Compensated Quantum and Topological Hall Effects of Electrons in Polyatomic Stripe Lattices

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The quantum Hall effect is generally understood for free electron gases, in which topologically protected edge states between Landau levels (LLs) form conducting channels at the edge of the sample. In periodic crystals, the LLs are imprinted with lattice properties; plateaus in the transverse Hall conductivity are not equidistant in energy anymore. Herein, crystals with a polyatomic basis are considered. For a stripe arrangement of different atoms, the band structure resorts nontrivially and exhibits strong oscillations that form a salient pattern with very small bandgaps. The Hall conductivity strongly decreases for energies within these bands, and only sharp peaks remain for energies in the gap. These effects are traced back to open orbits in the initial band structure; the corresponding LLs are formed from states with positive and negative effective mass. The partial cancellation of transverse charge conductivity also holds for different polyatomic stripe lattices and even when the magnetic field is replaced by a topologically nontrivial spin texture. The topological Hall effect is suppressed in the presence of magnetic skyrmions. The discussion is complemented by calculations of Hofstadter butterflies and orbital magnetization.

1. Introduction

The Hall effect in its quantized form,[1] was first described for free electrons, forming dispersionless Landau levels (LLs),[2] and first detected in 1980.[3] Later, Hofstadter butterflies[4–8] revealed essential differences in the shape of the LLs for electrons in a periodic lattice and their energy spacing. When this quantized Hall effect was observed in graphene in 2004,[9] research interest was renewed and calculations of Chern numbers showed for example that van Hove singularities (VHSs) of the initial band structure; the corresponding LLs are formed from states with positive and negative effective mass. The partial cancellation of transverse charge conductivity also holds for different polyatomic stripe lattices and even when the magnetic field is replaced by a topologically nontrivial spin texture. The topological Hall effect is suppressed in the presence of magnetic skyrmions. The discussion is complemented by calculations of Hofstadter butterflies and orbital magnetization.

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two different basis atoms and can be carried over to the THE in the presence of magnetic skyrmions (Section 3.3). We conclude in Section 4.

2. Model and Methods

To describe the QHE, we consider a tight-binding model on a square lattice with the Hamiltonian

$$ H = \sum_i f_i c_i^\dagger c_i + \sum_{\langle ij \rangle} t_{ij} c_i^\dagger c_j. \quad (1) $$

$c_i^\dagger$ and $c_i$ are spinless creation and annihilation operators (i and j site indices), $f_i$ is the onsite energy at site i, which dictates the superlattice unit cell. In the majority of this article, we consider a diatomic stripe lattice.

$t_{ij}$ is the nearest-neighbor hopping strength

$$ t_{ij} = e^{\varphi_{ij}}, \quad \varphi_{ij} = \frac{\varepsilon}{\hbar} \int_{r_i \rightarrow r_j} A(r) \cdot dr. \quad (2) $$

An external magnetic field $B = B e_z = V \times A$ with vector potential $A(r) = -B y e_z$ (Landau gauge) induces the complex phase factor. To preserve the periodicity of hopping amplitudes, the magnetic unit cell is enlarged compared with the onsite superlattice unit cell. The magnetic field has to be expressed by coprime integers $p$ and $q$: $p/q = \Phi/\Phi_0$ with $\Phi = Ba^2$ and $\Phi_0 = h/e$.\[4]

From the eigenvalues $E_n(k)$ and eigenvectors $|u_n(k)\rangle$ of the Hamiltonian (1), we calculate the Berry curvature

$$ \Omega_n(k) = \sum_{m \neq n} \frac{\langle u_m(k) | \nabla_k H(k) | u_n(k) \rangle \times (n \leftrightarrow m)}{|E_n(k) - E_m(k)|^2} \quad (3) $$

of band $n$ that enters the Kubo formula for the intrinsic transverse Hall conductivity

$$ \sigma_{xy}(E_F) = -\frac{e^2}{2\pi} \frac{1}{\hbar} \sum_{n} \int_{BZ} \Omega_n(k) F(E_n(k) - E_F) d^2k \quad (4) $$

as a Brillouin zone (BZ) integral.\[25] $F$ is the Fermi distribution function.

