Topological flat bands in a kagome lattice multiorbital system

Satoshi Okamoto, Narayan Mohanta, Elbio Dagotto & D. N. Sheng

Flat bands and dispersive Dirac bands are known to coexist in the electronic bands in a two-dimensional kagome lattice. Including the relativistic spin-orbit coupling, such systems often exhibit nontrivial band topology, allowing for gapless edge modes between flat bands at several locations in the band structure, and dispersive bands or at the Dirac band crossing. Here, we theoretically demonstrate that a multiorbital system on a kagome lattice is a versatile platform to explore the interplay between nontrivial band topology and electronic interaction. Specifically, here we report that the multiorbital kagome model with the atomic spin-orbit coupling naturally supports topological bands characterized by nonzero Chern numbers \( C \), including a flat band with \( |C| = 1 \). When such a flat band is 1/3 filled, the non-local repulsive interactions induce a fractional Chern insulating state. We also discuss the possible realization of our findings in real kagome materials.
Flat-band systems have been proposed as interesting theoretical models to prove the existence of ferromagnetic ordering with itinerant electrons. Theoretical developments in such flat-band systems have been made almost in parallel with those in the widely discussed topological insulators (TIs). The nontrivial topology of electronic bands in a kagome lattice, one of those flat-band systems, has been extensively studied.

The experimental quests for topological materials with kagome lattice have also been carried out. Many of such experimental efforts were stimulated by the prediction of Weyl semimetals, including intermetallic compounds involving Co$_3$Sn$_2$Te$_2$, Fe$_5$Se$_2$, and Mn$_3$B$_2$C, and van-der-Waals compounds. More recently, the coexistence of superconductivity and nontrivial band topology was reported in a kagome compound.

When a flat band is partially occupied by electrons, the Coulomb repulsive interactions could become dominant over the electronic kinetic energy. This situation is already realized in two-dimensional electron gases under applied magnetic fields, where flat bands correspond to Landau levels. Fractional quantum Hall (FQH) effects were thus discovered. An exact numerical analysis made an important contribution by demonstrating that quantum fluctuations are essential to stabilize FQH states over charge density wave states. Once the charge excitation gap is induced at a fractional filling, the property of FQH states is elegantly explained using effective theory.

Recently, further intriguing proposals were put forward by considering flat bands with nontrivial topology and repulsive interactions, whereby FQH states could be generated without having Landau levels, called fractional Chern insulators (FCIs). These proposals considered single-band models on a kagome lattice, checkboard lattices, a Haldane model on a honeycomb lattice, and a ruby lattice as well as multi-band models on a buckled honeycomb lattice, a triangular lattice, and a square lattice for the mercury-telluride TI. It was later revealed that quantum Hall states realized in flat band systems and those realized under an applied magnetic field are adiabatically connected.

When realized in real materials, FCI states have several flat bands with $(yz, x^2 - y^2)$ or $3z^2 - r^2$ character. We focus on a $(yz, xz)$ subset for simplicity and use $a = a$ for the $yz$ orbital and $b$ for the $xz$ orbital. With this basis, nearest-neighbor hopping integrals $t_{\sigma \sigma}$ can be parameterized using Slater integrals. Between site 1 and site 2, $t_{12}$ is diagonal in orbital indices as $t_{12}^{za} = t_z$ and $t_{12}^{zb} = t_x$, corresponding to $(d\bar{d})$ and $(d\bar{d})$, respectively, by Slater and Koster. Other components are obtained by rotating the basis $a$ and $b$ as shown in the Methods section. From now on, $t_{\sigma}$ is used as the unit of energy.

Because $yz$ and $xz$ are written using the eigenfunctions of angular momentum $l_z = \pm 1$ for $l = 2$ as $|yz\rangle = \frac{1}{\sqrt{2}}(|1\rangle + |-1\rangle)$ and $|xz\rangle = -\frac{1}{\sqrt{2}}(|1\rangle - |-1\rangle)$, respectively, the SOC $\lambda \vec{r} \cdot \vec{s}$ in the $(yz, xz)$ subset is written as

$$H_{soc} = \lambda \sum_{\sigma \sigma'} \left( \sigma_{\alpha \beta}^{\sigma \sigma'} \sigma_{\alpha \beta}^{\sigma \sigma'} + H.c. \right).$$

where $\sigma^z$ is the $z$ component of the Pauli matrices.

