Two-dimensional (2D) materials consisting of heavy atoms with particular arrangements may host exotic quantum properties. Here, we report a unique 2D semiconducting binary compound, a Sn$_2$Bi atomic layer on Si(111), in which hexagons are formed by bonding Bi with a triangular network of Sn. Because of the unique honeycomb configuration, the heavy elements, and the energy-dependent hybridization between Sn and Bi, 2D Sn$_2$Bi not only shows strong spin-orbit coupling effects but also exhibits high electron-hole asymmetry: Nearly free hole bands and dispersionless flat electron bands coexist in the same system. By tuning the Fermi level, it is possible to preserve both nearly free and strongly localized charge carriers in the same 2D material, which provides an ideal platform for the studies of strongly correlated phenomena and possible applications in nanodevices.

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The exploration of novel two-dimensional (2D) materials has attracted enormous research effort since the discovery of graphene [1,2], due to their promising properties and potential applications. Recently, the material family has been significantly enriched by adding a series of new members hosting diverse 2D quantum phases, including topology [3–5], ferromagnetism [6,7], ferroelectricity [8], charge density wave [9], superconductivity [10], etc. Current research predominantly focuses on 2D layered (like transition metal dichalcogenides [11]) or elemental (like graphene [12], silicene [13–15], germanene [16,17], stanene [18,19], borophene [20,21], blue phosphorous [22,23], antimonene [24], and bismuthene [25]) materials or others (like 2D alkali-earth metal hydroxides [26], graphene oxides, and silicene oxides [27–28]). In contrast, nonlayered binary compounds in the 2D limit could possess unusual chemical stoichiometry and atomic configuration and display extraordinary electronic properties, which greatly expands the opportunities for future research. This broad class of 2D materials, however, remains largely unexplored.

The development of new 2D nonlayered materials with high quality requires fine control of the growth dynamics at the atomic scale and, thus, represents a grand challenge to the research community. In practice, the fabrication and application of 2D materials both need a supporting substrate. Semiconducting substrates are usually preferable, as they enable the measurement of optical and transport properties within the substrate band gap. The exploration of novel 2D atomic layers on a silicon substrate is of particular interest, since it could be immediately applicable in the silicon industry. One intriguing direction is novel 2D binary materials composed of heavy elements, with strong spin-orbit coupling (SOC) effects and crystallized in a honeycomb lattice—a fascinating model lattice displaying exotic quantum effects, such as spin-valley coupling [29], Ising ferromagnetism [7], and quantum spin (anomalous) Hall effects [30,31]. In light of the recent successful growth of stanene and bismuthene and discoveries of 2D topological and superconducting phases therein [18,25,32,33], the experimental study of 2D Sn-Bi binary compounds is of crucial importance, which might offer rich material features beyond their elemental counterparts.

In this Letter, we report on the discovery of a novel 2D honeycomb structure of Sn$_2$Bi on a Si(111) surface. Combining scanning tunneling microscopy (STM) and
density functional theory (DFT) calculations, a honeycomb configuration is identified, in which hexagon Bi atoms are connected with each other by bonding with a triangular network of Sn atoms beneath. The scanning tunneling spectroscopy (STS) and angle-resolved photoemission spectroscopy (ARPES) measurements reveal a nearly 0.8 eV band gap. Moreover, hole gas bands in occupied states and flat band in unoccupied states are found to coexist in the same system. DFT calculations reveal that the unique honeycomb structure, where hexagon Bi atoms couple indirectly through Sn—Bi bonds, in combination with the energy-dependent hybridization between Sn and Bi results in this exotic electronic structure.

A silicon (111) wafer with a resistivity of 0.01–0.02 Ω cm was flashed to about 1250°C to obtain a clean (7 × 7) reconstruction in a UHV chamber. The β phase of Si(111)-√3 × √3-Bi (β-Bi/Si(111) for short) was prepared by the deposition of about 1.2 monolayer (ML) Bi on a Si(111)-√3 × √3 surface at room temperature and subsequent annealing at 670 K [34]. To prepare honeycomb Sn2Bi, Sn atoms were deposited on the β-Bi/Si(111) surface at 470 K. Then the sample was in situ transferred to the STM chamber and measured under a liquid-helium temperature. The STS were measured by a lock-in technique, in which an ac voltage of 10 mV and 676 Hz was superimposed on the tip bias. All the STM images were processed by WSxM software [35].