3. Results and Discussion

First, we recall the band structure and the Hall conductivity for the monatomic square lattice (Section 3.1). The results can be explained with the zero-field band structure straightforwardly. The stripe crystal (Section 3.2) dictates a different geometry of the superstructure unit cell, which leads to new effects in both the band structure and Hall conductivity. Finally, our findings are generalized for a polyatomic basis, different lattice types, and can be carried over to the THE of electrons in skyrmion textures (Section 3.3).

3.1. Preliminary Consideration: $f = 0$

As a prelude, we summarize the results for a monatomic lattice ($f_i \equiv f = 0$), i.e., a conventional QH system. For details, see the study by Göbel et al.\[13] The upcoming Hamiltonian of the stripe crystal (Equation (5)) can be used by setting $f = 0$.

The band structure of a QH system on a square lattice (Figure 2a) for $p = 1$ exhibits mainly flat bands (LLs) that are nearly equidistant for high and low energies, like for free electrons. This is due to the fact, that the band structure without magnetic field (the zero-field band structure, see Figure 2c) has a minimum (maximum) at $E = -4$ ($E = +4\hbar$), that resembles a free-electron (free-hole) parabola. Near a VHS where the density of states (DOS) diverges, the electrons do not behave "freely": the band spacing is reduced, and the LLs exhibit oscillations; these features are most pronounced for the bands close to the energy of the VHS $E_{VHS} = 0$ in the $\Gamma X$ direction.

The transverse Hall conductivity $\sigma_{xy}(E_F)$ in dependence of the Fermi energy decreases in steps of $e^2/h$ for every LL at energies below $E_{VHS}$ (Figure 2b); $\sigma_{xy}(E_F)$ is quantized. In this energy region, the bands carry a nonzero Chern number

$$ C = \frac{1}{2\pi} \int_{BZ} \Omega_\omega(k) d^2k = 1 \quad (5) $$

At $E_{VHS}$, for odd $q$ one band carries a total Chern number of $1 - q$ for even $q$. Two touching bands carry a joint $C$ of $2 - q$, which leads to a sign change in Hall conductivity: in addition to

![Figure 1. Main message exemplarily visualized for the THE of electrons due to a single skyrmion (noncollinear magnetic texture with out-of-plane components [gray] and inplane components [colored]). a) When the electron (blue ball) traverses the texture, its spin (arrow) aligns with the texture (same color) and gets deflected by the emergent magnetic field. b) In a diatomic lattice, characterized by different onsite energies $\pm f$ (vertical stripes), a mixed injection of electronic states with positive (blue) and negative (red) effective masses takes place; see text. Since both species feel the same force due to the emergent magnetic field, they are deflected into opposite directions. This leads to a cancellation in transverse charge transport.](image-url)
the LL character of these bands \(C = 1\) for one band for odd \(q\) and total \(C = 2\) for even \(q\) a large Chern number of \(-q\) is generated corresponding to the \(q\) band oscillations. Above \(E_{\text{VHS}}\), the Hall conductivity decreases again in steps of \(e^2/h\) back to zero.

The energy dependence of the Hall conductivity can be nicely reproduced using the band structure without magnetic field (zero-field band structure): In Figure 2c, a Fermi line below (above) \(E_{\text{VHS}}\) surrounds an electron (a hole) pocket. This corresponds to the negative (positive) sign in the conductivity. At the VHS, the electron pocket touches the BZ edge and becomes a hole pocket (Lifshitz transition). The effective mass changes sign and so does the Hall conductivity.\(^{13}\)

### 3.2. Compensation of the QHE for a Stripe Crystal

We proceed with the diatomic stripe crystal with alternating onsite energies along the \(y\) direction. Band structure and Hall conductivity show different features, which we explain with the orbits of the zero-field band structure.