As shown in Supplementary Note 1, an effective model for the $(xy, x^2 - y^2)$ doublet has the same form as the above $H_1 + H_{soc}$. By symmetry, there is no hopping matrix between the $(yz, xz)$

**Results**

**Theoretical model.** To begin with, we set up a multi-orbital tight-binding model on a kagome lattice

$$H_1 = - \sum_{(rr') \sigma} \sum_{\sigma \sigma'} \left( t_{rr'}^{\sigma \sigma'} c_r^\dagger \sigma c_{r'} \sigma + H.c. \right),$$

as schematically shown in Fig. 1. Here, $c_r$ is the annihilation (creation) operator of an electron at site $r$, orbital $\sigma$, and with spin $\sigma = \uparrow$ or $\downarrow$. As discussed by Meier et al., CoSn-type kagome systems have several flat bands with $(yz, x^2 - y^2)$ or $3z^2 - r^2$ character. We focus on a $(yz, xz)$ subset for simplicity and use $a = a$ for the $yz$ orbital and $b$ for the $xz$ orbital. With this basis, nearest-neighbor hopping integrals $t_{\sigma \sigma}$ can be parameterized using Slater integrals. Between site 1 and site 2, $t_{12}$ is diagonal in orbital indices as $t_{12}^{za} = t_z$ and $t_{12}^{zb} = t_x$, corresponding to $(d\bar{d})$ and $(d\bar{d})$, respectively, by Slater and Koster. Other components are obtained by rotating the basis $a$ and $b$ as shown in the Methods section. From now on, $t_{\sigma}$ is used as the unit of energy.

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**Fig. 1** Schematics of our theoretical model. a) Kagome lattice with three sublattices, labeled 1, 2, and 3. The two arrows are lattice translation vectors $a_1, a_2$. b) Local orbitals $a = yz$ and $b = xz$. Colored ellipsoids indicate regions of electron wave functions, where the sign is positive. c) Nearest-neighbor hopping integrals. $yz$ orbitals between site 1 and site 2 are hybridized via diagonal hopping $t_{\sigma \sigma}$, i.e., $\delta(z)$ bonding. Other hopping integrals between site 2 and site 3 and between site 1 and site 3 are obtained via the Slater rule as shown in the Methods section.
doublet and the other orbitals $xy$, $x^2 - y^2$, and $3z^2 - r^2$, but the \{xy, x^2 - y^2\} doublet and the $3z^2 - r^2$ singlet could be hybridized. As discussed briefly later, the degeneracy in the \{yz, xz\} doublet and in the \{xy, x^2 - y^2\} doublet could be lifted by a crystal field. Such band splitting is also induced by the difference between $t_6$ and $t_6$. Furthermore, all $d$ orbitals could in principle be mixed by the SOC. Including these complexities is possible but depends on the material and they usually induce smaller perturbations, therefore, here they are left for future analyses.

**Non-interacting band topology.** By diagonalizing the single-particle Hamiltonian $H_0 + H_{soc}$, one obtains dispersion relations as shown in Fig. 2. In the simplest case, where the hopping matrix $t_{rr}'$ does not distinguish $t_6$ and $t_6$ and the SOC is absent, the dispersion relation is identical to the one for the single-band tight-binding model, consisting of flat bands and graphene-like bands as shown by gray lines in Fig. 2a. Note that each band is fourfold degenerate because of two orbitals and two spins per site. Including SOC does not change the dispersion curve but simply shifts $\vec{T} \cdot \vec{s} = \pm 1/2$ bands (see Supplementary Note 1).

