

Functionalized bismuth films: Giant gap quantum spin Hall and valley-polarized quantum anomalous Hall states

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The search for new large band gap quantum spin Hall (QSH) and quantum anomalous Hall (QAH) insulators is critical for their realistic applications at room temperature. Here we predict, based on first-principles calculations, that the band gap of QSH and QAH states can be as large as 1.01 and 0.35 eV in an H-decorated Bi(111) film. The origin of this giant band gap lies in both the large spin-orbit interaction of Bi and the H-mediated exceptional electronic and structural properties. Moreover, we find that the QAH state also possesses the properties of a quantum valley Hall state, thus intrinsically realizing the so-called valley-polarized QAH effect. We further investigate the possibility of large gap QSH and QAH states in an H-decorated Bi($\bar{1}10$) film and X-decorated ($X = \text{F, Cl, Br, and I}$) Bi(111) films.

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Since their discovery [1,2], there has been growing interest in topological insulators (TIs) that host conducting surface states inside the bulk insulating gap. The gapless surface states are topologically protected by time-reversal symmetry (TRS) and are robust to nonmagnetic perturbations [3–5]. The first theoretically predicted [6] and experimentally observed [7] TI is a HgTe/CdTe quantum well structure that is a two-dimensional (2D) TI, also known as a quantum spin Hall (QSH) insulator. In a QSH insulator, pairs of dissipationless edge channels with opposite spins exist, leading to extraordinary properties and possible applications in low dissipation electronic devices. On the other hand, the realization of the quantum anomalous Hall (QAH) effect, which was first suggested to occur in a honeycomb lattice model [8], has been achieved recently in Cr-doped topological insulators (Bi,Sb)₂Te₃ [9] via suppressing one of the spin channels [10,11], but requires extremely low temperatures (30 mK). To obtain room temperature QSH- and QAH-based electronic devices, the search for novel materials with large band gaps as well as stable atomic and magnetic structures has been a fairly important topic in the field. In spite of extensive efforts so far [12–24], most of the known systems with the desired topological properties have a small band gap, which greatly obstructs their potential room temperature applications.

Valley polarization, as a new degree of freedom in honeycomb lattices, in addition to the intrinsic charge and spin, has received considerable attention in recent years [25,26]. The valley Hall conductivity can be nonzero when the inversion symmetry is broken, realizing the quantum valley Hall (QVH) effect characterized by the so-called valley Chern number [25]. Quite recently, a new quantum state, a valley-polarized QAH state that exhibits the electronic properties of both the QVH state and QAH state, has been predicted in silicene through tuning the extrinsic spin-orbit coupling (SOC) with broken TRS [27]. It provides a new way to design the dissipationless valleytronics. However, the presence of both inversion symmetry and TRS as well as the small SOC in pristine silicene makes experimental studies and possible applications difficult.

Generally, materials with strong SOC and simultaneously broken inversion and time-reversal symmetries, which exhibit nontrivial topological phases, are in high demand. As the heaviest atom with an effectively stable isotope and strong SOC [28], bismuth is an important ingredient for both two-dimensional (2D) and three-dimensional (3D) TIs, such as Bi(111) bilayers [13,16], Bi_{1-x}Sb_x [29,30], Bi chalcogenides [31,32], and TlBiSe₂ [33–35]. The Bi(111) bilayer has drawn much attention due to a relatively large band gap of the 2D system (~ 0.2 eV [13]) with the edge states observed experimentally [36]. Rather recently, Bi(111) bilayers were grown on different substrates [37–39]. On weakly interacting substrates, ultrathin, (111)-oriented films are unstable with respect to the transformation into another allotrope of Bi [40]. It grows in the black-phosphorous (A17) structure that resembles (110) layers of the bulk Bi (A7) structure [41] and turns out to be topologically trivial [42]. Recently, atomic hydrogen chemisorbed on 2D TIs, for example, graphene [43–45] and stanene [17], has been proven to be an effective way to modulate their properties. However, the band gap of graphene is quite small, and stanene is not a 2D TI in its equilibrium structure.

