Chemical migration and dipole formation at van der Waals interfaces between magnetic transition metal chalcogenides and topological insulators

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Metal and magnetic overlayers alter the surface of the topological insulator (TI) bismuth selenide (Bi$_2$Se$_3$) through proximity effects but also by changing the composition and chemical structure of the Bi$_2$Se$_3$ subsurface. The interface between Bi$_2$Se$_3$ and Mn metal or manganese selenide was explored using x-ray photoelectron spectroscopy revealing chemical and electronic changes at the interface. Depositing Mn metal on Bi$_2$Se$_3$ without an external source of Se shows unexpected bonding within the Mn layer due to Mn-Se bonding as Se diffuses out of the Bi$_2$Se$_3$ layer into the growing Mn film. The Se outdiffusion is further evidenced by changes in Bi core levels within the Bi$_2$Se$_3$ layers indicating primarily Bi-Bi bonding over Bi-Se bonding. No outdiffusion of Se occurred when excess Se is supplied with Mn, indicating the importance of supplying enough chalcogen atoms with deposited metals. However, Bi$_2$Se$_3$ core-level photoelectrons exhibited a rigid chemical shift toward higher binding energy after depositing a monolayer of MnSe$_2$, indicating a dipole within the overlayer. Stoichiometry calculations indicated that the monolayer forms MnSe preferentially over the transition metal dichalcogenide (TMD) phase MnSe$_2$, providing a consistent picture of the dipole formation in which a plane of Se anions sits above Mn cations. This study shows that chemical diffusion and dipole formation are important for Mn-Bi$_2$Se$_3$ and MnSe$_2$-Bi$_2$Se$_3$ and should be considered carefully for TMD/TI interfaces more generally.

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I. INTRODUCTION

Proximity effects between materials can induce new properties at their interface that would not otherwise exist separately. Magnetic overlayers brought into contact with topological insulators (TIs) such as Bi$_2$Se$_3$ can break time-reversal symmetry and affect spin properties leading to quantum phenomena, i.e., the quantum anomalous Hall effect (QAHE) [1,2] and skyrmions [3]. Three-dimensional (3D) ferromagnets such as $\alpha$-MnSe (111) or even two-dimensional (2D) materials such as 1T-MnSe$_2$ [4] are promising candidates to induce these surface magnetic and spin-related effects in Bi$_2$Se$_3$ [5]. A crucial component to realizing these advanced interface effects is an abrupt and clean interface between Bi$_2$Se$_3$ and the ferromagnetic overlayer, particularly for MnSe$_2$ [6], indicating a dipole within the overlayer. Molecular beam epitaxy (MBE) growth and subsequent transfers to measurement systems carried out completely in ultrahigh vacuum (UHV) provided a method to create and analyze high-quality interfaces without interlayer contamination or surface oxidation. Interlayer contaminants can alter the interface electronic and magnetic properties, and the electronic properties of both constituents will begin to degrade if exposed to oxygen. While UHV helps remove external contaminants, interdiffusion and other chemical interaction at the interface between Bi$_2$Se$_3$ and the magnetic overlayer can still occur. Many metals including Mn have been used to dope or alloy with Bi$_2$Se$_3$ [6] so that the chemical interaction between the two layers could in principle dominate the electronic and magnetic properties at their interface [7,8]. Furthermore, Mn and Bi$_2$Se$_3$ can form a magnetic ternary alloy Bi$_2$MnSe$_4$, which is also a promising system to investigate novel quantum and topological states [9–11].