Including orbital dependence as $t_6 \neq t_6$ without SOC instead splits the fourfold degeneracy except for two points at the $\Gamma$ point and two points at the $K$ point. Quite intriguingly, Dirac dispersions emerge from the topmost flat bands as shown by blue lines in Fig. 2a (see Supplementary Note 1 for more discussion). Turning on the SOC further splits such fourfold degeneracy, leading to nontrivial band topology. In this particular example, the spin component along the $z$ axis is conserved giving unique characteristics to this case. As shown in Fig. 2b, the spin up component of each band is characterized by a nonzero Chern number $C_{\uparrow}$. Because of the time-reversal symmetry, spin down bands have opposite Chern numbers. The topological property is also confirmed by gapless modes in the dispersion relation with the ribbon geometry, as shown in Fig. 2c. Here, there appear one (two) pair of gapless modes between the highest and the second highest (between the second lowest and the third lowest) bands, shown as red (blue) curves, corresponding to the sum of Chern numbers below the gap, $-1(-2)$. The other edge states are invisible because of the overlap with the bulk continuum.

A multi-orbital kagome model thus naturally shows quasi flat bands with nontrivial topology. However, close inspection revealed that, with $t_6 = 0.5$ and $\lambda = 0.2$, the minimum of the highest band at the $K$ point is slightly lower than the maximum of the second highest band at the $\Gamma$ point. Thus, instead of a TI, a topological semimetal is realized when the Fermi level is located between the highest band and the second highest band. In fact, there are ways to make the gap positive. Here, we consider second-neighbor hopping matrices $t_{rr}^{(2)}$. As explained in Supplementary Note 1, these are also parametrized by $s$-bonding (dd) and $d$-bonding (ddb), $t_{rr}^{(2)}$ and $t_{rr}^{(3)}$, respectively. For simplicity, we fix the ratio between $t_{rr}$ and $t_{rr}^{(2)}$ and between $t_6$ and $t_{rr}^{(2)}$ as $t_{rr}^{(2)}/t_{rr}^{(2)} = t_6/t_6 = r_2$, and analyze the sign and magnitude of the band gap $\Delta_{gap}$ between the highest band and the second highest band, as well as the flatness of the highest band defined by $\Delta_{flat} \equiv \epsilon_{min} - \epsilon_{max}$.

Figure 3a plots $\Delta_{flat}$ as a function of $t_6$ and $r_2$ with $\lambda = 0.2$. As mentioned previously, the perfectly flat band with $\Delta_{flat} = 0$ is realized at $t_6 = 1$ and $r_2 = 0$, but band gap $\Delta_{gap}$ is zero. The flatness is immediately modified by reducing $t_6$ from 1. As indicated by an open square in the plot, $t_6 = 0.5$ and $r_2 = 0$ gives $\Delta_{flat} \sim 0.88$ and negative band gap $\Delta_{gap} \sim -0.027$. Nonzero $r_2$ controls the relative energy between the zone center and the zone boundary. In particular, negative $r_2$ pushes up the energy at the $K$ point, hereby the flatness is recovered. Naturally, the flatness and the positive gap are correlated as indicated by red loops in the second and fourth quadrants because the separation between the highest band and the second highest band is fixed by the SOC strength. As indicated by a filled circle, $t_6 = 0.5$ and $r_2 = -0.2$ gives $\Delta_{flat} \sim 0.22$ and positive band gap $\Delta_{gap} \sim 0.17$. Corresponding dispersion relation is shown in Fig. 3b. The Chern numbers remain unchanged by this $r_2$.

**Many-body effects.** Having established the topological properties at the single-particle level, we turn our attention to many-body effects focusing on the highest-energy flat band. A unique property of the current model is that the topmost quasi flat band has Chern number $|C| = 1$. Thus, a large spin polarization can be induced by many-body interactions$^{27}$ or by a small magnetic field. Further intriguing possibilities are FCI states when a topological flat band has a fractional filling and the insulating gap
is induced by correlation effects\textsuperscript{44,46–52}. We examine such a possibility in our kagome model. Assuming the spin polarization in the highest band, we introduce local and nearest-neighbor Coulomb repulsive interactions as $H_{U} = U\sum_{l_{1}l_{2}l_{3}}n_{l_{1}}n_{l_{2}} + V_{U}(\text{tr})\sum_{l_{1}l_{2}}n_{l_{1}}n_{l_{2}}$, where $n_{l_{1}} = \psi_{l_{1}}^{\dagger}\psi_{l_{1}}$. Here $U$ is the effective Coulomb interaction given by $U = U^\prime - J$ with the interorbital Coulomb repulsion $U^\prime$ and the interorbital exchange interaction $J$. These interactions are then projected onto the highest band, leading to the effective Hamiltonian $H_{\text{eff}} = H_{I} + H_{\text{soc}} + H_{U}$.

