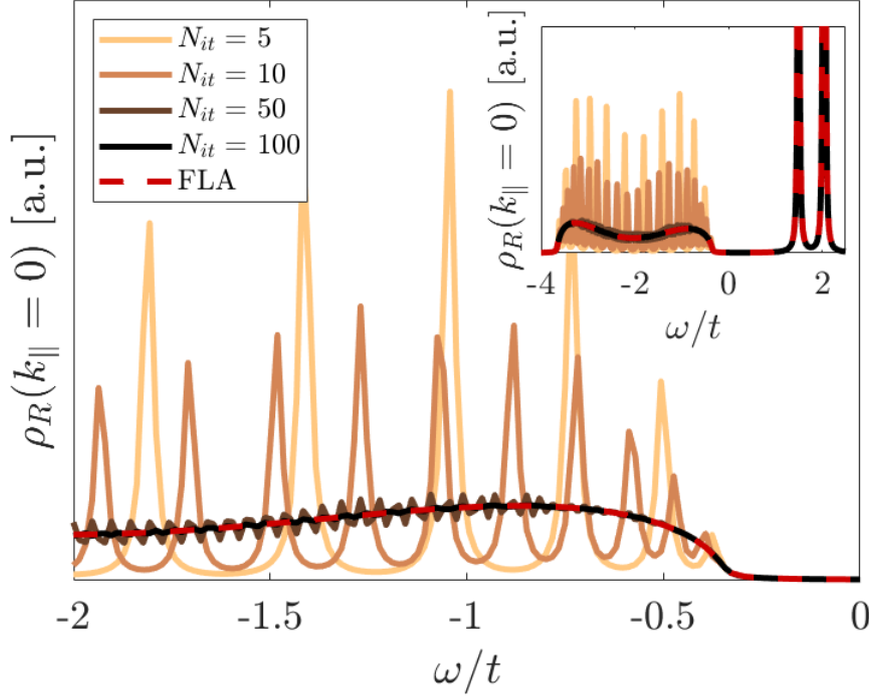


Figure 7: Open right boundary spectral density at $k_{\parallel} = \Gamma$ for the Checkerboard lattice model with $\eta = 0.02$ and the rest of parameters are the same as in Fig. 6. Solid lines represents the spectral density obtained by recursive GF for different number of recursive steps $N_{it} = 5, 10, 50$. Dashed red line is obtained using FLA. Main figure: top continuum valence bands contribution to the spectral density showing the discretization effect of the recursive method. Inset: All the contributions to the spectral density including the flat band at $\omega/t = 2$ and the topological chiral edge state at $\omega/t \approx 1.5$

296 at the first site $\Sigma_{L/R}(n = 1)$ can be defined to simulate the coupling to a doped continuum of
 297 the same material for better convergence.

298 From the polynomial decomposition of the Hamiltonian in Eq. (12) we can define the
 299 relevant matrices for the recursive method as

$$\hat{\mathcal{H}}_0 = \begin{pmatrix} \hat{H}_{m+1} & \hat{H}_m & \hat{H}_{m-1} & \cdots & \hat{H}_2 \\ \hat{H}_m^\dagger & \hat{H}_{m+1} & \hat{H}_m & \cdots & \hat{H}_3 \\ \hat{H}_{m-1}^\dagger & \hat{H}_m^\dagger & \hat{H}_{m+1} & \cdots & \hat{H}_4 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \hat{H}_2^\dagger & \hat{H}_3^\dagger & \hat{H}_4^\dagger & \cdots & \hat{H}_{m+1} \end{pmatrix}, \quad \hat{T}_{LR} = \begin{pmatrix} \hat{H}_1^\dagger & \hat{H}_2^\dagger & \hat{H}_3^\dagger & \cdots & \hat{H}_m^\dagger \\ \hat{0} & \hat{H}_1^\dagger & \hat{H}_2^\dagger & \cdots & \hat{H}_{m-1}^\dagger \\ \hat{0} & \hat{0} & \hat{H}_1^\dagger & \cdots & \hat{H}_{m-2}^\dagger \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \hat{0} & \hat{0} & \hat{0} & \cdots & \hat{H}_1^\dagger \end{pmatrix}. \quad (35)$$

300 Notice that the dimension of the recursive method goes as $N_r = Nn_n$ so for the usual nm
 301 case satisfies $N_r = N$ and $\hat{\mathcal{H}}_0 = \hat{H}_2$, $\hat{T}_{LR} = \hat{H}_1^\dagger$.

302 In Fig. 7 we illustrate the convergence of the continuum spectrum within the recursive GF
 303 method for the Checkerboard model at $k_{\parallel} = \Gamma$ with parameters as in Fig. 6 for several number
 304 of iterations compared to bGF obtained using FLA. While the recursive approach accounts well
 305 for discrete states, as boundary states, with few iterations, the number of recursive steps have
 306 to be greatly increased to properly converge the continuum spectrum into the semi-infinite
 307 limit [36]. In contrast, FLA provides an accurate description of both surface modes and con-
 308 tinuum spectra without further computational effort. It is worth mentioning that the recursive

309 method for all the lattice models in this publication takes from twice to four times more com-
 310 puting time than FLA for the same number of points in the spectral density and $N_{it} = 100$,
 311 for which, as shown in Fig. 7, the recursive calculation has not yet converged to a smooth
 312 continuum spectrum.