ARPES measurements were performed by the He Iα (hν = 21.218 eV) resonance lines and VG SCIENTA R4000 analyzer. The angular and energy resolutions were set to 0.2° and 30 meV, respectively. In the time-resolved ARPES measurements, we used 1.48-eV pump and 5.92-eV probe pulses with the repetition rate of 250 kHz. Samples were transferred in situ by a high-vacuum suitcase and measured at 300 K.

First-principles calculations were performed by the Vienna ab initio simulation package based on the density functional theory (DFT) [36], using the projector-augmented wave potential [37], Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional [38], and the plane wave basis with an energy cutoff of 250 eV. The model of the Si(111) substrate includes three Si bilayers, with the top bilayer relaxed during structural optimization and the bottom Si atoms saturated by hydrogen. A vacuum layer of ~10 Å was selected. Structural optimization was performed by applying a force convergence criterion of 0.01 eV/Å and a Gaussian smearing of 0.05 eV together with a 5 × 5 × 1 Monkhorst-Pack k grid for 2 × 2 superstructures on Si(111). Self-consistent electronic structure calculations were done by using an 11 × 11 × 1 k grid and including the SOC. STM images were simulated by the local density of states based on the Tersoff-Hamann approach [39]. The crystal structure analysis was by the particle swarm optimization (CALYPSO) algorithm [40,41].

![FIG. 1. A highly ordered honeycomb structure of a Sn-Bi overlayer on Si(111).](image)

(a) Large-scale STM image (V_{tip} = −2.0 V, I = 90 pA) of about 0.9 ML Sn on a β-Bi/Si(111) surface. An area of uncovered substrate is highlighted by the red dotted circle. (b) Atomic resolution STM image (V_{tip} = 1.0 V, I = 90 pA) taken on the Sn-Bi overlayer structure showing a uniform honeycomb lattice. (c) Height profile along the red arrow in (a), which indicates the height difference between the Sn-Bi overlayer and β-Bi/Si(111). (d) Line profile along the red arrow in (c). The periodicity of 0.77 ± 0.01 nm illustrates the formation of a 2 × 2 superstructure on Si(111). (e) Amplified STM image (V_{tip} = 1.0 V, I = 99 pA) of an area containing a boundary separating the Sn-Bi overlayer structure (bottom left) and β-Bi/Si(111) (upper right). (f) FFT patterns of a high-resolution STM image containing both a honeycomb Sn-Bi overlayer and uncovered β-Bi/Si(111). Red and blue dashed circles mark two sets of patterns that correspond to β-Bi/Si(111) and the Sn-Bi overlayer, respectively.
STM image containing coexisting areas of honeycomb structure and \(\beta\)-Bi/Si(111) shown in Fig. 1(e) indicates that the lattice of honeycomb structures is 30° rotated with respect to the lattice of \(\beta\) – Bi/Si(111). Combined with the measured lattice constant of 0.77 nm, it can be concluded that the honeycomb structure is a \(2 \times 2\) reconstruction with respect to the Si(111) substrate. The fast Fourier transform (FFT) of a high-resolution image including these two atomic structures simultaneously. The FFT of the \(\text{dI/dV}\) maps taken at the Si(111) reconstruction with \(\beta\)–Bi/Si(111) surface shown in Fig. 1(c) indicates a height difference of \(h\) \(\text{nm}\). Besides, the line profile across the boundary separating the honeycomb structure and \(\beta\)-Bi/Si(111), respectively. The data are well fitted by a red parabolic line, indicating the free-hole-like surface states.