Note that the $S_{z}$ conservation is not essential to realize FCI. For our case and most of others, including complexities which break $S_{z}$ conservation does not destroy FCI as long as the flat band has the nontrivial topology and is well separated from other bands, justifying projecting interaction terms onto the flat band and allowing for an accurate Lanczos calculation. While the computational cost would be expensive, direct calculations of multiband models with $S_{z}$-non-conserving terms would show FCI if the appropriate condition is fulfilled, but this possibility has not been fully explored yet.

The effective Hamiltonian $H_{\text{eff}}$ is diagonalized in momentum space. For this purpose, we discretize the momentum space into $N_{1} \times N_{2}$ patches and express the Hamiltonian in the occupation basis, i.e., the Hilbert space is built up by $|\psi_{l}\rangle = \prod_{l_{k}}|\psi_{l_{k}}\rangle|0\rangle$, where $\psi_{l_{k}}$ is the single-particle wave function for the highest flat band at momentum $k$, and the combination of $k$ is specified by $l$. Due to the translational symmetry and the momentum conservation of many-body interaction terms, $H_{\text{eff}}$ is subdiagonalized according to the total momentum $k_{\text{tot}} = \sum_{k} l$ modulo $b_{1}$ and $b_{2}$, with $b_{1,2}$ being two reciprocal lattice vectors. In this study, we take $N_{1} = 4$ and $N_{2} = 6$ and consider $\nu = 1/3$ filling, that is, the number electrons in the highest flat band is $N_{e} = 8$. Momentum sector will be specified using integer index $(k_{1}, k_{2})$ corresponding to the total momentum $k_{\text{tot}} = b_{1}k_{1}/N_{1} + b_{2}k_{2}/N_{2}$.

Figure 4a, b show the low-energy spectra of the interacting model with $t_{3} = 0.5$ and $r_{2} = 0$ and $t_{3} = 0.5$ and $r_{2} = -0.2$, respectively, with $U = 2$ and $V = 1$ as a function of total momentum. In (a), the energy spectrum has a unique ground state at total momentum $(k_{1}, k_{2}) = (0, 0)$ (note that this is to show the competition between the wide band width and the correlation effects on the highest band). When $r_{2}$ is introduced as $-0.2$, the highest band becomes flatter, leading to a drastic change in the energy spectrum. There appear three energy minima at $(0, 0)$, $(0, 2)$, and $(0, 4)$, respectively.
the numerical accuracy. The slight deviation from the ideal value and the sum of the three Chern numbers is exactly 1 within
Cλ are periodic. Along the in a discretized grid (n₁, n₃) for the GSM with tₙ = 0.5, r₂ = −0.2, λ = 0.2 with U = 2 and V = 1. Along the n₁ direction, these plots are periodic. Along the n₃ direction, plot (a) is continuously connected to plot (b), plot (b) is connected to plot (c), and plot (c) is connected back to plot (a). This also confirms the threefold GSM, where inserting one flux quantum along the b₂ direction shifts the sector (k₁, k₃) = (0, 0) to (0, 2), (0, 2) to (0, 4), and (0, 4) to (0, 0). By adding up the discretized values of F(k₁, k₃)(θ₁, θ₂), we obtain C(0,0) = 0.331489, C(0,2) = 0.330318, C(0,4) = 0.338193, and the sum of the three Chern numbers is exactly 1 within the numerical accuracy. The slight deviation from the ideal value C = 1/3 is ascribed to finite-size effects. This proves the existence of a v = 1/3 FCI phase with a quantized fractional Hall response σ₁∝ = 1/3e²/h, where e is the electron charge and h is the Planck constant. In our numerical analyses, we did not find a ground state with the threefold degeneracy and Chern number zero, thus excluding the charge density wave states. This is probably because the quantum effects make such states unstable, as discussed in ref. 42.