313 In order to compare the computational complexity of our technique one should have in
 314 mind that our method could be implemented in a partially analytical way, in the sense that
 315 we can provide an analytical expression for the characteristic polynomial for each of the cases
 316 that we study. The computational complexity is then limited to the evaluation of the roots
 317 of this polynomial which scales roughly as $O(M^2 \log M)$, where $M = 2m$ is the degree of the
 318 polynomial and the maximum degree possible is $M = 2m = 2n_n N$ (e.g., in a typical TB model
 319 up to nn , $M = 2N$ and for that its complexity goes as $\sim O(8N^2 \log N)$). On the contrary, the
 320 well-established recursive GF technique has $O(N_r^3 N_{it})$ complexity [36, 41], where N_{it} typically
 321 $\gg 1$ is the number of iterations required for convergence in a desired energy precision η and
 322 the term N_r^3 is due to matrix inversions where the recursive matrix dimension $N_r = Nn_n$ grows
 323 with the number of neighbours.

324 For larger matrix dimensions or higher degree polynomials that the ones analyzed in this
 325 paper, FLA might suffer from numerical instability in the computation of the polynomial co-
 326 efficients due to accumulated errors in the trace in Eq. (10) and from the recursive nature of
 327 the successive polynomial coefficients [42, 43]. However in Ref. [32] FLA was used to obtain
 328 the bGF of TB Hamiltonians that cannot be solved using symbolic approaches due to matrix
 329 dimension (e.g., $N = 12$ Hilbert space dimension). So, despite the potential instability of the
 330 method, it still can be used to efficiently solve the bGF problem of TB Hamiltonians beyond
 331 analytical approaches, at least for moderate dimensions.

332 6 Conclusions and outlook

333 In this work we have extended the boundary Green function method developed in Refs. [23, 32]
 334 to 2D lattices with hopping elements between arbitrary distant neighbors and solved the semi-
 335 analytical obstructions to compute the bGF for large systems, non-orthogonal lattice vectors
 336 or Hamiltonians with terms with momentum fractions. This was made by implementing the
 337 Faddeev-LeVerrier algorithm to compute the characteristic polynomial and the adjugate matrix,
 338 the building blocks to compute the bGF. As an illustration of the method we have analyzed
 339 the spectral properties of different topological 2D Hamiltonians showing the appearance of
 340 topological states.

341 With FLA we can compute the bGF for any TB model with a well-known algorithm and
 342 a simple implementation which provides the coefficients of the characteristic polynomial but
 343 also the adjugate matrix in the same process. Furthermore, FLA can be extended to obtain the
 344 generalized inverses of multiple-variable polynomials or particularly, two-variable polynomials
 345 [38–40].

346 In Ref. [44, 45] it is claimed that the classical Faddeev-LeVerrier algorithm for polynomial
 347 matrices in one variable has $O(N^3 N)$ computational complexity and it avoids any division by
 348 a matrix entry, which it is desirable from the convergence perspective in contrast to recursive
 349 approaches. Although the classical FLA is not the most efficient algorithm from the point of
 350 view of complexity (e.g. Berkowitz algorithm [46] is faster), it is a rather simple and general
 351 way to solve the inverse of a polynomial matrix problem. Despite the recursive nature of FLA,
 352 it can be easily modified to carry out the N matrix multiplications in parallel [43, 44, 47, 48].

353 As an outlook, the FLA method can be combined with interpolation approaches [45, 49, 50]
 354 to improve the stability of the algorithm when computing the bGF of TB systems with a large
 355 number of degrees of freedom and neighbours. Furthermore, this method and the recursive

356 methods could be combined to describe systems with regions with broken translational sym-
 357 metry like two semi-infinite translational invariant regions coupled by a disordered region. In
 358 addition, we foresee the application of the method to study higher order topological insula-
 359 tors [51] which requires projection onto the intersection of two or more edge surfaces.

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365 A Exact Hamiltonian diagonalization

366 From the matrices that define the recursive method in Eq. (35) we can also describe the total
 367 Hamiltonian for a finite system to compute an exact diagonalization and obtain the edge state
 368 spectrum.

$$\hat{H}_{TOT} = \begin{pmatrix} \hat{\mathcal{H}}_0 & \hat{T}_{LR} & \hat{0} & \cdots & \hat{0} \\ \hat{T}_{LR}^\dagger & \hat{\mathcal{H}}_0 & \hat{T}_{LR} & \cdots & \hat{0} \\ \hat{0} & \hat{T}_{LR}^\dagger & \hat{\mathcal{H}}_0 & \cdots & \hat{0} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \hat{0} & \hat{0} & \hat{0} & \cdots & \hat{\mathcal{H}}_0 \end{pmatrix}, \quad (36)$$

369 where the main diagonal has N_{sites} block elements and total dimension $N_d = N_{sites}Nn_n$ so for
 370 the usual nn case satisfies $N_d = N_{sites}N$ and $\hat{\mathcal{H}}_0 = \hat{H}_2$, $\hat{T}_{LR} = \hat{H}_1^\dagger$.

