Chern numbers hiding in time-of-flight images

Erhai Zhao,1,2 Noah Bray-Ali,2 Carl J. Williams,2 I. B. Spielman,2 and Indubala I. Satija1,2

1School of Physics, Astronomy, and Computational Sciences, George Mason University, Fairfax, Virginia 22030, USA
2Joint Quantum Institute, National Institute of Standards and Technology and University of Maryland, Gaithersburg, Maryland 20899, USA

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We present a technique for detecting topological invariants—Chern numbers—from time-of-flight images of ultracold atoms. We show that the Chern numbers of integer quantum Hall states of lattice fermions leave their fingerprints in the atoms’ momentum distribution. We analytically demonstrate that the number of local maxima in the momentum distribution is equal to the Chern number in two limiting cases, for large hopping anisotropy and in the continuum limit. In addition, our numerical simulations beyond these two limits show that these local maxima persist for a range of parameters. Thus, an everyday observable in cold atom experiments can serve as a useful tool to characterize and visualize quantum states with nontrivial topology.

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I. INTRODUCTION

Topological insulators and superfluids are many-body quantum systems with energy gaps in the bulk but topologically protected, gapless excitations on the boundary [1,2]. Each class of topological phases is characterized by a topological invariant which can be either an integer or a binary quantity. For example, the integer quantum Hall Hall numbers [3] are distinguished from other two-dimensional insulators with the same symmetry by the Chern number, an integer that captures the global properties of all occupied bands [4]. The Chern number coincides with the number of chiral edge modes and the Hall conductance in units of the conductance quantum. A central problem in exploring new forms of topological matter is how to extract the topological invariants from experiments and unambiguously determine the topological phase.

Experiments with ultracold atoms have brought us opportunities to create topological phases of matter in parameter regimes unreachable in solids. In addition to band-structure engineering using optical lattices and tuning the atom-atom interaction by Feshbach resonance, artificial magnetic fields for neutral atoms have been produced by either rotation [5] or atom-laser coupling [6,7], making way for realizing quantum Hall states of cold atoms [8]. Moreover, non-Abelian synthetic gauge fields [9,10] and spin-orbit coupling [11] have been demonstrated. These developments motivated theories on properties of ultracold topological matter, e.g., the fractional quantum Hall effect of interacting bosons [5,11,12], the quantum spin Hall effect of fermions [9], and the quantum anomalous Hall states of p-orbital fermions [13].

While cold atom systems offer new detection techniques and great tunability, it is challenging to measure the topological invariants directly. Measurements of local density of states of (the edge states) or mass transport are marred with complexity in contrast to solid-state systems. Thus, creative methods have to be invented. For example, Ref. [8] proposed to extract the quantized Hall conductance from the derivatives of the in situ atom density distribution by the Streda formula. Here, we show that the Chern number of a lattice-quantum Hall system of fermions can be simply determined from the structures (ripples) in the momentum distribution n(k) as measured in time-of-flight (TOF) images for a range of parameters. Our results transform n(k), a simple everyday observable of cold atom experiments, into an unexpected, useful tool for characterizing and visualizing topological states of matter.

Figure 1(a) summarizes the main results of this paper obtained for the Hofstadter model [14]. We highlight the parameter regimes where the structure in n(k) is most evident: in the continuum limit [Fig. 1(b)] or for highly anisotropic lattices [Fig. 1(c)]. In particular, we provide a transparent understanding of n(k) for large hopping anisotropy. In this limit, states with energy at the band edges dominate, and they are dimerized (their localized wave functions are peaked at pairs of spatially separated lattice sites). The size of these “Chern dimers,” in units of the lattice spacing d, is equal to the Chern number of the corresponding gap. As a result, n(k) varies sinusoidally [Fig. 1(c)] with a period determined by the Chern number. Furthermore, our detailed numerical studies verify that such distinctive features in the momentum distribution persist beyond the Chern dimer and the continuum limit [Figs. 2(c) and 2(d)]. However, the correspondence between the number of local maxima in n(k) and the Chern number is not always precise for arbitrary parameters.

