Topological Insulators from Electronic Superstructures

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The possibility of realizing topological insulators by the spontaneous formation of electronic superstructures is theoretically investigated in a minimal two-orbital model including both the spin-orbit coupling and electron correlations on a triangular lattice. Using the mean-field approximation, we show that the model exhibits several different types of charge-ordered insulators, which may lead to new types of topological phases, such as a charge density wave, which brings about a topological insulator (TI).

Quantum phenomena originating from the geometrical properties of electronic wave functions have been a central issue in modern condensed matter physics. The study of the anomalous Hall effect in ferromagnetic metals has revealed that the relativistic spin-orbit coupling (SOC) plays a crucial role in such phenomena.1) In particular, the strong SOC may bring about intriguing topological states of matter, such as the topological insulator (TI).2–4) The TI is a nontrivial band insulator, which is distinguished from conventional ones by a Z2 topological invariant under the time-reversal symmetry. It exhibits a peculiar metallic edge (or surface) state, which gives rise to the quantized spin Hall conductivity. Their band gap is dependent on electron correlations as well as the spin-orbit coupling, and even vanishes while showing the massless Dirac dispersion at the transition to a trivial charge-ordered insulator. Our results suggest a new route to realize and control topological states of quantum matter by the interplay between the spin-orbit coupling and electron correlations.

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is given by

$$H_{\text{ onsite}} = \frac{1}{2} \sum_{mnn' \sigma} U_{mnn'} \sum_{i} \sum_{\sigma'\sigma} c_{i\sigma}^\dagger c_{i\sigma'} c_{i\sigma'}^\dagger c_{i\sigma}. \quad (3)$$

Assuming the rotational symmetry of the Coulomb interaction, we set $U_{mnn'} = U$, $U_{mnnm} = U - 2J$, and $U_{mnnm} = U_{mnm} = J (m \neq n)$, where $U$ is the intraorbital Coulomb interaction and $J$ is the Hund’s coupling, respectively. We set $U = 1.0$ and $J/U = 0.1$ in the following calculations. For the intersite interaction, we consider the density-density repulsions given by

$$H_{\text{ intersite}} = V_1 \sum_{\langle i,j \rangle} n_i n_j + V_2 \sum_{\langle (i,j) \rangle} n_i n_j, \quad (4)$$

where $n_i = \sum_{\sigma} c_{i\sigma}^\dagger c_{i\sigma}$. The sum of $(i, j)$ $(\langle (i, j) \rangle)$ is taken for the nearest- (next-nearest-) neighbor sites.

To clarify the ground states of the two-orbital model given by Eqs. (1), (3), and (4), we use the mean-field approximation. In the mean-field calculation, we employ 12 sublattices [see Figs. 3(c) and 5(c)] and approximate the integration in the folded Brillouin zone by the summation over 64 × 64 $k$ points. We apply the Hartree-Fock approximation to the onsite interaction in Eq. (3) and the Hartree approximation to the intersite interaction in Eq. (4) to focus on charge order.\(^{18}\)

The mean fields are determined self-consistently, until they converge within a precision of less than $10^{-6}$. We investigate the ground state and find several interesting charge orders at the intersite interaction in Eq. (4) to focus on charge order.\(^{18}\)

In addition, we compute the spin Hall conductivity, which signals the nontrivial topological nature of the system; it can be quantized at a nonzero integer multiple value of $e/2\pi$ for two-dimensional TIs\(^{21}\) ($e$ is the elementary charge). Using the standard Kubo formula in the linear response theory, we calculate the spin Hall conductivity as

$$\sigma_{xy}^s = \frac{e}{2} \frac{1}{\Omega} \sum_{k,j,l} \frac{f(\epsilon_{ak}) - f(\epsilon_{lj})}{\epsilon_{ak} - \epsilon_{lj}} \frac{(\epsilon_{lj} + i\alpha k\hat{\kappa})\langle \hat{a}_{kj}\rangle\langle \hat{b}_{lk}\rangle\langle \hat{a}_{kj}\rangle\langle \hat{a}_{lj}\rangle}{\epsilon_{ak} - \epsilon_{lj} + \delta}, \quad (5)$$