In this Rapid Communication, based on first-principles calculations, we show that both the electronic and topological properties of ultrathin Bi films can be drastically modified when decorated by H. An H-decorated Bi(111) film [H-Bi(111)] exhibits a topological energy gap of 1.01 eV, which is much larger than those in known TIs. Besides the strong SOC [46], the lattice parameters also play a critical role in determining the giant band gap of the 2D system. For the case of Bi($\bar{1}10$) film [H-Bi($\bar{1}10$)], H decorating induces the realization of 2D TI phases with a direct gap of 0.34 eV. Based on Chern number calculations, we further demonstrate that the valley-polarized QAH effect can be realized in Bi thin films that are hydrogenated only on one side of the films.

Bulk bismuth shows two different bond lengths: 3.07 Å within (111) bilayers and 3.53 Å between them. Projected onto the (111) plane, the Bi bilayer forms a hexagonal lattice. In the Bi($\bar{1}10$) layers, two different bonds exist within a slightly buckled pseudosquare lattice. The crystal structures of H-Bi(111) and H-Bi($\bar{1}10$) are plotted in Fig. 1. In contrast to

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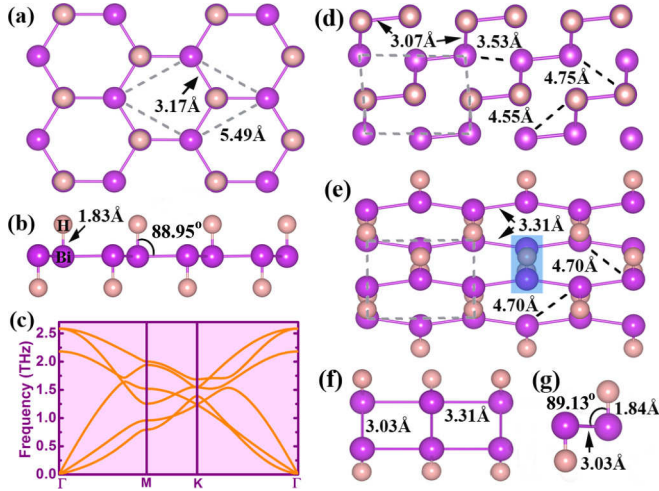


FIG. 1. (Color online) (a) Top and (b) side view of optimized structures for H-Bi(111). The corresponding phonon band structure is shown in (c). The top view of the unrelaxed and relaxed crystal structure of H-Bi(110) is shown in (d) and (e), respectively. (f) Side view of the relaxed crystal structure of H-Bi(110). (g) Zoomed-in view of the highlighted areas in (e). The unit cells are indicated by gray dashed lines. The numbers are interatomic distances ($d_{\text{Bi-H}}$ or $d_{\text{Bi-Bi}}$) and angles ($\theta_{\text{H-Bi-Bi}}$) between Bi (H) atoms.

decorated Sn [17], H-Bi(111) has a quasipolar geometry with the angle $\theta_{\text{H-Bi-Bi}}$ being slightly smaller than 90° . Therefore, the hexagonal lattice parameter increases from 4.54 to 5.49 Å, while $d_{\text{Bi-Bi}}$ is only expanded to 3.17 Å. H-Bi(110) forms a structure with an AB stacking of the pseudosquare layers, with $d_{\text{Bi-Bi}}$ of 3.03 Å connecting the atoms within these layers and 3.31 Å between the layers. In addition, it is interesting to note that Bi and H atoms within the same layer have a configuration that is quite similar to that of H-Bi(111). The H decoration changes the structure significantly, and the inversion symmetry is obtained as shown in Figs. 1(d) and 1(e). We further confirm the stability of H-Bi(111) by phonon calculations. The real phonon frequencies at all momenta shown in Fig. 1(c) confirm that the structures are stable.