II. ULTRACOLD FERMIONS IN SYNTHETIC MAGNETIC FIELD

We consider the Hofstadter model [14] of spinless fermionic atoms on a square optical lattice in the e_y-e_x plane:

\[ H = - \sum_{\langle \mathbf{s}, \mathbf{s}' \rangle} t_{\mathbf{s}, \mathbf{s}'} (\mathbf{e}^{i \theta_{\mathbf{s}, \mathbf{s}'}} \mathbf{f}_{\mathbf{s}}^\dagger \mathbf{f}_{\mathbf{s}'} + \text{H.c.}) - \sum_{\mathbf{s}} \mu_{\mathbf{s}} \mathbf{f}_{\mathbf{s}}^\dagger \mathbf{f}_{\mathbf{s}}. \]  (1)

Here, \( \mathbf{f}_{\mathbf{s}}^\dagger \) is the creation operator for a fermion at site \( \mathbf{s} = \{s_x, s_y\} \), the nearest-neighbor hopping strengths \( t_{\mathbf{s}} = t_{s_x, s_y} \), and \( t_{s_x, s_y} \) are real and positive, \( \theta_{\mathbf{s}, \mathbf{s}' = 0} = e \int_{\mathbf{s}} \mathbf{A} \cdot d\mathbf{l} \) depends on the vector potential \( \mathbf{A} \) for particles with charge \( e \), and \( \mu_{\mathbf{s}} \) is a local chemical potential. With cold atoms, the hopping anisotropy \( \lambda = t_\gamma/t_x \) can be tuned by varying the intensity of the lasers, giving rise to the optical lattice potential [6]. For an applied magnetic field \( \mathbf{B} = Be_x \), normal to the two-dimensional (2D) plane, \( \alpha = d^2 e B / h \) is the magnetic flux per plaquette in

\[ \lambda = t_\gamma/t_x \]  (a)

In cold atom experiments the product \( e \) enters the Hamiltonian, and neither \( e \) nor \( A \) is separately defined.
FIG. 1. (Color) Relation between the Chern number $c_r$ and structure in TOF data as a function of magnetic flux per unit cell $\alpha$ in units of the flux quantum $h/e$ and hopping anisotropy $\lambda$. (a) Schematic map of the $\alpha$-$\lambda$ plane showing the Chern dimer and the continuum regimes. Numerical results for the remainder of the parameter space will be shown in Figs. 2 and 3. (b) Isotropic, nearly-continuum: $\lambda = 1, \alpha = 0.05$ and (c) Chern dimer: $\lambda = 1000, \alpha = 0.45$. The data in (b) is for an isotropic $\lambda = 1$ lattice at $\alpha = 0.05$, and the data in (c) is for an anisotropic $\lambda = 1000$ lattice at $\alpha = 0.45$. $k_x$ is measured in units of $1/d$, where $d$ is the lattice spacing.

units of the flux quantum. We study homogeneous gases with $\mu_s = \mu_r$ and also finite systems in the presence of an isotropic harmonic trap potential, captured by $\mu_s = \mu_0 - m\omega^2 d^2 s^2 / 2$.

Artificial magnetic fields have been implemented for bosons [15], and it is straightforward to apply these techniques to fermions. With their smaller fine-structure splittings, the alkali fermions ($^6$Li and $^{40}$K) are more susceptible to heating from spontaneous emission than some bosons ($^{87}$Rb and $^{133}$Cs), but, for the $h/t_{x,y} \sim 240 \mu$s time scales$^2$ associated with the major energy gaps in the spectrum [see Fig. 2(a)], the expected $\approx 200$-ns lifetime in $^{40}$K should be acceptable; see Ref. [9] for an alternate solution.

We first summarize the well-known properties of the homogeneous Hofstadter model [Eq. (1)] with commensurate flux $\alpha = p/q$, where the integers $p,q$ are relatively prime. In the Landau gauge,$^3$ suitable for current experiments [6], the vector potential $A = \alpha x e_y$ scales the unit cell along $e_x$ by a factor of $q$ to a magnetic unit cell; this also creates a magnetic Brillouin zone (MBZ) with $k_x \in (-\pi q, \pi q)$ and $k_y \in (-\pi, \pi]$. The Hofstadter model is diagonal in the single-particle basis $\Psi_{s_x,s_y} = e^{i(k_x k_s x + k_y s_y)} \Psi_{s_x} (k_y)$, provided $\psi_{s_x}$ satisfies [14]

$$\hat{e}^{k_y} \psi_{s_x+1} + e^{-ik_y} \psi_{s_x-1} + 2\lambda \cos(2\pi s_x \alpha + k_y) \psi_{s_x} = -E_s \psi_{s_x},$$

$^2$Estimated assuming a $4E_s$ deep lattice from retroreflected $\lambda = 2d = 800$-nm lasers, giving $t_{x,y} = 0.085E_s$, where the single-photon recoil energy is $E_r = h^2 / 2m \lambda_L^2 = h \times 7.8$ kHz and $m$ is the atomic mass.

