Silicene is a monolayer of silicon atoms forming a two-dimensional honeycomb lattice, which shares almost every remarkable property with graphene. The low energy structure of silicene is described by Dirac electrons with relatively large spin-orbit interactions due to its buckled structure. The key observation is that the band structure is controllable by applying electric field to silicene. We explore the phase diagram of silicene together with exchange field $M$ and by applying electric field $E_z$. There appear quantum anomalous Hall (QAH) insulator, valley polarized metal (VPM), marginal valley polarized metal (M-VPM), quantum spin Hall (QSH) insulator and band insulator (BI). They are characterized by the Chern numbers and/or by the edge modes of a nanoribbon. It is intriguing that electrons have been moved from a conduction band at the K point to a valence band at the K' point for $E_z > 0$ in the VPM. We find in the QAH phase that almost flat gapless edge modes emerge and that spins form a momentum-space skyrmion to yield the Chern number. It is remarkable that a topological quantum phase transition can be induced simply by changing electric field in a single silicene sheet.

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Silicene consists of a honeycomb lattice of silicon atoms with two sublattices made of A sites and B sites. The states near the Fermi energy are $\pi$ orbitals residing near the K and K' points at opposite corners of the hexagonal Brillouin zone. We refer to the K or K' point also as the K$_{\eta}$ point with the valley index $\eta = \pm 1$. We take a silicene sheet on the $xy$-plane, and apply the electric field $E_z$ perpendicular to the plane. Due to the buckled structure the two sublattice planes are separated by a distance, which we denote by $2\ell$ with $\ell = 0.23\text{Å}$. It generates a staggered sublattice potential $\propto 2\ell E_z$ between silicon atoms at A sites and B sites.

The silicene system is described by the four-band second-nearest-neighbor tight binding model,

\begin{equation}
H = -t \sum_{\langle i,j \rangle \alpha \beta} c_{i\alpha}^\dagger c_{j\beta} + i\frac{\lambda_{SO}}{3\sqrt{3}} \sum_{\langle i,j \rangle \alpha \beta} \nu_{ij} c_{i\alpha}^\dagger \sigma_{\alpha \beta}^{z} c_{j\beta} \\
+ i\lambda_{R1}(E_z) \sum_{\langle i,j \rangle \alpha \beta} c_{i\alpha}^\dagger (\sigma \times \hat{d}_{ij})_{\alpha \beta}^{z} c_{j\beta} \\
- i\frac{2}{3}\lambda_{R2} \sum_{\langle i,j \rangle \alpha \beta} \mu_{\alpha} c_{i\alpha}^\dagger (\sigma \times \hat{d}_{ij})_{\alpha \beta}^{z} c_{j\beta} \\
+ \ell \sum_{i,\alpha} \mu_{i} E_z c_{i\alpha}^\dagger c_{i\alpha} + M \sum_{i,\alpha} c_{i\alpha}^\dagger \sigma_{z} c_{i\alpha},
\end{equation}

where $c_{i\alpha}^\dagger$ creates an electron with spin polarization $\alpha$ at site $i$, and $\langle i,j \rangle$ run over all the nearest/next-nearest neighbor hopping sites. We explain each term. (i) The first term represents the usual nearest-neighbor hopping with the transfer energy $t = 1.66\text{eV}$. (ii) The second term represents the effective SO coupling with $\lambda_{SO} = 3.9\text{meV}$, where $\sigma = (\sigma_{x}, \sigma_{y}, \sigma_{z})$ is the Pauli matrix of spin, with $\nu_{ij} = +1$ if the next-nearest-neighboring hopping is anticlockwise and $\nu_{ij} = -1$ if it is clockwise with respect to the positive $z$ axis. (iii) The third term represents the first Rashba SO coupling associated with the nearest neighbor hopping, which is induced by external electric field$\left[\frac{14}{15}\right]$. It satisfies $\lambda_{R1}(0) = 0$ and becomes of the order of $10\mu\text{eV}$ at the critical electric field $E_z = \lambda_{SO}/\ell = 17\text{meV} \cdot \text{Å}^{-1}$. (iv) The forth term represents the
second Rashba SO coupling with $\lambda_{R2} = 0.7\text{meV}$ associated with the next-nearest neighbor hopping term, where $\mu_i = \pm 1$ for the A (B) site, and $d_{ij} = d_{ij}/|d_{ij}|$ with the vector $d_{ij}$ connecting two sites $i$ and $j$ in the same sublattice. (v) The fifth term is the staggered sublattice potential term. (vi) The sixth term represents the exchange magnetization: Exchange field $M$ may arise due to proximity coupling to a ferromagnet such as depositing Fe atoms to the silicene surface or depositing silicene to a ferromagnetic insulating substrate, as has been argued for graphene$^{[13–15]}$. The Hamiltonian (1) can also be used to describe germanene, which is a honeycomb structure of germanium$^{[5, 6]}$, where various parameters are $t = 1.3\text{eV}$, $\lambda_{SO} = 43\text{meV}$, $\lambda_{R2} = 10.7\text{meV}$ and $\ell = 0.33\text{Å}$.