The calculated band structures for H-Bi(111) and H-Bi(110) are plotted in Fig. 2. In the case without SOC, as shown in Figs. 2(a) and 2(c), the systems are gapless and show a semimetallic character with one band crossing exactly at the K point for the H-Bi(111) case and slightly away from the Y point for the case of H-Bi(110). This is different from other Bi-based TIs, such as the Bi(111) bilayer [13] or the Bi_2Se_3 [31], but is quite similar to graphene [1]. Taking SOC into account, a band gap opens [Figs. 2(b) and 2(d)]. Different from both the Bi-based TIs and graphene, the Dirac-related bands have contributions mainly from the p_x and p_y orbitals while the p_z orbital is removed away from the Fermi level by H, resulting in the large band gap. A similar mechanism was recently reported for a Bi/Si system [47]. To identify the band topology, the $\mathbb{Z}_2$ invariant is investigated by evaluating the wave function parities at four time-reversal invariant momentum (TRIM) points [29], i.e., the Γ and three M points for H-Bi(111) and Γ , X, Y, and M for H-Bi(110) [insets of Figs. 2(a) and 2(c)]. The product of the parities of the occupied bands at TRIM k , δ_k , is given in the insets of Figs. 2(b) and 2(d), together with the

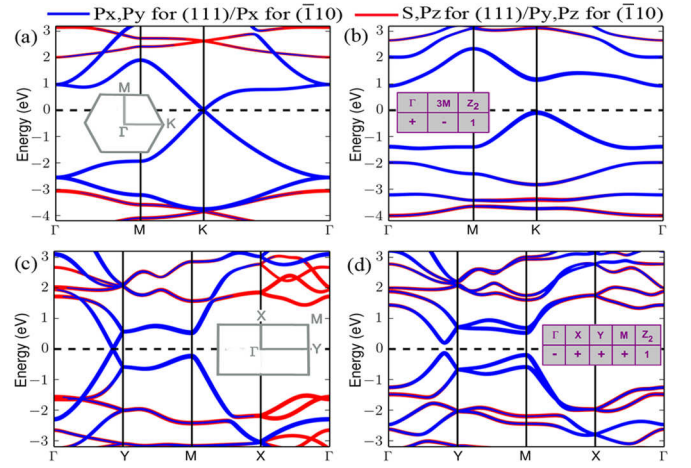


FIG. 2. (Color online) Orbital-resolved band structures for (a), (b) H-Bi(111) and (c), (d) H-Bi(110), without [(a), (c)] and with [(b), (d)] SOC, weighted with the s , p_x , p_y , and p_z characters. Dark (blue) colors mark states that contribute to the fundamental band gap. The Fermi level is indicated by the dashed line. The insets in (a) and (c) show the 2D Brillouin zone, and those in (b) and (d) show the products of the parities of all occupied bands at the time-reversal invariant momenta and the $\mathbb{Z}_2$ number.

$\mathbb{Z}_2$ number (ν) determined by $(-1)^\nu = \prod_k \delta_k$. We verify that both the H-Bi(111) and H-Bi(110) are QSH insulators [not for H-Bi(110) and H-Bi(A17)]. The QSH state is further explicitly confirmed by the emergence of the gapless edge states in thin nanoribbons of the bilayers [48].

To utilize the QSH effect in room temperature QSH-based electronic devices, a large band gap is needed. As shown in Figs. 2(b) and 2(d), an indirect band gap of 1.01 eV and a direct band gap of 0.34 eV are obtained for H-Bi(111) and H-Bi(110), respectively, which are large enough for practical applications at room temperature. For H-Bi(111), a similar energy gap was quite recently reported by Song *et al.* [46]. In order to test the stability and understand the mechanisms of such a giant band gap, we expose the electronic structure of the bilayers to variations of different parameters. Figure 3(a) shows the variation in the energy gap at the Γ point (E_Γ), K point (E_K), and the global energy gap (E_{global}) of H-Bi(111) as the SOC strength λ is varied. Such a variation can be realized