$^3$Our analysis can be generalized to the symmetric gauge which is appropriate for rotating gases; see, e.g., S. Powell et al., Phys. Rev. A 83, 013612 (2011).
with energy in units of $\hbar \omega_c$. The index $r = 1, 2, ... q$ labels linearly independent solutions, and $\psi_{\alpha s,\nu} = \psi_{\alpha s} e^{ik_l(s-z)}$. The single energy band at $\alpha = 0$ splits into $q$ sub-bands, and the energy spectrum has $q - 1$ gaps [14]. When the chemical potential $\mu$ is inside the $r$th energy gap, the system is an integer quantum Hall state with Chern number $c_r$, defined by the Thouless-Kohmoto-Nightingale-Nijs formula [4] and related to the quantized Hall resistance.

Usually we consider TOF images as approximate representations of the momentum distribution $n(k)$ obtained by imaging the 2D atomic column density after a $t_{\text{TOF}}$ period of free expansion. In most cases, the final spatial coordinate of an atom is $x = r_{\text{TOF}}$, where $v$ is the velocity [16]; however, vector potentials complicate the interpretation of TOF images because the velocity is $v = \hbar k/m - e \lambda B/m$. In Ref. [17], we showed that suddenly turning the gauge field to zero is equivalent to a brief electric field that takes the initial momentum $\tilde{h}k$ and maps it to a new final velocity $v' = \hbar k/m$. As a result, detected TOF images do approximate momentum distributions. The crystal momentum distribution

$$\tilde{n}(k) = \sum_x \langle f_x^i f_x^j \rangle e^{ik_l(s-z)}$$

(3)

can be reconstructed from TOF images using the recipe in Ref. [18]. Here, the average $\langle \cdots \rangle$ is over the ground state. We also define a one-dimensional (1D) crystal momentum distribution by integrating $\tilde{n}(k)$ along $e_y$:

$$n(k) = \int_{-\pi}^{\pi} dk_y \tilde{n}(k).$$

(4)

For homogeneous systems, we computed $n(k)$ by numerically diagonalizing Eq. (2) using a sufficiently large unit cell, while for finite trapped systems we diagonalized Eq. (1). Our numerical results are complemented by analytic analysis for low flux and large anisotropy. In what follows, we discuss in turn the behaviors of $n(k)$ in different parameter regimes, starting with an isotropic lattice and low flux.

III. OSCILLATIONS IN MOMENTUM DISTRIBUTION

Figures 1(b) and 2(b) show $n(k)$ in this isotropic low flux limit, at $\alpha = 0.05$ and $\lambda = 1$. As the chemical potential is increased up to $\mu = 0$, successively higher Hofstadter bands are filled, with Chern numbers $c_r = r$, where $r = 1, 2, 3, ...$ labels the highest filled band. Correspondingly, $n(k)$ has exactly $c_r$ local peaks on top of a smooth envelope; i.e., counting peaks in TOF images unambiguously determines the Chern number and therefore which quantum Hall state the system is in. To understand this, we recall that, for $\alpha \ll 1$, the cyclotron orbit is large compared to the lattice spacing, so the system resembles a free Fermi gas in magnetic field, and the narrow bands in the Hofstadter spectrum map to Landau levels. For $r$-filled Landau levels, $n(k)$ is the summed square modulus of the momentum-space wave functions:

$$n(k) \propto \sum_{\nu=0}^{r-1} (2\nu')^{-1} H^2_r(k_x, \ell_B) e^{-i(k_x s)^2},$$

(5)

where $H_r$ are the Hermite polynomials and $\ell_B = \sqrt{\hbar/eB}$ is the magnetic length. Due to the node structure of $H_r$, $n(k)$ has exactly $r$ local maxima located at the zeros of $H_{r+1}$. This is reflected in our numerical results as plotted in Fig. 1(b) for the lattice system. As is depicted by the shaded regions in Fig. 1(a), this association is valid in the entire low flux regime and is insensitive to $\lambda$.

At larger $\alpha$ the Landau level picture breaks down, but the correspondence between the number of local peaks and the Chern number appears to persist for a range of parameters. Figures 2(c) and 2(d) show examples of $n(k)$ for large flux $\alpha = 0.45$ and $\lambda = 1$, where well-resolved peaks do remain. More generally, as $\alpha$ increases (for $\mu$ residing within the same energy gap), the distinctive features of $n(k)$ are generally retained, but, compared to the low flux regime, the local peaks (except at $k_x = 0$) move to higher momenta and can be subsumed into the background structure. In some instances, these peaks turn into shoulderlike structures.