The Hamiltonian in Landau gauge for \(q\) basis atoms reads

\[
\begin{pmatrix}
    h_1 - f & h_1^{(+)} & 0 & \ldots & 0 & h_1^{(-)} \\
    h_1^{(-)} & h_2 + f & h_2^{(+)} & \ldots & 0 & 0 \\
    0 & h_3^{(-)} & h_3 - f & \ldots & 0 & 0 \\
    \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\
    0 & 0 & 0 & \ldots & h_{q-1} - f & h_{q-1}^{(+)} \\
    h_1^{(+)} & 0 & 0 & \ldots & h_{q}^{(-)} & h_{q} + f
\end{pmatrix}
\]

(6)

with \(h_j = 2t \cos (\alpha k_y - 2\pi n^j q)\), \(h_j^{(\pm)} = \pm \text{e}^{\pm i \alpha k_y}\).

### 3.2.1. Band Structure

Starting again at \(f = 0\) (Figure 3a), we identify \(q\) LL oscillations in \(\Gamma X \Gamma\) direction, as already described in Section 3.1. Increasing \(f\) leads to \(q/2\) additional oscillations with a twice as large period, corresponding to the \(q/2\) atoms per sublattice. These modulations are most pronounced near \(E = \pm 2t\) and are independent of \(k_y\), which is at variance with the omnipresent oscillations that are strongest near \(E_{\text{VHS}}\).

For large \(f\), the amplitudes of these oscillations increase until bands seem to intersect Figure 3d–i. Close-ups (insets) show that the bands neither intersect nor touch. However, for \(f \to \infty\) the band spacing converges to zero; the band energies, given by the diagonal elements

\[
E_n = 2t \cos \left(ak_x - \frac{2 \pi n^j}{q}\right) + (-f)^n
\]

(7)

of the Hamiltonian, form a “weaving pattern.” For large \(f\) (e.g., \(f = 2t\) in Figure 3g) the block separation is clearly visible and the weaving pattern dominates the entire band structure of each block.

The origin of these patterns can be found in the zero-field band structure. The rectangular BZ accounting for the diatomic basis imposes a back-folding of the initial zero-field band structure. The single band in Figure 4a turns into two bands (blue and red) in Figure 4b that overlap in energy. At the BZ edge at \(k_y a = \pm \pi/2\), the bands are degenerate.

If \(f\) is increased, the band degeneracy is lifted and the bands’ slope becomes zero at the BZ edge,

\[
E_{12}(k) = \pm \sqrt{f^2 + 2t^2[1 + \cos(2k_y a)] + 2t \cos(k_x a)}
\]

(8)

In the energy range \(|E| < 2t\), in which the bands overlap initially, the zero-field band structure becomes deformed and
exhibits open orbits, which seem to introduce the weaving pattern. For $f > 2t$, the zero-field bands do not overlap and block separation sets in (Figure 3g–i).

3.2.2. Hall Conductivity

The weaving pattern brings about peculiar effects in the Hall conductivity (red in Figure 5). For a small $f (f = 0.2t$ in Figure 5a), a smooth shape of $\sigma_{xy}(E_F)$ is recognizable. The oscillations in the band structure (Figure 3b) cause drops in the modulus of the conductivity due to the conventional intrinsic Hall effect with opposite sign. However, the signal remains quantized in the bandgaps. "U-shaped" valleys appear at energies within the bands.

Upon increasing $f$, these valleys become deeper (Figure 5b) until the conductivity is almost zero (Figure 5d) but with very sharp peaks ("spikes") in the tiny bandgaps (insets in Figure 3) caused by topologically protected edge states. For sizable $f$ (Figure 5c–f), the conductivity in such gaps eventually changes sign. For $f \geq 2t$, the block separation is accompanied by a sign reversal of the transverse conductivity roughly in the middle of each block (despite being vanishingly small apart from the very small bandgaps). In the limit $f \to \infty$, the blocks themselves become antisymmetric.