Discussion
In this work, we have considered an itinerant electron model on a kagome lattice with twofold degenerate orbitals per site. However, each site has C₂ rotational symmetry, rather than C₃ or C₄. Thus, the degeneracy between the two orbitals (yz and xz) can be lifted. In our tight-binding model, a difference in the hopping amplitude between tₙ and tₙs in fact lifts such degeneracy, leading to the splitting of the band structure. Thus, adding local crystal field splitting, which respects the underlying lattice symmetry, would not fully destroy the topological property found in this work, while the position of topological or flat bands would be modified depending on model parameters. As a number of kagome materials and tuning the Fermi level to a topological flat band by chemical substitution or gating, might be a promising route to observe the phenomena predicted here. The sign and the magnitude of the parameter t₂ could depend on details of the material, such as the species of ligand ions, and might be further controlled by compressive or tensile strain. First principles calculations would help to construct realistic material-dependent models. It is anticipated that the separation between {yz, xz}, {xy, x²−y²}, or 3z²−r² subsets will be enhanced by reducing the film thickness compared with that in the bulk so that one can focus on one of the subsets only. In addition to a kagome lattice, topological flat bands appear in dice and Lieb lattices. Study of FCI states in such lattice geometries and material search is another important direction.

To summarize, we have demonstrated the close interplay between the spatial frustration and the orbital degree of freedom in a kagome lattice. With the relativistic SOC, such an interplay not only affects the band dispersion, but also induces nontrivial topology. Specifically, we showed that the original flat bands in a kagome lattice become dispersive and topologically nontrivial. When such topological bands are fractionally occupied by electrons, many-body interactions drive further intriguing phenomena, i.e., fractional Chern insulating states. Our work may bridge the gap between idealized theoretical studies and real materials.

Methods
Non-interacting (yz, xz) model. Here we deduce the hopping matrices of the (yz, xz) model in the Slater–Koster approximation.
For nearest-neighbor bonds, in addition to the diagonal matrix k₆ in the main text, we have

\[ i_{13}^{(2)} = \frac{1}{4} \begin{bmatrix} 0 & 3t_{x} + 3t_{y} & \sqrt{3}(t_{x} - t_{y}) \\ \sqrt{3}(t_{x} - t_{y}) & 3t_{x} - 3t_{y} & 0 \\ t_{x} + 3t_{y} & \sqrt{3}(t_{x} - t_{y}) & 0 \end{bmatrix} \]

Similarly, second neighbor hopping matrices can be written as

\[ i_{13}^{(2)} = \frac{1}{4} \begin{bmatrix} 0 & 3t_{x} + 3t_{y} & \sqrt{3}(t_{x} - t_{y}) \\ \sqrt{3}(t_{x} - t_{y}) & 3t_{x} - 3t_{y} & 0 \\ t_{x} + 3t_{y} & \sqrt{3}(t_{x} - t_{y}) & 0 \end{bmatrix} \]

where subscript (2) is introduced to highlight the difference from the nearest-neighbor bonds. These are schematically shown in Fig. 6. \( i_{ij}^{(2)} \) and \( i_{ij}^{(3)} \) correspond to \( (dHH) \) and \( (ddHH) \), respectively, by Slater and Koster.
Fig. 6 Second neighbor hopping matrices. \(yz(xz)\) orbitals between site 1 and site 2 are hybridized via diagonal hopping \(t^{(2)}_{23}\), i.e., \(\sigma(x)\) bonding. Other hopping integrals between site 1 and site 3 and between site 2 and site 3 are given by \(t^{(2)}_{13}\) and \(t^{(2)}_{23}\), respectively, obtained via the Slater rule\(^{6}\).