Intriguingly, as the hopping anisotropy becomes large, small wiggles on a background turn into regular and pronounced oscillations regardless of $\alpha$. Figures 2(c) and 2(d) illustrate the progression of $n(k)$ from $\lambda = 1$ to 10, showing pronounced oscillations develop with increasing $\lambda$. Most importantly, the simple rule reappears in the limit of $\lambda \gg 1$: the number of oscillations in $n(k)$, i.e., the number of peaks for $k_x \in [-\pi, \pi]$, is again exactly equal to the Chern number $c_r$. As is emphasized in Fig. 1(c), $n(k)$ asymptotically approaches

$$n(k) \propto r \pm \frac{1}{2} \cos(c_r k_x),$$

(6)

for chemical potential inside the $r$th gap. The one-to-one correspondence, or mapping, between the number of oscillations and the Chern number for anisotropic lattices [Fig. 1(a); Chern dimer regime] is one of our key observations, which we now explain analytically.

IV. CHERN DIMERS

The $\lambda \gg 1$ limit is tractable in perturbation theory [19], where the solutions to Eq. (2) are simple. When $t_z = 0$ (i.e., $\lambda \to \infty$), there are $q$ bands each with dispersion $E_r(k_x) = -2t_r \cos(2\pi r x + k_x)$, $r \in \{1, ..., q\}$ (see Fig. 3, where $p = 1$ and $q = 5$). These eigenstates are extended along $e_x$ and localized to single sites as a function of $x$. As defined in Eq. (2), the $r$th band describes states $\psi_{s,\nu}^r \delta_{s, r}$ localized at site $s_r = r$ and displaced in $k_x$, as identified in Fig. 3. The bands intersect at numerous degeneracy points $k^* \in \{r \pm \ell \}$ for states $r = s_r$ and $r' = s_r'$. The spatial separation along $e_x$ of these localized states is $\ell = s_r' - s_r$. These band crossings occur at $q - 1$ discrete energies (right axes of Fig. 3), with exactly $q$ crossings having the same $\ell$ at each energy. For finite $\lambda$, $t_z$ hybridizes the states at each crossing, leading to $q(q - 1)$ avoided crossings and $\ell$...
gaps in the band structure. Near each degeneracy point the system is well described by the effective Hamiltonian [19):

$$\hat{H}_2(k) = \Delta \left[ \sin(k_x \ell) \hat{\sigma}_x + \sin(k_y \ell) \hat{\sigma}_y \right] + \nu \left( k_x^2 - k_y^2 \right) \hat{\sigma}_z.$$  (7)

where \(\Delta\) and \(\nu\) are the Pauli matrices in the basis of localized states \(|r\rangle\) and \(|r'\rangle\), and the \(k\)-independent constants are \(\Delta = (-1)^{r_{2n+1}} \sqrt{2 \left( E_r(k_x^2) + E_r(k_y^2) \right) - E_{R}(k_x^2)} \) and \(\nu = 2 \lambda \left[ \sin(2 \pi \alpha r + k_x^2) - \sin(2 \pi \alpha r' + k_y^2) \right] \). The eigenstates are Chern dimers, superpositions of \(|r\rangle\) and \(|r'\rangle\) with wave function

$$\psi_{n_r} = \frac{1}{\sqrt{2}} \left( \delta_{n_r,r} + e^{i \beta_n} \delta_{n_r,r'} \right).$$  (8)

where \(\beta_+ = -\ell (k_x + \pi) + \pi\) and \(\beta_- = -\ell (k_x + \pi)\) are relative phases for the upper and lower band edges, respectively. Starting from Eq. (7), it is straightforward to show [19] that the Chern number of each band gap is \(c_r = \ell\). Thus, for sufficiently anisotropic hopping, dimerized states form at the band edges with the spatial extent along \(c_r\) equal to the Chern number of the corresponding gap.