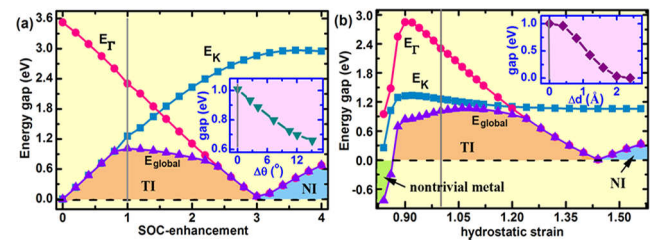


FIG. 3. (Color online) The calculated energy gaps at the Γ point (E_Γ), K point (E_K), and the global energy gap (E_{global}) of H-Bi(111) as a function of (a) SOC strength and (b) hydrostatic strain. A phase transition from TI to normal insulator (NI) occurs, accompanied by a gap closing and reopening at Γ . The insets in (a) and (b) show E_{global} vs $\theta_{\text{H-Bi-Bi}}$ and $d_{\text{Bi-H}}$, respectively.

experimentally by alloying Bi with isoelectronic Sb [30]. Starting from a calculation without SOC, it can be seen that E_Γ closes with increasing λ , while E_K opens accordingly. The transition from a direct ($E_K = E_{\text{global}}$) to an indirect band gap occurs when the relative SOC strength (λ/λ_0 , where λ_0 is the actual SOC strength) exceeds 0.8. At λ_0 , the global energy gap reaches its maximum and stays rather constant, since both the highest occupied band at the K point and the lowest unoccupied band at the Γ point are downshifted with increasing λ . With further enhancing SOC, the band gap becomes direct at the Γ point, and then decreases rapidly. A band gap closing and reopening occurs at $\lambda/\lambda_0 = 3.0$, thus marking a band inversion and phase transition from a TI to a normal insulator (NI) that is confirmed by our topological analysis.

The lattice constant a , the angle $\theta_{\text{H-Bi-Bi}}$ (or bond length $d_{\text{Bi-Bi}}$), and the Bi-H separation $d_{\text{Bi-H}}$ are three key structural parameters that define the lattice of H-Bi(111). They might be altered, e.g., by epitaxial constraints imposed by a substrate. To reveal their influence, we show the variation in the band gap as a function of strain a/a_0 , where a_0 is the equilibrium lattice constant, $\Delta\theta$ (change of $\theta_{\text{H-Bi-Bi}}$ from the equilibrium value), and Δd (change of $d_{\text{Bi-H}}$) in Fig. 3(b), the insets of Fig. 3(a), and the insets of Fig. 3(b), respectively. The large (>0.6 eV) indirect band gap is robust for a/a_0 ranging from 0.86 to 1.24, showing high adaptability in various application environments. When further compressed or expanded, the global band gap decreases rapidly. Under compression we observe the transition to a nontrivial metal, while with expansion a phase transition from TI to NI occurs when the gap closes and reopens at the Γ point at $a/a_0 = 1.44$, similarly to the case of strong SOC enhancement. By applying hydrostatic strain, both $\Delta\theta$ and Δd are determined from a full optimization of the internal atomic coordinates. As shown in the insets of Fig. 3, the global band gap decreases with an increase of both $\Delta\theta$ and Δd .

Having established the existence of a stable QSH phase, we focus now on the possibility of realizing the QAH effect in functionalized Bi films. The essential ingredient for the transition from the QSH to the QAH phase is the breaking of TRS, since the QSH phase can be considered as two copies of the QAH states that are coupled together by TRS [11]. To date, most of the research on magnetically doped TIs has been focused on transition-metal doping [9–11,49]. In the following, we show theoretically that the realization of ferromagnetic ordering and QAH states without transition-metal doping can be achieved when the hydrogen atoms are removed from one side of H-Bi(111) while keeping the other side hydrogenated (what we call half-hydrogenated Bi or semihydrogenated Bi). Experimentally, semihydrogenated layers might be accessible when one side of the Bi film is protected by a substrate and the other one is exposed to hydrogen. For graphene, half-hydrogenated layers have been successfully fabricated [50].