371 B Faddeev-LeVerrier algorithm

372 We include here a simple pseudocode description of the classic FLA [27–31] to obtain the
 373 coefficients of the characteristic polynomial \bar{C} and the polynomial description of the adjugate
 374 matrix \hat{M} of the secular equation $[\omega\hat{\mathbb{I}} - \hat{H}]$ from a constant matrix (Algorithm 1).

Algorithm 1 Classic Faddeev-LeVerrier algorithm

Input: $\hat{H} \in \mathbb{C}^{n \times n}$ where $n \geq 2$

Output: (\bar{C}, \hat{M})

- 1: $\bar{C}_n = 1, \hat{M}_1 = \hat{\mathbb{I}}, k \leftarrow 2$
 - 2: $\bar{C}_{n-1} = -\text{tr}\{\hat{H}\}$
 - 3: **while** $k \leq n$ **do**
 - 4: $\hat{M}_k \leftarrow \hat{H}\hat{M}_{k-1} + \bar{C}_{n-k+1}\hat{\mathbb{I}}$
 - 5: $\bar{C}_{n-k} \leftarrow -\frac{1}{k}\text{tr}\{\hat{H}\hat{M}_k\}$
 - 6: $k \leftarrow k + 1$
 - 7: **end while**
-

375 We also describe the modified FLA for two variable polynomials in (ω, z) where the matrix
 376 itself $\hat{\mathcal{H}}(z)$ is a polynomial matrix [38–40] given as an entry the polynomial decomposition in
 377 z of the Hamiltonian as in Eq. (12) (Algorithm 2).

Algorithm 2 Two-variable Faddeev-LeVerrier algorithm

Input: $\hat{H}_1, \hat{H}_2, \dots, \hat{H}_{2m+1} \in \mathbb{C}^{n \times n}$ where $n \geq 2$

Output: (\bar{C}, \hat{M})

- 1: $\bar{C}_n = 1, \hat{M}_1 = \hat{\mathbb{1}}, k \leftarrow 2$
- 2: $\bar{C}_{n-1,1} = -\text{tr}\{\hat{H}_1\}, \bar{C}_{n-1,2} = -\text{tr}\{\hat{H}_2\}, \dots,$
 $\bar{C}_{n-1,2m+1} = -\text{tr}\{\hat{H}_{2m+1}\}$
- 3: **while** $k \leq n$ **do**
- 4: **for** $i \leftarrow 1 : 2m(k-1) + 1$ **do**
- 5: **if** $i \leq 2m(k-2) + 1$ **then**
- 6: $\hat{M}_{k,i} \leftarrow \hat{M}_{k,i} + \hat{H}_1 \hat{M}_{k-1,i}$
- 7: **end if**
- 8: **if** $i \geq 2$ **and** $i \leq 2m(k-2) + 2$ **then**
- 9: $\hat{M}_{k,i} \leftarrow \hat{M}_{k,i} + \hat{H}_2 \hat{M}_{k-1,i}$
- 10: **end if**
- 11: ...
- 12: **if** $i \geq 2m + 1$ **and** $i \leq 2m(k-2) + 2m + 1$ **then**
- 13: $\hat{M}_{k,i} \leftarrow \hat{M}_{k,i} + \hat{H}_{2m+1} \hat{M}_{k-1,i}$
- 14: **end if**
- 15: $\hat{M}_{k,i} \leftarrow \hat{M}_{k,i} + \bar{C}_{n-k+1} \hat{\mathbb{1}}$
- 16: **end for**
- 17: **for** $i \leftarrow 1 : 2mk + 1$ **do**
- 18: **if** $i \leq 2m(k-1) + 1$ **then**
- 19: $\bar{C}_{n-k,i} \leftarrow \bar{C}_{n-k,i} - \frac{1}{k} \text{tr}\{\hat{H}_1 \hat{M}_{k,1}\}$
- 20: **end if**
- 21: **if** $i \geq 2$ **and** $i \leq 2m(k-1) + 2$ **then**
- 22: $\bar{C}_{n-k,i} \leftarrow \bar{C}_{n-k,i} - \frac{1}{k} \text{tr}\{\hat{H}_2 \hat{M}_{k,i}\}$
- 23: **end if**
- 24: ...
- 25: **if** $i \geq 2m + 1$ **and** $i \leq 2m(k-1) + 2m + 1$ **then**
- 26: $\bar{C}_{n-k,i} \leftarrow \bar{C}_{n-k,i} - \frac{1}{k} \text{tr}\{\hat{H}_{2m+1} \hat{M}_{k,i}\}$
- 27: **end if**
- 28: **end for**
- 29: $k \leftarrow k + 1$
- 30: **end while**

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