Especially for large $f$ the Hall conductivity is suppressed over a large energy range. An intuitive argument, stemming from the synthetic case $f/t \to \infty$, is that the stripes in the lattice formed by different onsite energies make the system quasi-1D and prohibit any transverse charge transport (cf. Equation (6)). For any finite $f/t$, the spikes correspond to topologically protected edge channels in the gaps.

In the following paragraphs, we establish an explanation using open orbits in the zero-field band structure that lead to a mixing of electron states of positive and negative effective mass. We claim that these two species of electrons are deflected into opposite transverse directions, meaning that electrons do not move one dimensionally, as speculated earlier, but rather their transverse motion is compensated (in an analogy to a spin Hall scenario). In a later section of the article, we prove this claim by visualization of the scattered electrons' density. The established interpretation allows also to explain the complicated behavior of the transverse conductivity for small $f/t$ ratios.
3.2.3. Relation to Open Orbits in the Zero-Field Band Structure

The evolution of the zero-field band structure with $f$ is accompanied by a change of the character of the Fermi lines as the bands partially overlap for $f \leq 2t$. In any case, the lower band exhibits one electron pocket for low energies (blue lines in Figure 4c), open orbits (gray) between the VHSs (purple), and one hole orbit (red) for energies above the VHSs. The upper band behaves similarly. This segmentation allows to understand the shape of $\sigma_{xy}(E_F)$. A scheme for estimating the envelope function is given in the following paragraphs.

Near the edges of the energy spectrum the zero-field band structure does not exhibit open orbits. Here the transverse conductivity can be estimated by the enclosed “area” (in reciprocal space) of Fermi lines in the zero-field band structure. The fermion character of a closed Fermi line determines the sign of the Hall conductivity: electron like gives a negative sign and hole like is positive.[13] This approximation (blue) describes well the

Figure 5. QH conductivity for the diatomic stripe crystal with $p/q = 1/20$. The explicitly calculated transverse Hall conductivity $\sigma_{xy}(E_F)$ (red) is compared with the approximation of the envelope (blue, see text for details). In green-shaded energy intervals, closed and open electron and hole orbits overlap. The unit of quantization is $\sigma_0 = e^2/h$.

Figure 6. Orbit characterization of the zero-field band structure (Figure 4) and explanation of the approximation used for the Hall conductivity $\sigma_{xy}(E_F)$ in Figure 5. Each panel shows a left and a right block, corresponding to the lower or upper band of the zero-field band structure, respectively. The white line shows the DOS in arbitrary units with two VHSs each (the two peaks). Between the corresponding energies, the Fermi line is open (Figure 4c). Near the minimum (maximum), the Fermi line is a closed electron (hole) pocket visualized in blue (red). For the case $f = 0$, electron and hole states are separated at $E = 0$. For $f > 0$, the border is shifted to $E = -f (E = +f)$ for the lower (upper) band. This energy is always located between the VHSs in the open-orbit regime. In the overlapping areas, an LL may be formed from electron-like and hole-like states. In the closed-orbit regime, the fermion character with the higher DOS dictates the character of the LL. Energy regions with trivial fermion character are colored in red (hole) or blue (electron) and a darker shade in the open-orbit regime. When electron-like and hole-like states mix in the open-orbit regime, the fermion character is not trivial, and the region is colored green, allowing for a sign change of the Hall conductivity peaks in Figure 5.
explicitly calculated conductivity (red; Equation (4)) in the corresponding energy range. For $f < 2t$, closed electron and hole orbits are also present in the middle of the energy range (Figure 6c–g). In this case, the fermion character with the higher DOS dictates the sign (Figure 5a–g), which automatically leads to a sign change at $E = 0$.