To understand the origin of the spin polarization which we encounter, the calculated total density of states (DOS) and partial DOS for semihydrogenated Bi(111) without SOC are shown in Fig. 4(a). We clearly see that the spin polarization is mainly carried by the p_z states of the unhydrogenated Bi atoms due to the breaking of the orbital network (see also the spin density plots in the Supplemental Material [48]).

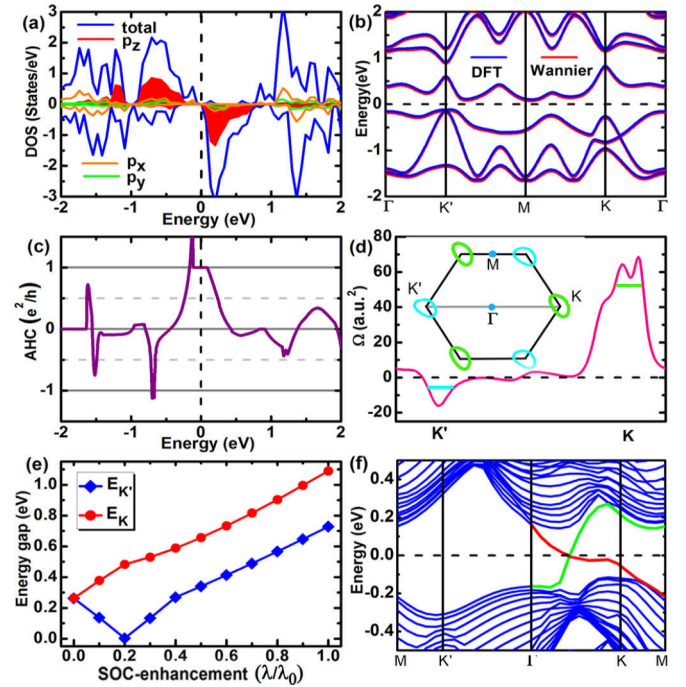


FIG. 4. (Color online) (a) Total density of states (DOS) and partial DOS of the unhydrogenated Bi atoms (decomposed into $6p_x$, $6p_y$, and $6p_z$ states) of half H-decorated Bi(111) without SOC. Positive and negative values indicate spin-up and spin-down channels, respectively. (b) Wannier and first-principles band structures with SOC for half H-decorated Bi(111). The Wannier results have been shifted down by 30 meV for visibility. (c) Anomalous Hall conductivity as a function of the position of the Fermi level E_F . (d) Berry curvature distribution of the occupied bands in the K - Γ - K' direction. The inset shows the contour of the Berry curvature distribution (as marked in main panel) around valleys K and K' . (e) The energy gaps at valleys K (E_K) and K' ($E_{K'}$) as a function of SOC strength. (f) Band structure of zigzag-terminated half H-decorated Bi(111) exhibiting valley-polarized QAH states. The states located at different edges are indicated by different colors.

The Fermi level is pinned inside the gap between the filled spin-up and empty spin-down Bi $6p_z$ states, leading to a magnetic moment of $1.0\mu_B$ per unit cell. Furthermore, non-spin-polarized density functional theory (DFT) calculations for semihydrogenated Bi(111) show that the spin-polarized state is more stable by about 93.3 meV, proving that the ground state is magnetic. The calculated band structure in the presence of SOC is shown in Fig. 4(b), from which we conclude that our system is a magnetic insulator with a sizable indirect energy gap of 0.35 eV. Calculation of the magnetic anisotropy energy shows that the easy magnetization axis points out of plane and it is 1.19 meV lower in energy than the in-plane spin orientation. The existence of an insulating, magnetic ground state does not necessarily result in a strong magnetic coupling and, furthermore, in a QAH phase. To access the nature of the exchange coupling between the spin moments of unhydrogenated Bi atoms, which are separated by 5.49 Å, we compute the total energies of ferromagnetic (E_{FM}) and antiferromagnetic (E_{AFM}) states, finding that the ferromagnetic order is favored by $\Delta E = E_{\text{AFM}} - E_{\text{FM}}$ of 21.79 meV [48].