Surface-sensitive techniques such as x-ray photoelectron spectroscopy (XPS) can provide insight into the impact of Mn-containing overlayers on the electronic structure of Bi$_2$Se$_3$ and how Mn incorporates with Bi$_2$Se$_3$ at the MnSe$_2$ interface. We used XPS connected via a UHV suitcase to a MBE system to investigate chemical interaction at the interface between a selenide transition metal dichalcogenide (TMD) and Bi$_2$Se$_3$. We compared the chemical effects on pristine Bi$_2$Se$_3$ of Mn-containing overlayers in two different scenarios: (i) extreme Se-deficient case, i.e., Mn deposited alone onto Bi$_2$Se$_3$, and (ii) Mn deposited with excess Se. In the case of pure Mn metal on Bi$_2$Se$_3$ we saw that Mn reacts with the Bi$_2$Se$_3$ surface where Mn bonds with Se extracted by diffusion from the Bi$_2$Se$_3$ substrate. In contrast, when excess Se was provided, there was little chemical interaction observed at the interface between MnSe$_{2-x}$ and Bi$_2$Se$_3$. Instead, MnSe$_{2-x}$ on Bi$_2$Se$_3$ rigidly shifted subsurface Bi$_2$Se$_3$ core levels toward higher binding energy, indicating the
formation of a negative surface/interface dipole. The presence of this dipole is likely due to the growth of primarily \( \alpha \)-MnSe instead of the 1T-MnSe\(_2\) 2D phase, which is supported by topographic scanning tunneling microscopy (STM) images and spectroscopy. Our XPS core level analysis combined with controlled depositions, air-free transfers, and surface analysis provides a consistent picture of chemical diffusion and dipole formation at the Mn selenide/Bi\(_2\)Se\(_3\) junction, revealing interfacial effects that may be relevant to TMD/ TI heterostructures in general.

II. METHODS

A. Molecular beam epitaxy

Prior to Bi\(_2\)Se\(_3\) sample growth, Al\(_2\)O\(_3\)(0001) substrates were annealed in the air at 1000 °C for 3 h. The substrates were then transferred to a home-built molecular beam epitaxy (MBE) chamber and heated to a growth temperature of 250 °C. Bi and Se were evaporated from standard Knudsen cells. A Se/Bi flux ratio (by thickness) of 20:1 was used during the growth with the Bi cell at 575 °C and the Se cell at 180 °C. Bi\(_2\)Se\(_3\) was deposited at a growth rate of 1 nm/min. After the growth, the samples were cooled down to room temperature in vacuum. A 20 nm Se capping layer was deposited on top of Bi\(_2\)Se\(_3\) to prevent oxidation before taking the sample out of the chamber.

Both Mn-Bi\(_2\)Se\(_3\) and MnSe\(_{2-x}\)-Bi\(_2\)Se\(_3\) were grown in a separate MBE chamber (Veeco GEN930). The Se-capped Bi\(_2\)Se\(_3\) samples were mounted onto 18 mm flag style paddles and loaded into the MBE chamber using special adapters to the 75 mm uniblock sample plates. The samples were first gently annealed at 170 °C to remove the Se capping. Reflection high-energy electron diffraction (RHEED) was used to monitor the sample surface and ensure that the cap removal is complete. During deposition, both Mn and Se were evaporated from a Knudsen cell. For the Mn-Bi\(_2\)Se\(_3\) sample, Mn was deposited at room temperature with a beam flux of \(5\times10^{-9}\) torr. For the MnSe\(_{2-x}\)-Bi\(_2\)Se\(_3\) sample, Mn and Se were codeposited at 250 °C with a flux ratio of \(\sim 1:5\) by beam equivalent pressure.

B. In vacuo transfer and XPS measurement

To prevent oxidation, all samples were transferred between the MnSe\(_{2-x}\) growth chamber and XPS chamber using an ultrahigh vacuum (UHV) suitcase tool that carries the 18 mm sample paddles. XPS measurements provided chemical composition and bonding information for the decapped Bi\(_2\)Se\(_3\) surface and two overlayer samples, 1.5 nm of Mn metal and 1 monolayer (ML) of MnSe\(_{2-x}\). XPS was performed using a PHI VersaProbe 5000\textsuperscript{TM} system equipped with a Scanning XPS Microprobe X-ray source (h\(\nu_{\text{AlK}α}=1486.6\) eV; full width at half-maximum \(\leq 0.5\) eV), and a hemispherical energy analyzer with a pass energy of 23.5 eV and a 0.05 eV step. To minimize the effects of charging, the XPS system is equipped with a two-stage sample surface neutralization system consisting of a 10 eV electron flood gun accompanied by a 10 eV Ar\(^+\) ion beam. Photoelectrons were collected at a takeoff angle of 45°. Chemical composition was determined using the PHI MultiPak analysis suite to determine relative atomic concentration using relative sensitive factors [12] for Bi 5d, Bi 4f, Se 3d, and Mn 2p core levels of 26.089, 180.178, 14.492, and 53.366, respectively. Additional curve fitting was performed using asymmetric Gaussian or Voigt line shapes for metallic and nonmetallic states, respectively.