where $\Omega$ is the system volume, $f$ is the Fermi distribution function, $\epsilon_{ak}$ and $|\alpha k\rangle$ are the eigenvalues and eigenvectors of the $\alpha$th electronic band with the wave vector $k$ in the mean-field solution, respectively, and $\delta$ is the infinitesimal positive parameter. In the calculation of Eq. (5), we take the summation over $512 \times 512$ $k$ points and set $T = 10^{-3}$ and $\delta = 10^{-3}$. The current operator is defined by $j_z = \partial H_{\text{MF}} / \partial \epsilon_z$, where $H_{\text{MF}}$ is the mean-field Hamiltonian. We define the spin current operator as $j_z^s = \langle j_z, \sigma_z \rangle / 2$, where $\sigma_z$ is the z-component of the Pauli matrices for spin; note that $H_{\text{MF}}$ for our mean-field solutions commutes with $\sigma_z$. Hereafter, we denote $\tilde{\sigma}_z^s \equiv \sigma_z^s / (e/2\pi)$ as the normalized spin Hall conductivity. We note that, for two-dimensional systems whose Hamiltonian commutes with $\sigma_z$, $\tilde{\sigma}_z^s$ is directly related to the $Z_2$ topological invariant.\(^{2}2\)

First, we consider the situation where $V_1$ is dominant rather than $V_2$, and thus, set $V_2 = 0$. In this case, the system shows

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**Fig. 1.** (Color online) (a) Schematic picture of edge-sharing octahedra. The large (red) spheres inside the octahedra denote the transition metal cations and the small (gray) ones on the vertices indicate the ligand ions. (b) Atomic $d$ orbital levels of the transition metal cations under the octahedral and trigonal crystalline electric fields corresponding to (a). The $d_{\alpha\beta}$ orbitals are further split by the SOC; see the text for details. (c) Schematic picture of the triangular lattice of transition metal cations. $\eta_1, \eta_2$ (or $\eta_1, \eta_2, \eta_3$) and spin $\sigma$ are the orbital levels of the transition metal cations under the octahedral and trigonal crystalline electric fields corresponding to (a). The $d_{\alpha\beta}$ orbitals are further split by the SOC; see the text for details. (c) Schematic picture of the triangular lattice of transition metal cations. $\eta_1, \eta_2$ (or $\eta_1, \eta_2, \eta_3$) and spin $\sigma$ are the orbital levels of the transition metal cations under the octahedral and trigonal crystalline electric fields corresponding to (a). The $d_{\alpha\beta}$ orbitals are further split by the SOC; see the text for details. (d) Energy levels and hopping processes in the two-orbital model in Eq. (1).

\[ \frac{1}{2} \sum_{m,m'=+1} (m\sigma) c_{km\sigma}^\dagger c_{km'\sigma}, \quad (1) \]

where $c_{km\sigma}^\dagger (c_{km\sigma})$ is the creation (annihilation) operator of an electron for the wave vector $k$, orbital $m = \pm 1$, and spin $\sigma = \pm 1$. $t_0$ and $t_1$ are the intra- and interorbital hopping elements between nearest-neighbor sites, respectively. The factors $\gamma_{\alpha k}$ ($\alpha = 0, \pm 1$ in the mean fields in Eq. (1) are given by

\[ \gamma_{\alpha k} = \sum_{n=1,2,3} 2 c^{2(n-1)\eta_1} \cos (k \cdot \eta_n), \quad (2) \]

which originate from the direction-dependent overlaps between $|x\rangle$, $|y\rangle$, and $|z\rangle$ orbitals. Hereafter, we take the triangular plane as the $x'y'$ plane, and set the primitive translational vectors for the triangular lattice as $\eta_1 = (1, 0)$, $\eta_2 = (1/2, \sqrt{3}/2)$, and $\eta_3 = (-1/2, \sqrt{3}/2)$ [see Fig. 1(c)]. $\lambda$ is the SOC constant, which splits the energy levels of $|m, \sigma\rangle$ into two Kramers doublets, $|$1; 1, +1$\rangle$, $|$1; 1, -1$\rangle$, and $|$1; 1, -1, -1$\rangle$, and $|$1; 1, +1$\rangle$, $|$1; 1, -1$\rangle$, where $t_0 = t_1 = 0$, as shown in Fig. 1(b).