To identify the topological properties and predict a stable QAH state resulting from the sizable energy gap, we calculate the anomalous Hall conductivity $\sigma_{xy} = (e^2/h)C$, where C , quantized and known as the first Chern number in the case of an insulator, can be obtained as an integral of the Berry curvature of occupied state $\Omega(\mathbf{k})$ over the Brillouin zone [48]. The anomalous Hall conductivity as a function of band filling is calculated and presented versus the position of the Fermi level in Fig. 4(c). When the chemical potential is located within the energy gap, the Chern number of all occupied states indeed acquires an integer value of +1, confirming the QAH effect in semihydrogenated Bi(111). However, the semihydrogenation of Bi(111) leads to the simultaneous breaking of TRS and inversion symmetry, and the band structures at valleys K and K' have different patterns [Fig. 4(b)]. Valleys K and K' are distinguishable, and the valley-polarized QAH state, which exhibits properties of both the QAH state and the quantum valley Hall (QVH) state [25,27], is obtained. For further insight, the Berry curvature of all occupied bands along the K - Γ - K' path is plotted in Fig. 4(d). It is clearly visible that the Berry curvature distribution is localized in the vicinity of K and K' , and it has an opposite sign around the two valleys. The evolution of the energy gap at K and K' , shown in Fig. 4(e) as a function of the SOC strength, reveals the underlying physics of the formation of the QVH state. At K' , the energy gap closes and reopens as the SOC is increased, while that at the K point always opens, indicating that a topological phase transition occurs at the K' but not at K , resulting in different valley-resolved Chern numbers, i.e., $C_K = 1$ and $C_{K'} = 0$. To further confirm the valley-polarized QAH state, the edge states of zigzag-terminated half H-decorated Bi(111) at valleys K and K' are calculated and shown in Fig. 4(f). The number of edge states in each valley indeed corresponds to the corresponding valley Chern number.

For device applications, it is important to make sure that, given a large enough bulk band gap of the lattice-matching substrate, which is aligned with the Bi-originated 2D gap, the predicted topological properties are preserved [47]. While, owing to the enlarged lattice constant, both Bi(111) [36] and Bi chalcogenides [37–39] are not suitable for this purpose, we demonstrate this by taking MoS₂ ($\sqrt{3} \times \sqrt{3}$) as an example substrate, which fits hydrogenated Bi nicely both in the lattice

constant as well as in the alignment of the band gaps. We confirmed that magnetism for the semihydrogenated Bi is maintained and the topological properties for both fully and semihydrogenated cases are unchanged [48].

Nontrivial topological QAH states occur also for not fully hydrogenated Bi($\bar{1}10$) under strain [48]. Removing half of the hydrogen atoms in one of the two layers results in a magnetic ground state, but this state shows a metallic character with a global energy gap of -0.46 eV at the equilibrium lattice constant. Under hydrostatic strain, a transition from metal to insulator occurs, leading to the realization of the QAH state. Similar to the H-Bi(111), the QSH states form in the F-, Cl-, Br-, and I-decorated Bi(111), with gigantic energy gaps of 1.10, 0.93, 0.88, and 0.87 eV, respectively [48]. But for the half-decorated cases, the films show a metallic character even with the effect of strain. This is different from the half I-decorated Sn, which is reported to realize the QAH effect with a large energy gap [18].

In summary, by performing DFT calculations for fully and semihydrogenated Bi(111) and Bi($\bar{1}10$), we have demonstrated that both QSH and QAH states with giant band gaps can be realized. Especially in H-decorated Bi(111), the band gaps for the QSH and QAH states reach 1.01 and 0.35 eV, respectively. We further predict that the QAH state in semihydrogenated Bi(111) is quite different from the normal one. It exhibits the properties of both the QAH state and QVH state, realizing a different quantum state, called a valley-polarized QAH insulator. Both the QSH and valley-polarized QAH states survive even if the decorated Bi bilayers are on an appropriate substrate. Our results are of importance for further theoretical and experimental studies of topological insulators both from the point of view of fundamental exploration and as well as practical applications at room temperature.

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