![FIG. 1. Mn 2p core level spectrum of deposited Mn metal indicating unexpected Mn-Se bonding as shown by satellite features in green shifted to higher binding energy as indicated by the arrows.](image-url)
FIG. 2. (a) Bi 4f core level showing the binding energy difference before (black) and after (red) 1.5 nm Mn deposition. Depositing 1.5 nm of Mn causes Bi core levels of Bi₂Se₃ to shift toward higher binding energy. (b) Se 3d and Bi 5d core levels. The Se 3d peaks do not exhibit the same 1.74 eV shift as the Bi core levels.

C. STM measurement

STM measurements were performed at 5 K using a CreaTeC LT-STM. Samples were transferred from the MBE growth chamber to the STM system through a UHV suitcase to preserve the as-grown quality of the films. No additional preparation of the surface was performed prior to measurement. Images were obtained in constant current mode with a setpoint of 50–500 pA and sample bias in the range of 0.5–1 V. Tunneling spectra were obtained by adding a 20–50 mV modulation voltage at 1063 Hz to the dc bias and measured using a lock-in amplifier. Image processing and analysis were performed using WxSM [13].

III. RESULTS

A. Bare Bi₂Se₃ surface

The bare surface of decapped Bi₂Se₃ was first studied to check for surface contamination and establish a baseline for Bi and Se XPS peak centers and stoichiometry. Wide energy range XPS surveys and detailed region scans showed no detectable core level signals other than the expected Bi and Se levels and related Auger electron features. The decapped surface showed sharp Bi 5d, Bi 4f, and Se 3d doublets with binding energies (BE) consistent with previously measured Bi₂Se₃ core levels [14,15]. The stoichiometry of the Bi₂Se₃ surface (Table I) was obtained by comparing the relative peak areas below the Bi 5d, Bi 4f, and Se 3d spectra where the corrected peak areas for Bi 5d and 4f were averaged together. The Bi:Se ratio was found to be 1.03 ± 0.02 indicating a Se-deficient surface (Table I). The XPS spectra also show a pair of small broad features near the 285 eV region where a C 1s line would be. However, neither feature is a C 1s feature but instead they are related to Se Auger electron transitions [16]. Since the O 1s line was below the detection threshold of the survey scan, a detailed region scan was not performed. The lack of surface contamination of the samples transferred via a UHV suitcase confirms the quality of the suitcase transfer and its ability to maintain a clean, oxide-free sample surface.

B. Mn-Bi₂Se₃

To study the interaction of Mn metal with the Bi₂Se₃ surface, we deposited Mn metal only, e.g., without Se, onto Bi₂Se₃. The wide range survey scan [Fig. S1(a) of the Supplemental Material] shows virtually identical peaks to the bare Bi₂Se₃ surface with the addition of Mn-related peaks, notably the Mn 2p core level. Detailed region scans of the Mn 2p core level [Fig. 1(a)] show additional satellite features shifted to higher BE relative to the metallic Mn peak features. The shift to higher BE indicates Mn-Se bonding as evidenced by the increase in BE and charge state of the Mn ions. Additionally, the Mn 2p line shape is consistent with past measurement and modeling of manganese octahedrally coordinated to form MnX₆ clusters [17,18].

There are two likely sources for the Se observed in the Mn metal overlayer: (i) Se pulled from the Bi₂Se₃ layers below, or (ii) residual Se from the growth chamber. The Bi core levels provide insight into the source of this Se in the Mn
FIG. 4. (a) Bi 4f core level showing the binding energy difference before (black) and after (blue) 1 ML MnSe$_{2-x}$ deposition. (b) Se 3d and Bi 5d core levels. The Se 3d peaks do not exhibit the same 1.74 eV shift as the Bi core levels. Depositing 1 ML of MnSe$_{2-x}$ causes the Bi and Se core levels to rigidly shift 0.84 eV toward higher binding energy.