The energy levels and hopping processes are schematically shown in Fig. 1(d). Note that a similar model was studied on a honeycomb lattice.\(^{15,16}\) Although the honeycomb lattice model exhibits topologically nontrivial states even in the noninteracting case,\(^{16}\) our triangular lattice model in Eq. (1) does not for any values of the parameters $t_0$, $t_1$, and $\lambda$. Hereafter, we set $t_0 = 0.5$ and $t_1 = 0.25$, which are reasonable when considering the $d$-$d$ direct and $d$-$p$-$d$ indirect hoppings in the Slater–Koster scheme.\(^{17}\)

In addition to the one-body part, we take into account both the onsite and intersite Coulomb interactions. The onsite one

\[ + \frac{\lambda}{2} \sum_{m,m'=+1} (m\sigma) c_{km\sigma}^\dagger c_{km\sigma}, \quad (1) \]
an interesting behavior at 1/3 filling. Figure 2(a) shows the ground-state phase diagram obtained by the mean-field approximation while changing $\lambda$ and $V_1$. We find three different phases in this parameter region: paramagnetic metal (PM) for small $V_1$ and two charge-ordered insulators (COIs) for large $V_1$. In both COIs, the local charge density is disproportionated to form a honeycomb superstructure; in the COI in the larger $V_1$ region, the local charge density is lower at the sites belonging to the honeycomb network than at the isolated sites, while they are opposite in the COI in the intermediate $V_1$ and large $\lambda$ region [see the schematic pictures in Fig. 2(a)]. The local charge densities are plotted in Fig. 2(b). We call the former (latter) the honeycomb type-A(B) COI.

In the small $\lambda$ region, there is a transition from the paramagnetic metal to the honeycomb type-A COI with increasing $V_1$. This is easily understood by considering that the electrons tend to avoid each other under large $V_1$ and the lowest energy configuration at 1/3 filling is given by the type-A charge ordering. On the other hand, in the large $\lambda$ region, the type-B COI appears between the type-A COI and PM phases. The intervening type-B COI is stabilized by the synergy between the strong SOC and intersite Coulomb repulsion. This is understood by considering the large $\lambda$ limit as follows. As the bands for the Kramers pair are largely split from each other, the two-orbital model at 1/3 filling reduces to a single-band model at 2/3 filling for the lower-energy band. In the single-band model at 2/3 filling, the lowest energy state under $V_1$ is given by the type-B charge ordering, which explains why the type-B COI is stabilized in the large $\lambda$ and $V_1$ region in Fig. 2(a). We note that all the phase boundaries in Fig. 2(a) are of first order with discontinuous changes in local charge densities.

Figure 2(c) shows $\lambda$ dependences of the energy gap and normalized spin Hall conductivity at $V_1 = 0.6$. The results clearly indicate that both CO states are gapped insulators, while the type-A COI has a larger gap than the type-B COI in this parameter region. Remarkably, in the type-B CO state, the band gap once closes around $\lambda \sim 3.2$. The gapless line inside the type-B CO phase is shown by the dashed line in Fig. 2(a). The result indicates that the type-B CO phase may include two different insulating states separated by the gapless boundary. Indeed, as shown in Fig. 2(c), the normalized spin Hall conductivity in the type-B COI is quantized at $-1$ for $\lambda \lesssim 3.2$, while it changes discontinuously to zero when crossing the gapless point. Therefore, the gapless boundary in the type-B COI corresponds to a topological transition between a TI for smaller $\lambda$ and a trivial band insulator for larger $\lambda$.

To clarify the electronic states in the type-B COI further, we show the electronic band structure of the mean-field solution for the type-B COI near the gapless boundary in Figs. 3(a) and 3(b) [the unit cell and Brillouin zone are shown in Figs. 3(c) and 3(d), respectively]. As shown in Fig. 3(a), the Kramers doublets are split by the strong SOC into two ‘copies’ of three bands; the highest band in the lower three bands hybridizes with the lowest one in the higher three bands, resulting in a small gap at $\varepsilon \sim 1.1$. The three bands in each copy are composed of two subsets, reflecting the honeycomb CO superstructure; the lower two bands comprise the dispersive bands similar to those of the single-band model on the honeycomb lattice, and the remaining higher band is less dispersive as it comes from the isolated sites in the honeycomb hexagons. The lower honeycomb-like bands are occupied (the Fermi level is set at zero). This result supports the above discussion for the origin of the type-B COI.

Figure 3(b) shows more details of the band structures near the Fermi level at 1/3 filling around the $\Gamma$ point at $V_1 = 0.6$. With increasing $\lambda$, the band gap at 1/3 filling decreases and closes at $\lambda \sim 3.242$. In the gapless state, the low-energy dispersions are well approximated by the massless Dirac cone. The Dirac cone is gapped out again by further increasing $\lambda$. We note that, although this topological transition appears to share the fundamental mechanism with that found for the sim-
The kagome CO state is interesting from the topological viewpoint, as discussed below. In Fig. 4(b), we plot the $V_2$ dependence of the charge density at each sublattice, the band gap, and the normalized spin Hall conductivity at $\lambda = 0.6$.