In this paper we derive the topological phase diagram in the $E_z-M$ plane and make its physical interpretation. The topological quantum numbers are the Chern number $C$ and the $\mathbb{Z}_2$ index. If the spin $s_z$ is a good quantum number, the $\mathbb{Z}_2$ index is identical to the spin-Chern number $C_s$. They are defined when the state is gapped and when the Fermi level is taken within the gap, and given by

$$
C = C_+ + C_- \quad \text{and} \quad C_s = \frac{1}{2}(C_+ - C_-),
$$

where $C_{\pm}$ is the summation of the Berry curvature in momentum space over all occupied states of electrons with $s_z = \pm 1$. They are well defined even if the spin is not a good quantum number$^{[15, 21]}$. In the present model the spin is not a good quantum number because of spin mixing due to the Rashba couplings $\lambda_{R1}$ and $\lambda_{R2}$, and the resulting angular momentum eigenstates are indexed by the spin chirality $s = \pm 1$. We can calculate these numbers at each point in the $E_z-M$ plane by using the standard formulas$^{[13–15]}$.

We present our result on the phase diagram in Fig.1 We show later how to derive the phase boundaries based on the low-energy Dirac theory. We have also calculated the band structure of a silicene nanoribbon with zigzag edges, which we give in Fig.2 for typical points in the phase diagram. The topological numbers are $(C, C_s) = (0, 0)$ in the BI phase, $(0, 1)$ in the QSH phase, $(2, 0)$ in the QAH phase with $M > 0$ and $(-2, 0)$ in the QAH phase with $M < 0$. In all these states the band gap is open, where the Fermi level is present, and they are insulators.

![Phase diagram](image1.png)

**FIG. 1:** (Color online) Phase diagram in the $E_z-M$ plane. Heavy lines represent phase boundaries, where the system becomes metallic. Chern and spin-Chern numbers ($C, C_s$) are well defined and given in insulator phases. Dotted lines represent the points where the band gap closes, which are within the VPM phase. A circle shows a point where the energy spectrum is calculated and shown in Fig.2.

![Band structure](image2.png)

**FIG. 2:** (Color online) The band structure of a silicene nanoribbon at marked points in the phase diagram (Fig.1). The vertical axis is the energy in eV, and the horizontal axis is the momentum. We can clearly see the Dirac cones representing the energy spectrum of the bulk, as explained in Fig.3. Lines connecting the two Dirac cones are edge modes. The spin $s_z$ is practically a good quantum number, which we have assigned to the Dirac cones. An enlarged figure of the QAH state is given in Fig.4.

We first discuss the system at $E_z = 0$ and compare our results with those previously obtained in graphene$^{[13–15]}$. The main difference is the appearance of almost flat edge modes in our system (Fig.3). This occurs because the Rashba interactions are different between these two systems. We have $\lambda_{R1} = 0$ for $E_z = 0$ and $\lambda_{R2} = 0$ at the K and K’ points in silicene, but $\lambda_{R1} \neq 0$ and $\lambda_{R2} = 0$ in graphene. Nevertheless, the difference is only quantitative. As far as the topological properties are concerned, there exists no difference. Indeed, in these two systems, the Chern number is identical in each corresponding phase together with quantized Hall conductivity, and the edge states support the edge current. However, the group velocity of the edge modes is extremely small due to the almost flat gapless modes in silicene.