**Non-interacting Berry curvature.** The band-dependent Berry curvature of non-interacting electrons is given as a function of momentum \(\mathbf{k}\) as

\[
\Omega_{\mathbf{k}} = -i \sum_{\{\mathbf{m}\}} \left\langle \hat{\mathbf{v}}^\dagger_{\mathbf{m}} \hat{\mathbf{v}}_{\mathbf{m}} \right\rangle \left( \epsilon_{\mathbf{k}} - \epsilon_{\mathbf{m}} \right)^2,
\]

where, using the Hamiltonian matrix in momentum space \(\hat{H}_{\mathbf{k}}\), \(\hat{\mathbf{v}}_{\mathbf{k}}\) is given by

\[
\hat{\mathbf{v}}_{\mathbf{k}} = \partial \hat{H}_{\mathbf{k}} / \partial \mathbf{k}.\]

With this Berry curvature, the band-dependent Chern number \(C_{\mathbf{k}}\) is given by

\[
C_{\mathbf{k}} = \frac{1}{2\pi} \int d^2 k \Omega_{\mathbf{k}}.
\]

where the momentum integral is taken in the first Brillouin zone.

**Many-body Chern number.** The many-body Chern number is computed by introducing a twist boundary condition to a single-particle wave function as \(\psi(\mathbf{r} + N_{\mathbf{a}}) = e^{i\theta}(\mathbf{r})\), where \(N_{\mathbf{a}}\) are the numbers of unit cells along lattice translation vectors \(\mathbf{a}_{m,n}\), with phase factors \(\theta_{m,n}\). This corresponds to inserting magnetic fluxes. When one flux quantum is inserted, \(\theta_{1,1}\) changes from 0 to 2\(\pi\) and discretized momentum \(\mathbf{k}\) moves from its original position to its neighbor along the \(\mathbf{b}\) direction with the momentum shift given by \(\Delta \mathbf{k} = bN_{\mathbf{b}}/N_{\mathbf{a}}\).

Many-body Chern number of the ground state \((\mathbf{k}_1, \mathbf{k}_2)\) is computed via

\[
C_{(\mathbf{k}_1, \mathbf{k}_2)} = \frac{1}{2\pi} \int_{0}^{\mathbf{b}} d\mathbf{b}_1 \int_{0}^{\mathbf{b}_2} d\mathbf{b}_2 F_{(\mathbf{k}_1, \mathbf{k}_2)}(\theta_1, \theta_2)\]

where \(F(\theta_1, \theta_2)\) is the Berry curvature given by

\[
F_{(\mathbf{k}_1, \mathbf{k}_2)}(\theta_1, \theta_2) = \text{Im} \left\langle \frac{\partial \Phi_{(\mathbf{k}_1, \mathbf{k}_2)}}{\partial \theta_1} \frac{\partial \Phi_{(\mathbf{k}_1, \mathbf{k}_2)}}{\partial \theta_2} - \frac{\partial \Phi_{(\mathbf{k}_1, \mathbf{k}_2)}}{\partial \theta_2} \frac{\partial \Phi_{(\mathbf{k}_1, \mathbf{k}_2)}}{\partial \theta_1} \right\rangle.
\]

Here, \(\partial \Phi_{(\mathbf{k}_1, \mathbf{k}_2)} / \partial \theta_{1,2}\) is the many-body wave function obtained using single-particle wave functions with a twist boundary condition \(\psi(\mathbf{r})\) after the Fourier transformation to momentum space. The momentum index \((\mathbf{k}_1, \mathbf{k}_2)\) will be omitted in the following discussion for simplicity.

Partial derivative of a wave function with respect to \(\theta_{1,2}\) is approximated by a finite difference as

\[
\frac{\partial \Phi(\theta_1, \theta_2)}{\partial \theta_{1,2}} \approx \frac{1}{\Delta \theta_{1,2}} \left[ \Phi(\theta_1 + \Delta \theta_{1,2}, \theta_2) - \Phi(\theta_1, \theta_2) \right].
\]

Here, the vector notation is used for \(\theta = (\theta_1, \theta_2)\), and \(\Delta \theta = (\Delta \theta_1, 0)\) or \((0, \Delta \theta_2)\). Then, it is required to compute a product of two wave functions as \(\Phi(\theta) (\Phi(\theta'))\) with \(\theta = \theta'\). Because we are using a multilayered model projected onto the flat band, special care is needed, as detailed in Supplementary Note 2.

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Author contributions

S.O. designed the research and carried out numerical calculations and wrote the manuscript with the input from all the authors. N.M. supported the construction of the model Hamiltonian. E.D. and D.N.S. supported many-body numerical calculations.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to Satoshi Okamoto.

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