Next, we take into account the next-nearest-neighbor repulsion $V_3$. We find that $V_3$ leads to different types of electronic superstructures around 1/3 filling. In particular, here, we discuss interesting CO states appearing at 3/8 filling. Figure 4(a) shows the ground-state phase diagram at $V_1 = 0.6$ while changing $\lambda$ and $V_2$. In the small $V_2$ region, the system exhibits a honeycomb type-A CO metal (COM) as well as PM, whose charge patterns are also seen in the 1/3 filling case above. When increasing $V_2$, we find two new CO phases: kagome and stripy CO phases [see the schematic picture in the phase diagram in Fig. 4(a)]. In the kagome CO state, charge density is disproportionate and so the charge-poor sites comprise a kagome superstructure, as plotted in Fig. 4(b). (The charge-poor sites have a very small charge disproportionation among them, which does not affect the following topological nature of this phase.) On the other hand, the stripy CO state has a four-sublattice order, where charge density is disproportionate into three groups: charge-rich, charge-poor, and intermediate at one, two, and one sublattices, respectively [see Fig. 4(b)].

The kagome CO state is intriguing from the topological viewpoint, as discussed below. In Fig. 4(b), we plot the $V_2$ dependence of the energy gap at $\lambda = 0.6$. The result shows that the kagome CO phase is metallic in the small $V_2$ region but becomes insulating with increasing $V_2$. The band gap is opened by the cooperation between the SOC and $V_2$ [see also Fig. 4(a)]. This is explicitly shown in the band structures for $V_2 = 0.15$ in Figs. 5(a) and 5(b) at $\lambda = 0.0$ and $\lambda = 0.6$, respectively [the unit cell and Brillouin zone are shown in Figs. 5(c) and 5(d), respectively]. Although the bands near the Fermi level do not have a gap at $\lambda = 0.0$, they are separated by a gap for $\lambda = 0.6$. We find that the kagome COI is a TI by calculating the normalized spin Hall conductivity, as shown in Fig. 4(b). Although the normalized spin Hall conductivity is already nonzero in the honeycomb type-A and kagome COM phases for smaller $V_2$, it is quantized at a nonzero integer number, $\Delta S_{xy} = -1.0$ in the kagome COI.

Finally, let us discuss our results. We found two different types of COIs which are topologically nontrivial: the honeycomb type-B and kagome COIs. The important physics here is the role of the SOC under the electronic superstructures. As remarked above, the noninteracting model including the SOC [Eq. (1)] does not exhibit any topological nature owing to the high symmetry of the triangular lattice. The formation of the honeycomb and kagome superstructures activates the hidden SOC effect and changes the system into TIs. This is, for instance, understood from the relationship between the electronic states of the honeycomb type-B COI in the present model and the PM in the honeycomb-lattice model studied in a previous work, as discussed above. The situation is distinct from other interaction-driven TIs, the so-called topological Mott insulators, where the atomic SOC does not play an important role.

Similar mechanisms activating the SOC effect by superstructure formation were discussed for spatial inversion symmetry breaking, which induces the antisymmetric SOC. Thus, our results point to a much broader route to activate the nontrivial SOC physics and realize topological states of matter, with the aid of the change of spatial symmetry by electronic correlations. This has richer implications, since, in addition to charge ordering, the superstructure formation can be caused by other degrees of freedom, e.g., magnetic ordering in spin-charge coupled systems and bond ordering in electron-phonon coupled systems. Interestingly, there are many candidate materials exhibiting various superstructures, e.g., delafossite-type oxides and transition metal dichalcogenides. In particular, the latter compounds are intriguing, as they show a variety of charge density waves with longer periodicities accompanied by lattice distortions. The topological nature in these interesting states with electronic superstructures is left for a future study.
Acknowledgment The authors thank Satoru Hayami, Hiroaki Kusunose, Takahiro Misawa, and Youhei Yamaji for constructive suggestions. Y.S. is supported by the Japan Society for the Promotion of Science through the Program for Leading Graduate Schools (MERIT). This work was supported by Grants-in-Aid for Scientific Research (Nos. 24340076 and 15K05176), the Strategic Programs for Innovative Research (SPIRE), MEXT, and the Computational Materials Science Initiative (CMSI), Japan.

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18) Imaginary terms from the Fock decoupling of the intersite interactions may prefer an interaction-driven TI, the so-called topological Mott insulator.\textsuperscript{9,20–22} In this study, however, we focus on the charge ordering as a trigger for TIs. We confirmed that the result remains qualitatively the same, when adopting the three-site unit cell as shown in Fig. 3(c) and the Hartree-Fock approximation for the intersite interactions in the case of 1/3 filling.
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