Our most important result is the VPM phase, which appears in such regions that $E_zM \neq 0$ and occupies a major part of the phase diagram. A part of the conduction (valence) band is below (above) the Fermi level at the K (K’) point for $E_z > 0$, as is observed in Fig.2 (VPM). Hence, electrons are moved from the K valley to the K’ valley, as implies the valley polarization. The phase is characterized by the property that it is a metallic state though gaps are open both at the K and K’.
points. We note that the Chern and spin-Chern numbers are ill-defined in the VPM phase, since the Fermi level does not lie inside the band gaps at the K and K' points simultaneously. There exist M-VPM states on phase boundaries indicated by heavy lines in the phase diagram, where the conduction and valence bands touch the Fermi surface at the K and K' points, respectively, for $E_f > 0$. On the other hand, in SVPM states the conduction and valence bands touch the Fermi surface both at the K and K' points. We expect topological quantum critical phenomena in these states.

In order to explore the physics underlying the phase diagram, we analyze the low-energy effective Hamiltonian derived from the tight binding model. It is described by the Dirac theory around the $K, K'$ point as

$$H_0 = \hbar v_F (\eta k_x \tau_x + \eta k_y \tau_y) + \eta \tau_z h_{11} + \ell E_z \tau_z + M \sigma_z + \lambda_{R1} (\eta \tau_x \sigma_y - \tau_y \sigma_x)/2$$

(2)

with $h_{11} = \lambda_{SO} \sigma_z + a \lambda_{R2} (k_y \sigma_x - k_x \sigma_y)$, where $\tau_a$ is the Pauli matrix of the sublattice pseudospin, $v_F = \frac{\sqrt{3}}{a}$ is the Fermi velocity, and $a = 3.86\text{Å}$ is the lattice constant.

The Hamiltonian $H_+$ explicitly reads

$$
\begin{pmatrix}
E(1,1) & \hbar v_F k_+ & ia \lambda_{R2} k_- & 0 \\
\hbar v_F k_+ & E(1,-1) & -i \lambda_{R1} & -ia \lambda_{R2} k_- \\
-ia \lambda_{R2} k_+ & i \lambda_{R1} & E(-1,1) & \hbar v_F k_- \\
0 & ia \lambda_{R2} k_+ & \hbar v_F k_+ & E(-1,-1)
\end{pmatrix}
$$

(3)

in the basis $\{ \psi_{A+}, \psi_{B+}, \psi_{A\downarrow}, \psi_{B\downarrow} \}$, where $k_\pm = k_x \pm ik_y$, and the diagonal elements are

$$E(s_z, t_z) = \lambda_{SO} s_z t_z + \ell E_z t_z + M s_z,$$

(4)

with the spin $s_z = \pm 1$ and the sublattice pseudospin $t_z = \pm 1$. They are not good quantum numbers in general. However, since $\lambda_{R1}$ and $\lambda_{R2}$ are very small with respect to the other parameters, it is a good approximation to set $\lambda_{R1} = \lambda_{R2} = 0$ in most cases. Thus the spin $s_z$ is almost a good quantum number in general. An exceptional case occurs when two Dirac cones collapse and cross each other, forming a QAH state after taking into account the effect of $\lambda_{R1} \neq 0$, as we soon discuss.