layer. Both Bi 5d and 4f core levels [Figs. 2(a) and 2(b)] exhibit a 1.74 eV shift toward lower BE after Mn metal was deposited. However, such a large negative BE shift is not seen in the Se 3d core level [Fig. 2(b)]. This large negative BE shift in bismuth core levels has been previously identified as Bi$_2$ layer formation within the bismuth selenide [14,19]. A careful deconvolution of the Bi 4f core level spectra (Fig. 3) reveals a small feature at higher BE compared to the dominant Bi 4$f_{7/2}$ peak. This smaller feature has BE consistent with the Bi$_2$Se$_3$ previously observed, indicating that Mn is not fully converting the subsurface Bi$_2$Se$_3$ into bismuth bilayers. Such changes in the bismuth core levels were not observed in earlier XPS studies on Mn-doped Bi$_2$Se$_3$ [20]. This is evidence that the Se in the Mn layer has diffused out of the Bi$_2$Se$_3$ leaving behind Bi$_2$ bilayers rather than Se pulled from the surrounding chamber.

C. MnSe$_{2-x}$-Bi$_2$Se$_3$

There is a very different interface picture on the side of Bi$_2$Se$_3$ when Mn is deposited with excess Se in a growth targeted for one unit-layer of MnSe$_2$. Figure 1(b) shows that the Mn 2p core level of MnSe$_{2-x}$ shows a similar peak structure but shifted 1.4 eV toward higher BE compared to that of the deposited Mn metal on Bi$_2$Se$_3$ in Fig. 1(a). The similar deconvolved peak structure confirms that the Mn metal BE shown above is consistent with Mn-Se bonding. However, all the photoelectron BE measured from Bi$_2$Se$_3$ after depositing 1 ML MnSe$_{2-x}$ exhibit a rigid 0.8 eV chemical shift toward higher BE compared to bare Bi$_2$Se$_3$ [Figs. 4(a) and 4(b)]. This shift toward higher observed BE indicates the presence of a negative interface dipole. Given the increase in BE, such a dipole represents an extra potential barrier at the free surface that escaping core level photoelectrons most overcome, shown schematically in Fig. 5.
We used the relative peak areas below deconvolved Mn 2p, Se 3d, and averaged Bi 4f and 5d core level features to calculate the surface stoichiometries shown in Table I. Based on the Bi:Se ratio of 1.03 as observed for the bare Bi2Se3, the ratio of Mn:Se is 1.0 ± 0.1, indicating a Se deficiency, and the chemical phase of the manganese selenide overlayer is more likely MnSe instead of MnSe2. The formation of MnSe versus the van der Waals material MnSe2 helps explain the origin of the observed dipole. A single monolayer of MnSe would form a negatively charged plane of Se ions on positively charged Mn creating the dipole that retards exiting electrons and causes the core level shift to higher BE.

Consistent with these XPS results, STM surface characterization suggests that α-MnSe(111) is the dominant phase within the MnSe2−x interfacial region. Atomic resolution of a typical α-MnSe(111) island is shown in Fig. 6(a). The inset shows a Fourier transform of the area confirming well-ordered structures with triangular symmetry and a lattice constant of 3.97 ± 0.04 Å consistent with α-MnSe(111) [22]. The topographic line profiles and dI/dV spectroscopy [Figs. 6(c) and 6(d)] indicate that α-MnSe(111) islands have an apparent height of 2.9 ± 0.2 Å and an energy gap of 700 meV. The observed gap is smaller than the 3 eV gap expected for bulk α-MnSe [22], but the measured step height and atomic lattice constant is in good agreement with other measurements of α-MnSe(111) [8], consistent with our expected XPS stoichiometry. Figure 6(b) also shows that a small secondary phase distinct from α-MnSe(111) exists on the surface. In comparison to α-MnSe, these small (<10 nm diam) islands show apparent heights of 5.8 ± 0.1 Å and exhibit a distinct dI/dV spectrum [Fig. 6(d)] with a much smaller gap. This is suggestive of 1T-MnSe2 formation, but these smaller secondary phase regions are too small and low density to measure with XPS for further characterization.