For energies that are characterized by open orbits, the conductivity is reduced—for large $f/t$ ratios, this reduction is drastic which could mean that LLs formed from open orbits do not contribute to transverse transport at all. However, this would not explain the spikes and why for small $f/t$ the transverse conductivity is sizable over the whole energy range. For this reason, let us revisit the case $f/t = 0$. When treating the system in the native unit cell (one basis atom, as all atoms are equivalent) the zero-field band structure is a single band that exhibits only closed orbits upon cutting at different energies (Figure 2c). As shown in Section 3.1 the conductivity does not exhibit U-shaped valleys, i.e., it is not suppressed. When treating the system in the diatomic basis (even though this is not necessary here), the zero field-band structure is backfolded and open orbits appear, as shown in Figure 4. Due to the equivalence of the two systems, we can clearly tell that these open orbits behave exactly like closed orbits—which seems to be partially reminiscent even for larger $f/t$ ratios explaining the conductivity peaks in the small gaps between LLs.

The scheme for approximating the envelope therefore accounts also for states characterized by an open orbit when counting the occupied states to determine $\sigma_{x y}(E_F)$. The open orbits have either electron or hole character depending on their energy in relation to $\pm f$ (cf. Figure 6). This single assumption reproduces the peaks if open-orbit energy regimes of one band do not overlap with closed-orbit regimes of the other zero-field band (for $f > 1.5t$; Figure 5h and 6h). Furthermore, it can explain the mostly unsuppressed conductivity in the open-orbit regime for small $f/t$ (Figure 5a). In all cases, this approximation seems to determine the envelope function of $\sigma_{x y}(E_F)$.

Going more into detail, one case deserves special attention. For $f/t < 1.5$ we find small energy ranges where the two zero-field bands exhibit open and closed orbits of different characters for the same energy (highlighted green in Figure 5 and 6). Consequently, in these cases, the transverse conductivity may change its sign. The envelope function can be constructed by accounting for “both” orbits as either electron like or hole like. For this reason, the two branches of the envelope function have a positive sign and a negative sign in these energy ranges, respectively. The envelope function can be comprehended in detail when comparing Figure 5 and 6.

Summarizing at this point, we find a strongly diminished QH conductivity in a stripe bipartite lattice. Deflection of electrons is suppressed by the two inequivalent sublattices except for small energy gaps where topologically protected edge channels allow for transport, like for a conventional QH system with $f = 0$.

The quantization of open-orbit states is not as straightforward as for closed orbit states. Our findings suggest that they behave like closed orbits at the very edges of the LLs, so that $\sigma_{x y}$ in the bandgaps exhibits quantized peaks as for closed orbits. The Chern numbers of those bands (except for those corresponding to a sign change in the Hall conductivity) are still $C = 1$. However, the Berry curvature only has a considerable magnitude near the band extrema. “Within the bands” (apart from the LL edges) the conductivity drops nearly to zero, accounting for the mixed electron and hole characters of open orbits.

3.2.4. Hofstadter Butterfly

The full field dependence of the energy spectrum of a QH system is commonly described by so-called Hofstadter butterflies.

For $f < 0$, we obtain the well-known Hofstadter butterfly of the square lattice (Figure S2a, Supporting Information). At the very left of that figure ($\Phi/\Phi_0 = 1/q$), black dots represent the LL behavior we have shown in Section 3.1: equidistant, flat bands in the low- and high-energy ranges, and very narrow bands with increased band width near the VHS. The general case of $p/q$ can be reduced to known cases ($p = 1$) by an expansion into continuous fractions. A magnetic flux of $(p/q)\Phi_0$ with $p > 1$ is reduced to a flux of $(1/q)\Phi_0$ after the expansion, where each of the $q$ bands comprises a bundle of several bands. This “bunching” leads to the fractal nature of a Hofstadter butterfly. The exact band distribution can be found via the expansion into continuous fractions or from the Diophantine equation.

For small $f = 0.5t$ (Figure 7a), the energy spectrum generally looks similar to the Hofstadter butterfly for $f = 0$ (Figure S2a, Supporting Information). The introduced band oscillations lead to a denser appearance and condense in complicated patterns with additional “lines” (e.g., in the extended topmost structure). A symmetric deformation resembles the spread out zero-field band structure.