We diagonalize the Hamiltonian (3) and obtain four energy levels. When two energy levels coincide, the band gap becomes zero, as found in Fig.2(M, VMP3, VMP2, SVPM). This occurs at the K and K' points, where $k_\pm = 0$. Let us temporarily neglect $\lambda_{R1}$ because it is very small. Then, the band closes when $E(s_z, t_z) = E(s'_z, t'_z)$ with (4). They yield four lines described by $E_z = \pm \lambda_{SO}/\ell$ for $|M| \leq \lambda_{SO}$, $M = \pm \lambda_{SO}$ for $|E_z| > \lambda_{SO}/\ell$, and two lines by $M = \pm (\lambda_{SO}/\lambda_{SO}) E_z$ outside the square. They are illustrated by dotted lines in Fig.1. These lines are modified by the nonzero effect of $\lambda_{R1}$, but the modification is too small to be recognized in Fig.1. See also Fig.3 for the typical order of correction.

The Hamiltonian can be diagonalized analytically in some cases. First, along the $E_z$-axis in the phase diagram [Fig.1], we have already demonstrated [7] that a topological phase transition occurs along the $E_z$-axis from the QSH insulator [Fig.2(QSH)] to the band insulator [Fig.2(BI)]. The critical point is given by

$$E_c = \pm \frac{2 \lambda_{SO}}{\ell} \left[ \sqrt{1 + (\alpha/\ell)^2} - 1 \right],$$

(5)

where we have set $\lambda_{R1}(E_z) = \alpha E_z$ with $\alpha = 10^{-3}\text{Å}$. Note that the effect of $\lambda_{R1}$ is negligible, $(\alpha/\ell)^2 = 10^{-4}$. The
SVPM realizes at the critical point, where helical currents flow in the bulk.

Second, along the $M$-axis, the first Rashba interaction vanishes ($\lambda_{R1} = 0$), and the energy spectrum reads

$$\mathcal{E} = \pm \sqrt{a^2 \lambda_{R2}^2 k^2 + \left(M - s_s \sqrt{\lambda_{SO}^2 + h^2 v_F^2 k^2} \right)^2}.$$  \hspace{1cm} (6)

We study a topological phase transition along the $M$-axis based on this formula (6). When $M = 0$, there are two spin-degenerate Dirac cones for conduction and valence bands with a gap between them [Fig.2(QSH1)]. As $M$ increases, the spin-up (spin-down) Dirac cones are pushed upward (downward) [Fig.2(QSH2)]. When $|M| \leq \lambda_{SO} (1 + a^2 \lambda_{R2}^2 / h^2 v_F^2)$, the band gap is given as $\Delta = |M - s_s \lambda_{SO}|$ at $k = 0$, and it closes at $M = s_s \lambda_{SO}$. This is a topological phase transition point [Fig.2(M)]. Let us temporally assume $\lambda_{R2} = 0$. Then, as $|M|$ increases further, the two Dirac cones cross each other making a circle around each $K$ point. Actually, the Rashba interaction ($\lambda_{R2} \neq 0$) mixes up and down spins, turning the crossing points into the anticrossing points, and opens a gap to form the QAH insulating state [Fig.2(QAH) and Fig.3].

When $|M| > \lambda_{SO} (1 + a^2 \lambda_{R2}^2 / h^2 v_F^2)$, the gap is given by

$$\Delta = a \lambda_{R2} \sqrt{M^2 / h^2 v_F^2 + a^2 \lambda_{R2}^2 / h^2 v_F^2 - \lambda_{SO}^2 / h^2 v_F^2}.$$ \hspace{1cm} (7)

We finally determine the phase boundary. It is determined as a boundary between insulating and metallic states. The Chern and spin-Chern numbers are quantized in insulating states, while they are ill-defined in metallic states. As we have seen, each Dirac cone moves upward or downward oppositely at the $K$ and $K'$ points. Because of this phenomenon the system can become metallic though the gap is open both at the $K$ and $K'$ points. This is the VPM state. It occurs when one valence band crosses the Fermi level. The condition yields four heavy lines ($M = \pm \lambda_{SO} \pm \ell E_2$) in the phase diagram (Fig.1). On the other hand, the gap formula (7) determines the boundary between the QAH phase and the VPM phase, which are the parabolic curves in the phase diagram (Fig.1). In passing we comment that the VPM phase is metallic in nature and does not have mobility gap. Thus the transition from insulator to VPM might accompany a mobility gap closing.

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