IV. ANALYSIS

A. Mn-Bi2Se3

Pure Mn metal interacts strongly at the interface with Bi2Se3 where Se diffuses out of Bi2Se3 into the Mn overlayer creating clusters of MnSe2−x as indicated by the satellite features in the Mn 2p core level [Fig. 1(a)]. The removal of Se from Bi2Se3 also leaves behind metallic Bi bilayers below the interface as evidenced by the strong 1.74 eV negative binding energy shifts of the Bi 5d and 4f core levels. This Se outdiffusion has been observed in other Bi2Se3-metal interfaces with Cr and Fe overlayers [15]. The difference in chemical reactivity between MnSe and MnSe2 may also factor into the MnSe formation. The chemical equations for Mn reaction at the surface of Bi2Se3 are Mn+(1/3)Bi2Se3 → MnSe+(2/3)Bi for MnSe, and Mn+(2/3)Bi2Se3 → MnSe2+(4/3)Bi for MnSe2. The heat of reaction ΔHR per metal atom for MnSe (HF = −155.6 kJ/mol) [23] versus Bi2Se3 (HF = −140.2 kJ/mol) [24] is −108.8 kJ/mol or −1.12 eV/metal atom compared with MnSe2(HF = −180.5 kJ/mol) versus Bi2Se3, which yields −87 kJ/mol or −0.90 eV/metal atom, which is less reactive.

B. MnSe2−x-Bi2Se3

There is less chemical reactivity at the interface when Se is supplied alongside Mn during deposition. There is no evidence of Se outdiffusion or metallic bismuth formation below the MnSe2−x/Bi2Se3 interface as there was in the case of Mn metal. However, the existence of a single monolayer of MnSe2−x produces a 0.74 eV rigid shift toward higher binding energy observed in Bi2Se3 core level photoelectrons. This increase in binding energy is due to an interface dipole created by the dominant manganese monoselenide phase that formed over the MnSe2 phase. α-MnSe (111) forms a plane of negatively charged Se ions at the free surface sitting above positive Mn ions, which creates the observed negative dipole that slows escaping core level photoelectrons. Smaller regions of MnSe2 also exist on the Bi2Se3 surface, but these regions are too small and low density to compare results between MnSe and MnSe2 regions. Presumably, such a dipole would not exist in a predominantly MnSe2/Bi2Se3 interface.
V. OUTLOOK

VdW heterostructures of 2D magnets on topological insulators hold promise for many spintronic studies including spin-orbit torque [25,26], skyrmion magnetic textures [27,28], the proximity-induced quantum anomalous Hall effect [29–31], and intrinsic magnetic topological insulators [32–34]. Our results highlight the importance of chemical analysis and atomic-scale structural characterization to help realize the desired heterostructures with abrupt interfaces and minimal chemical migration. The coexistence of tall islands (2D magnet MnSe$_2$ on topological insulator Bi$_2$Se$_3$) and short islands (MnSe covalently bonded to Bi$_2$Se$_3$ to form a septuple vdW layer) indicates a path to make either distinct 2D magnet/topological insulator heterostructures or intrinsic MnBi$_2$Se$_4$ by appropriate optimization of the growth conditions [35]. More generally, this provides a method for synthesizing and characterizing high-quality TMD/TI vdW heterostructures with a variety of functionalities for the TMD (e.g., semiconductor, valleytronic, superconductor, etc.).

VI. CONCLUSIONS

We have used surface-science techniques to explore the interface chemistry between Bi$_2$Se$_3$ and Mn-containing overlayers, and we found surface reactions and diffusion between constituents that depend significantly on the Se conditions during deposition. Mn metal deposited without excess Se induces Se outdiffusion from the Bi$_2$Se$_3$ substrate. This behavior suggests that Mn should be supplied with excess selenium to avoid altering the Bi$_2$Se$_3$ on which the Mn is deposited. When Se is supplied with Mn, no outdiffusion of Se occurs. Rather, Bi$_2$Se$_3$ core level photoelectrons exhibit a rigid shift toward higher binding energy, indicating a surface dipole caused by the formation of primarily α-MnSe (111) with smaller regions of MnSe$_2$. The existence of this dipole can be useful to indicate the phase of the grown manganese selenide overlayer when exact stoichiometry analysis is challenging or impossible due to a shared core level. Further, this work points to the importance of supplying excess chalcogen during deposition to avoid chemical bonding with substrate chalcogen atoms. This evidence for chemical diffusion and dipole formation at the Mn selenide/Bi$_2$Se$_3$ junction reveals interfacial effects that may be relevant to TMD/TI heterostructures in general.

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