For $f = 3t$ (Figure 7b), block separation is clearly established. The two blocks of the butterfly appear confined by almost straight lines, whereas for small $f$, the energy spectrum had the famous butterfly shape. The two blocks are almost densely filled for large $f/t$, which is explained by the weaving pattern that dominates the entire band energy range. The origin of the few white lines (unoccupied energies) within this “filling” can be understood from the band structure for $\Phi/\Phi_0 = p/q = 3/20$ (Figure 8). The associated continued fraction

$$\frac{3}{20} = \frac{1}{6 + \frac{1}{1 + \frac{1}{1}}}$$

(9)

yields only six groups of bands. An increased $f$, as compared with the case $p/q = 1/20$, is thus needed to make the bands touch. The white spaces (bandgaps) in the Hofstadter butterfly vanish for an even larger $f$.

3.2.5. Other Lattice Types

The results for the stripe crystal can be carried over to the general case of $b = 2, 3, \ldots, q$ basis atoms in the “structural” basis ($b$ has to be a factor of $q$). Choosing all of the $b$ onsite energies differently separates the band structure into $b$ blocks, in which each block exhibits the weaving pattern with $qb$ band oscillations. As an example, we choose $b = 3$ (Figure 9).

For the diatomic case $b = 2$, the bands showed $q/2$ oscillations for each of the two blocks in $\Gamma\Gamma$ direction. The oscillations of the upper block were shifted half an oscillation length in $\Gamma\Gamma$. 
Now, for \( b = 3 \), we find three blocks with \( q/3 \) oscillations. The bands look again almost identical when comparing the blocks but they are shifted by \( 1/3 \) (2/3) of an oscillation length for the middle (upper) block in \( \Gamma \) direction.

The behavior of the Hall conductivity \( \sigma_{xy}(E_F) \) can be deduced in analogy to the case \( b = 2 \). We now find \( b \) blocks that exhibit the same shape in the limit \( f \gg t \).

The established findings of the suppressed Hall conductivity and the weaving pattern in the band structure are not limited to stripe square lattices. We find similar results for triangular lattices with stripes of different onsite energies and expect these features for other polyatomic lattices. They appear as long as open orbits are present in the zero-field band structure. As we show in the next section, the results are not even limited to the QHE due to an external magnetic field; we also find a suppressed THE in the presence of a skyrmion crystal and even for single skyrmions.

One exceptional lattice is the checkerboard-type diatomic square lattice, as shown in the Supporting Information (Figure S1 and S2). Here, the zero-field band structure is folded back exactly at the straight orbit including the VHS, where the orbits transition from electron like to hole like. For this reason, the back-folded zero-field band structure consists of one purely electron-like band and one purely hole-like band. In this case, the introduction of \( f > 0 \) does not generate open orbits but leads to a trivial splitting of the zero-field band structure at \( E_{VHS} = 0 \). In the presence of an external magnetic field, we find bands without the weaving pattern and an unsuppressed QH conductivity. The lack of open orbits further substantiates our aforementioned argumentation.

A suppression of the Hall conductivity arising from open orbits can alternatively be established upon considering a monatomic lattice with anisotropic hoppings \( t_x \neq t_y \). Figure 10 shows the results of a square lattice with \( p/q = 1/20 \), where each atom has the same onsite energy but the hopping amplitudes are \( t_x = 2t \) and \( t_y = t \).
In this section, we replace the external magnetic field by a skyrmion crystal to discuss the topological contribution to the Hall effect. In addition to skyrnion crystals in stripe lattices, formed by a polyatomic basis from different elements, we have in mind multiparticle systems. Consider, for example, a skyrnion layer on top of a heavy metal layer with a lattice mismatch. At the interface, lattice relaxation may arise and stripe reconstructions like for Si(100) may form. The electrons in the skyrnion layer on top will then feel onsite differences distributed in a stripe shape. Another scenario is that the skyrnion layer itself becomes buckled which leads to stripe-shaped oscillations in the atomic distance to the layer below which translates to oscillations in the onsite energy.