The formation of dimers of size \(\ell = c_r\) at the edges of the \(r\)th gap implies a “hidden spatial correlation” described by correlation function \(C(l) = \sum_{n_r} \langle f_{l+n_r} \rangle \), which is peaked at \(l = \ell\). For chemical potential in the \(r\)th gap, \(C(l)\) asymptotes to a delta function \(C(l) \rightarrow \delta(l - \ell), \) as \(\lambda \rightarrow \infty\). Because \(\beta_+ - \beta_- = \pi\) for each crossing, the net contribution to \(C(l)\) from dimerized states associated with gaps fully below the Fermi energy is zero. Since \(C(l)\) and \(n(k)_r\) are fourier transform pair, \(n(k)_r = \sum_{l} C(l) \cos(c_r k_l) / \sqrt{N}\); this yields the \(\cos(c_r k_l)\) term in Eq. (6). The \(r\) offset in Eq. (6) comes from the \(C(l = 0)\) contribution.

V. EFFECT OF TEMPERATURE AND TRAP

Our numerical simulation based on Eq. (1) confirms that the structures in \(n(k)_r\) persist, and remain visible, for trapped systems at finite temperatures. Due to the finite band gap of the quantum Hall insulator, thermal fluctuations only slightly smear the peak (or oscillation) structure in \(n(k)_r\) as long as \(T\) is small compared to the gap of interest, which is of the order \(t/2\) for the major gaps in the Hofstadter spectrum. Figure 4 shows examples of \(n(k)_r\) for realistic trap and temperature parameters. Thus, our proposal of detecting the Chern number is experimentally feasible.

VI. \(Q\)

Finally, we point out a general connection between \(\tilde{n}(k)\) and the \(Q\) matrix [20], which plays a fundamental role in the classification of topological insulators and in turn is related to the Chern number of a quantum Hall insulator. For a band insulator with band structure \(E_{\alpha}(k)\) (with energy measured from the chemical potential), the spectral projection operators \(P_\alpha(k)\) and \(P_- (k)\) project onto states above or below the chemical potential, respectively. The \(Q\) operator \(Q(k) = P_+(k) - P_- (k)\) describes a “flattened” Hamiltonian that is smoothly deformed from the original [20]. The momentum distribution is related to the spectral projection operator of the filled bands \(\rho(k) = P_- (k)\). In matrix form, \(\rho\) is related to \(Q\) by

$$Q_{ab} (k) = \delta_{ab} - 2 \rho_{ab} (k).$$  (9)

As an example, for the Hofstadter model at flux \(\alpha = p/q, \rho_{ab}(k) = \langle f_{a}(k) f_{b} (k) \rangle \), where \(a, b \in \{1, \ldots, q\} \) are the site indices within the magnetic unit cell and \(k\) is the crystal momentum within the MBZ. \(\tilde{n}(k)\) as defined in Eq. (3) is simply \(\tilde{n}(k) = \sum_{ab} \rho_{ab} (k)\). By Eq. (9), for given \(k_x = k_x^*\), the behavior of \(\tilde{n}(k_x, k_x^*)\) as a function of \(k_x\) is determined by the \(Q\) matrix along cut \(k_y = k_y^*\) in the BZ. So is the 1D momentum distribution \(n(k_x)\). On the other hand, the Chern number is nothing but the winding number of the off-diagonal block of the \(Q\) matrix in the chiral basis, also known as the transition function, along a cut that splits the BZ [20]. Thus, on general grounds, phase winding responsible for a nonzero
Chern number implies nontrivial structures in the momentum distribution function. Explicitly, for the asymptotically exact effective Hamiltonian $\hat{H}_2(k)$, the $Q$ matrix is $\hat{Q}(k) = (v^2 k_x^2 + \Delta^2)^{-1/2} \hat{H}_2(k)$. The Chern number $c_r = \ell$, for the off-diagonal block of the $Q$ matrix is simply $\sim e^{ik_x \ell}$. Meanwhile, from the $Q$ matrix, we obtain $n(k_x) = q[1 - \text{sgn}(\Delta x) \cos(k_x \ell)]$. This proves analytically our previous result Eq. (6). The oscillations adiabatically evolve into local peaks for the isotropic lattice, as explicitly shown in Figs. 2(c) and 2(d).

In contrast with Chern numbers, $n(k_x)$ is not a topologically invariant quantity. Still, the clear signatures of Chern numbers in TOF images that we find here make momentum distributions into an unexpected tool for exploring topological states of matter. In addition to edge states and Hall conductance, ripples in the momentum distribution and dimerized states of anisotropic lattices whose spatial extent encodes the topological quantum numbers provide a new way to characterize and visualize nontrivial topological states. Our study opens the possibility of using momentum distributions to explore other topological states of matter, including topological superfluids.

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