Differential conductance \((\text{dI/dV})\) measurements taken on the honeycomb structure shown in Fig. 2(a) reveal a semiconducting gap of about 0.8 eV (from 0 to 0.8 eV above the Fermi level). The \(\text{dI/dV}\) maps taken at the energy range \((-0.6, 0)\) eV show clear standing-wave patterns, and an example is shown in Fig. 2(b), which contains the fluctuation of density of states (DOS) and atomic structures simultaneously. The FFT of the \(\text{dI/dV}\) map shows an isotropic circle at the center. Considering that the standing waves are due to quasiparticle interference (QPI) in the valence band, we measure the values of scattering vector \(q\) along the \(\Gamma-K\) direction at different energies and draw the \(E-q\) dependence in Fig. 2(d). The data can be fitted well by a parabolic curve, indicating the existence of nearly free 2D hole gas in the surface of the honeycomb structure. The fitted valence band maximum (VBM) is located at \(E_0 = -65\) meV, with effective mass \(m^*/m_0 = -0.14\) (\(m_0\) is the free electron rest mass). The slightly different VBM values measured by QPI fitting and \(\text{dI/dV}\) curves (at about 0 eV) are due to the band bending induced by the STM tip [42].

To illustrate the band structure of the honeycomb lattice more explicitly, ARPES measurements were performed on a sample with 95% of the surface covered by the Sn-Bi overlayer. Band dispersions along the \(M-\Gamma-K-M\) directions are shown in Fig. 3(a). Two surface bands \((S_1\) and \(S_2\)) with the vertex overlapping at the \(\Gamma\) point can be distinguished unambiguously. The VBM is located at \(-0.2\) eV (referenced to the Fermi level), which is \(-0.2\) eV lower than that in the \(\text{dI/dV}\) curve [Fig. 2(a)]. This discrepancy may arise from the photohole doping effect in ARPES and the tip-induced band bending in STS [42,43]. The constant energy contour at the VBM shows three spots in Fig. 3(b), corresponding to a \(2 \times 2\) superstructure. Besides, time-resolved ARPES with a pump-probe method was performed to determine the information of unoccupied states. Interestingly, only a nearly flat band along the \(M-K\)
Sn-rich and Bi-poor conditions (see details in Ref. [44]). The red (blue) balls represents contributions of Sn (Bi) atoms. After systematically searching for over 5000 potentials of Sn (Bi) bulk, we obtained a semiconducting band gap and a strong electron-hole asymmetry (i.e., nearly free holes and strongly localized electrons). In order to understand the atomic structure of the honeycomb Sn-Bi overlayer, we performed first-principles DFT calculations. The key issue is to find the right atomic model that can reproduce all the essential experimental results: (i) It is an atomically thin structure in a honeycomb model that can reproduce all the essential experimental data (background). (b),(c) STM images of honeycomb Sn$_2$Bi taken at ($V_{up}=1.0$ V, $I=90$ pA) and ($V_{up}=-1.0$ V, $I=91$ pA). (d), (e) Simulated STM images of honeycomb Sn$_2$Bi taken at $-1.0$ eV below and $+0.46$ eV above the calculated VBM, respectively.

FIG. 5. Comparison of experimental results and theoretical calculations of honeycomb Sn$_2$Bi. (a) Electron dispersion along the $K$-$\Gamma$-$K$ direction from QPI measurements (yellow dots), DFT calculations (red balls), and second derivatives of ARPES data (background). (b),(c) STM images of honeycomb Sn$_2$Bi taken at ($V_{up}=1.0$ V, $I=90$ pA) and ($V_{up}=-1.0$ V, $I=91$ pA). (d), (e) Simulated STM images of honeycomb Sn$_2$Bi taken at $-1.0$ eV below and $+0.46$ eV above the calculated VBM, respectively.

Figs. 4(b) and 4(c), which includes 1/2 ML Bi in the first layer and 1 ML Sn in the second layer above Si(111). The stacking geometry of Sn$_2$Bi/Si(111) is elaborated in Supplemental Material [44]. This structure is very stable, because all the Si, Sn, and Bi atoms satisfy the octet rule by forming four, four, and three chemical bonds, respectively. A honeycomb lattice is formed by the top Bi atoms that are located above the $T_4$ and $H_3$ sites of Si(111). The low-lying Sn atoms are either at the boundary or center of the Bi hexagon. Note that all the honeycomb Bi atoms are bridged by neighboring Sn atoms, and there is no direct bond between Bi atoms, resulting in an unusual honeycomb structure. The 2D Sn$_2$Bi is expected to be stable at elevated temperatures according to $ab$ initio molecular dynamic simulations [44].