The spins of the electrons of Hamiltonian (1) \( \{ c_i \} \) are now spinor-valued operators couple to a skyrnion texture \( \{ s_i \} \)

\[
H_{\text{Sk}} = m \sum_i s_i \cdot \left( c_i^\dagger \sigma_i c_i \right)
\]

(\( \sigma \) vector of Pauli matrices) via Hund’s coupling. We model skyrnions in a ferromagnetic surrounding like in the study by Göbel et al.\(^{[10]} \) The topological charge is \( N_{\text{Sk}} = -1 \), as the center spin shall point into negative \( z \) direction. This leads to a negative emergent field\(^{[19,31]} \)

\[
B_{\text{em}}^{(z)}(\mathbf{r}) \propto n_{\text{Sk}}(\mathbf{r}) = s(\mathbf{r}) \cdot \frac{\partial s(\mathbf{r})}{\partial x} \times \frac{\partial s(\mathbf{r})}{\partial y}
\]

(\( B_{\text{em}}^{(z)} \) is the component in the \( z \) direction). This effective magnetic field is found to be negative in a continuous formulation, which effectively accounts for the noncollinearity of the skyrnion spin texture.

In this respect, the QHE and the THE are related\(^{[12,13,24]} \) and many of the aforementioned findings also appear for the THE of electrons in skyrnion crystals. We would like to emphasize that the emergent field was not used for any calculations and merely serves to easily interpret the THE generated by the spin chirality of the skyrnion texture.

Bandgaps and Chern numbers are similar to the LLs in the QH system. The main difference compared with the QH system is that for electrons in a skyrnion crystal, the bands are dispersible so that the widths of global bandgaps are decreased. The reason for this is the inhomogeneity of the emergent field, in contrast to the homogeneous external magnetic field in the QH case. However, the average value of the emergent field can be related to the magnetic field of a QH system \([ B_{\text{em}} = B \neq B_{\text{em}}(\mathbf{r}) ] \); in this sense, a skyrnion crystal with \( n_0 \) sites in the unit cell corresponds to the QH scenario of \( p/q = \pm N_{\text{Sk}}/n_0 \).

The two signs correspond to the two blocks in the band structure. They occur due to the introduction of the electron spin: one block corresponds to electrons with their spin aligned parallel with respect to the texture and one where the spins are aligned antiparallel. The two blocks are shifted by \( \pm m \), respectively, for \( m = 5t \) (chosen throughout this article).

When the onsite energy of a bipartite lattice is introduced, the band structure (Figure 11a) exhibits similar features as for the QH system. Each of the two blocks begins to split up again like for the one block in the QH system. Most importantly, we find again the weaving pattern. This time the unit cell has to be chosen quadratic (here \( 6 \times 6 \) lattice sites). In analogy to the QH system, one observes three oscillations in \( \Gamma \chi \Gamma \) direction; due to the inhomogeneity of the emergent field, the oscillations are no more perfectly periodic. However, the weaving pattern still leads to the suppression of the topological Hall conductivity \( \sigma_{xy}(E_F) \) nearly to zero in the same energy range as for the corresponding QH system shifted by \( \pm m \) (Figure 11b). The aforementioned established interpretation using open orbits of the zero-field band structure holds also in this scenario.

To further investigate the mixing of electronic states with positive and negative effective mass, we use a Green’s function approach and compute the scattered electron’s density for a

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Figure 10. Monatomic lattice with anisotropic hopping \( t_x = 2t \) and \( t_y = t \). a) The zero-field band structure with electron orbits (blue), hole orbits (red), and open orbits (white). The three regimes are separated by orbits containing VHSs (purple). b) Band structure when an external field \( (p/q = 1/20) \) is applied. c) The corresponding Hall conductivity in units of quantization \( \sigma_0 = e^2/h \).
single skyrmion in a finite ferromagnetic sample (corresponding to the sketch in Figure 1), like in a racetrack storage device.\textsuperscript{[12–15]} For this purpose, the program package KWANT\textsuperscript{[36]} is used similar to the example presented in the documentation of this code (Section 2.7; cf. also the studies by Yin et al., Hamamoto et al., and Göbel et al.\textsuperscript{[37–40]}) A bias voltage imposes an electric current, where electrons move from the left to the right terminal. To visualize the effect of the skyrmion on the deflection, the background electron density (the scattered electron density for a purely ferromagnetic system) is subtracted from the scattered electron density of the skyrmion system.