We make comparisons between the theory and experiments. The height difference between the honeycomb Sn$_2$Bi and the $\beta$-Bi/Si(111) substrate is 0.18 nm from the theory, in good agreement with the STM value of $0.19\pm0.01$ nm. Moreover, the calculated band structure [Fig. 4(e)] shows two parabolic valence bands centered at the $\Gamma$ point ($S_1$ and $S_2$) and two nearly flat conduction bands minimum along the $M$-$K$ direction, which are separated by an indirect semiconducting gap, consistent with the ARPES measurement. According to the determined band edges, an indirect energy gap of about 0.87 eV can be derived from the ARPES data, which is consistent with the gap value measured by STS ($\sim0.8$ eV). The theoretical gap value of 0.36 eV, typically underestimated by DFT PBE, meanwhile is comparable to the experimental value of $\sim0.8$ eV. Also, the band dispersions of $S_1$ and $S_2$ fit well with the STS measurements and ARPES data [Fig. 5(a)]. In addition, the simulated STM images [Figs. 5(d) and 5(e)] show triangular (honeycomb) patterns for unoccupied (occupied) states as observed in experiments. At the unoccupied states, the brightness contrast of the two
nearest spots in Figs. 5(c) and 5(e) can be explained by the different registry of the Bi atoms above silicon [H₃ and T₄ sites on Si(111)]. In summary, a quantitative agreement between the theory and experiments is established.

According to a band analysis, the valence and conduction bands around the Fermi level are mainly contributed by heavy elements Bi and/or Sn [Fig. 4(e)], which implies strong SOC effects in this system. The effects are emphasized by comparing band structures excluding and including the SOC [Figs. 4(d) and 4(e)]. The SOC generates a giant Rashba splitting for the lowest conduction band near the Γ point as well as a large band splitting of ~0.5 eV at the Γ point between the S₁₂ and S₃₄ bands. This SOC-induced band splitting can be well described by DFT PBE, as it is insensitive to material details. Indeed, ARPES observed such a kind of band splitting in Fig. 5(a); thus, the strength of the SOC of this material is experimentally quantified.

In contrast to the elemental honeycomb lattice, the Sn₂Bi binary honeycomb lattice shows a strong electron-hole asymmetry. The unique feature is inherently related to the couplings between Bi and Sn atoms. In the honeycomb structure, every Bi atom bonds to three Sn atoms to form two inequivalent trimers around a Sn atom at hollow sites. Band dispersions thus depend critically on the strength of hybridization between Bi and Sn. The hybridization is strong for the top of valence bands, which shows the character of 2D light hole gas with the effective mass \( m^* ≈ 0.14m_0 \), but quite weak for the bottom of the conduction bands, leading to nearly flat bands [44]. Therefore, nearly free and strongly localized charge carriers can be both realized in the same material by tuning the Fermi level.

Electrons in a dispersionless flat band with quenched kinetic energy are highly degenerate, which provides an ideal platform to realize the strongly correlated electronic states. In 2D materials, flat bands are predicted theoretically in square (lieb) lattices [53,54], kagome lattices [55,56], and honeycomb (hexagonal) lattices [57,58] but observed only on some artificial systems [54]. By tuning the Fermi level of the novel 2D system, transport measurements in flat bands may bring surprising properties like ferromagnetism and superconductivity [59,60]. Furthermore, the nearly free-hole-like carriers in the \( S_1 \) and \( S_2 \) bands with spin splitting indicate that the 2D Sn₂Bi with a semiconducting gap will be a good candidate for high-performance nanoelectronic devices.

Like Sn₂Bi/Si(111), similar 2D materials \( X_2Y \) (\( X = \) group IVA, \( Y = \) group VA) could be grown on substrates that satisfy the conditions of surface lattices commensurate with that of the 2D overlayer, small lattice mismatch, and one-electron dangling bond per surface site. Candidate substrates include (111) surfaces of rhombohedral and cubic structures as well as (001) surfaces of hexagonal structures. For instance, we expect that the 2D Sn₂Bi could be grown on Ge(111) and ZnS(111). This offers opportunities to explore new 2D binary compounds and novel quantum phenomena.

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