The result is shown in Figure 12: for \( f = 0 \) (monatomic lattice) and \( E_F = 3.1t \), the zero-field band structure exhibits one closed electron pocket. Electrons with a positive effective mass align their spin with the skyrmion texture and are therefore deflected to the bottom due to the negative emergent field. For nonzero \( f \), open orbits appear in the zero-field band structure. We find that electrons are deflected to the bottom and top due to coupling to the skyrmion (Figure 12b). Blue stripes (deflected electrons) are visible in both transverse directions. This finding motivated the schematic Figure 1 from the introduction.

The blue stripes also show white vertical lines in this case (every second lattice site in the horizontal direction appears white). This means that electrons are mainly “living” on one of the two sublattices of the bipartite lattice. We find that for the two different stripes, electrons live on the opposite sublattice, which means that the mixing of the electronic character (positive and negative effective mass) appears separated in real space.

The classical formula for the deflection of electrons by the Lorentz force
\[
\hat{r} = -\frac{e}{m^*} \hat{r} \times B_{\text{em}}
\] (12)
gives two possibilities to suppress the THE. The first is to mix two spin species as these feel opposite emergent fields \( \pm B_{\text{em}} \). This happens in antiferromagnetic skyrmions\textsuperscript{[41–45]}—a skyrmion consisting of two sublattices with mutually reversed spins. The THE vanishes, and a topological spin Hall effect emerges: electrons with spins aligned parallel to the texture are deflected into one transverse direction, whereas electrons with oppositely aligned spins are deflected into the other direction. For weak Hund’s coupling (\( m^*/t < 4 \)),\textsuperscript{[130,135]} the same can occur for conventional skyrmion crystals at specific Fermi energies.

The second possibility is presented in this article: the introduction of onsite energies that can lead to a mixing of electronic states with positive and negative effective masses \( m^* \) (see Equation (12)). Here, the spin of the electrons is aligned parallel with respect to the texture, so a spin Hall effect cannot emerge. Instead, \( m^* > 0 \) states are deflected in one transverse direction, whereas \( m^* < 0 \) states are deflected in the opposite transverse direction. Conceptually, this situation describes an “effective mass Hall effect” similar to the vanishing Hall effect in compensated metals.

4. Conclusions

Using a tight-binding model, we investigated the influence of a homogeneous magnetic field on band structure and Hall conductivity of a stripe square lattice featuring a diatomic basis. The band structure exhibits a weaving pattern, leading to a suppression of the Hall conductivity for energies within the strongly dispersive LLs. In the bandgaps (that become tiny for \( f \gg t \)),
the Hall conductivity is still quantized, with the consequence of sharp peaks in $\sigma_{xy}(E_F)$ at the corresponding energies.

These features are traced back to open orbits in the zero-field band structure, that exhibit positive and negative orbit curvatures. Electrons with a positive effective mass live on one sublattice and are deflected in one direction, whereas electrons with a negative effective mass live on the other sublattice and are deflected in the opposite transverse direction, yielding a compensated transverse charge deflection. A similar argument holds for the orbital magnetization. It is strongly suppressed when the Hall conductivity is suppressed. This result is exemplarily shown for an electronic system in the presence of an external magnetic field in the Supporting Information (Figure S3).

The results and their interpretation can be generalized to other polyatomic stripe lattices. Furthermore, the results can be carried over to the THE of electrons in topologically nontrivial spin textures like skyrmion crystals, but also bimeron crystals and even isolated topological objects in a ferromagnetic environment. If the Fermi energy is within a specific range, electrons in these textures on a polyatomic stripe lattice exhibit no net transverse charge transport, but—conventionally speaking—an effective mass Hall effect is present.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

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