RSC Advances

PAPER

Cite this: RSC Adv., 2017, 7, 32294

Received 11th May 2017 Accepted 19th June 2017

DOI: 10.1039/c7ra05317g

rsc.li/rsc-advances

1. Introduction

The layered compound $Bi₂Se₃$ has been used for thermoelectric (TE) energy conversion for years.^{1,2} After being revealed as a topological insulator (TI), $Bi₂Se₃$ has become much more sought-after than ever before.³ Featuring a large bulk band gap (\sim 0.3 eV) and simple Dirac-cone shaped surface states, $Bi₂Se₃$ is advocated as the most promising TI for room temperature applications.^{4,5} Some fingerprints of $Bi₂Se₃$ TI surface states have already been detected in electronic transport measurements, such as B-A interference effects,⁶ anti-localization behaviors,⁷ and unique anisotropic magneto-resistance.⁸ These findings have promised novel quantum devices of $Bi₂Se₃$ beyond its traditional TE applications. Furthermore, high quality $Bi₂Se₃$ thin films can be readily grown on substrates such as Si ,⁹⁻¹¹ GaAs,¹² and InP^{13,14} at rather low temperatures $(\sim 500 \text{ K})$, which sensibly promotes bottom-up on-chip integration of $Bi₂Se₃$ devices.

However, as a key step towards device realization, patterning $Bi₂Se₃$ thin films is still of challenge. Physical bombardment of argon ions and reactive ion etching offer very little selectivity of the sample material over photoresist and carbonization of photoresist masks due to long term exposure to high energy ion beams may ensue. The complicated operation also hinders the popularity of dry-etch in processing weakly-bonded $Bi₂Se₃$. As compared to dry etching approaches, wet etching is an effective alternative due to its high etching rate, low cost and high selectivity. The etchants with different etching velocity, surface

Patterning $Bi₂Se₃$ single-crystalline thin films on Si(111) substrates using strong oxidizing acids

Lei Gao, ^{Dac} Handong Li,^{*a} Wuyang Ren,^b Gaoyun Wang,^a Hui Li,^a Zhihua Zhou,^a Haining Ji,^a Xiaobin Niu^a and Zhiming Wang^b

Acidic potassium dichromate solutions $(K_2Cr_2O_7-H_2SO_4$ and $K_2Cr_2O_7-HCl$) are applied for patterning single crystalline Bi₂Se₃ thin films on Si(111) substrates. In solutions with appropriate component proportions, vertical walls and mesa-shaped structures on the etching profiles of (001) Bi₂Se₃ films can be achieved. Stoichiometric etching behavior is noted for Bi₂Se₃ in K₂Cr₂O₇–H₂SO₄ etchant, while incongruently dissolution of Bi₂Se₃ in K₂Cr₂O₇–HCl is observed which leaves a Se deficient layer on the etched film surface. The chemical reaction kinetics of Bi_2Se_3 in the two different etchants are also discussed

roughness, and some other characteristics can further be worked out for various purpose. Among etchants ever employed for $Bi₂Se₃$ and other chalcogenides, iodine and/or bromine based chemistries promise selective surface etching at limited rate.¹⁵ For fast chemical cutting of bismuth chalcogenides, the most popular wet etchant ever used is aqueous solutions of $HNO₃$ and HCl mixture (diluted aqua regia).^{16–21} However, these highly corrosive solutions could cause a vigorous uncontrollable reaction with $Bi₂Se₃$, resulting in severe edge undercutting. The diluted aqua regia etchants become even detrimental for handling thin films because they either cause pattern disappearance or trigger the delamination of weakly adhering $Bi₂Se₃$ films from bottom substrates.²² **PAPER**
 **PARE PERECONSISTED BI₂Se₃ single-crystalline thin films on

Creative and all distance Signal Signal S**

In this work, oxidizing etchants based on $K_2Cr_2O_7-H_2SO_4$ and $K_2Cr_2O_7$ –HCl solutions are used for patterning Bi_2Se_3 thin film electrodes on Si, respectively. $K_2Cr_2O_7-H_2SO_4$ –HCl ternary system has been verified suitable for using in etching solutions of covalent GaAs and InP wafers before.²³ It's demonstrated that the simpler $K_2Cr_2O_7-H_2SO_4$ and $K_2Cr_2O_7-HCl$ binary aqueous solutions can also produce mesa-shaped structures with vertical walls of layered $Bi₂Se₃$ thin films in this research. Superior surface morphologies of the etched $Bi₂Se₃$ (001) over the unetched ones are obtained in solutions with proper acidities. However, incongruently etching behaviors on the $Bi₂Se₃$ surfaces are noted that a Se depletion surface layer is formed on $Bi₂Se₃$ etched in the $K₂Cr₂O₇$ –HCl solutions.

2. Experimental section

Single crystalline $Bi₂Se₃$ thin films grown on (111)-oriented Si substrates by physical vapor deposition are adopted in this study. Details of thin film growth can be found in our previously published work.¹¹ The thickness of $Bi₂Se₃$ films involved in this study is \sim 300 nm. Photoresist etching mask is made by standard photolithography. About 1 μ m thick resist (AZ5214) is

YAL SOCIETY
CHEMISTRY

[&]quot;State Key Laboratory of Electronic Thin Films and Integrated Devices, School of Microelectronics and Solid-State Electronics, University of Electronic Science and Technology of China, Chengdu 610054, China. E-mail: hdli@uestc.edu.cn

^bInstitute of Fundamental and Frontier Sciences, University of Electronic Science and Technology of China, Chengdu 610054, China

c Microsystem and Terahertz Research Center, Chinese Academy of Engineering Physics, Chengdu 610200, China

Table 1 H^+ concentration of $K_2Cr_2O_7$ –HCl (H₂SO₄) etchants

Etch solution	Proportion	$[H^+] \pmod{L^{-1}}$
$K_2Cr_2O_7$ -HCl	1:1	6
$K_2Cr_2O_7-HCl$	1:2	8
$K_2Cr_2O_7-HCl$	1:3	9
$K_2Cr_2O_7-H_2SO_4$	1:1	9
$K_2Cr_2O_7-H_2SO_4$	1:2	12
$K_2Cr_2O_7-H_2SO_4$	1:3	13.5

masked on each $Bi₂Se₃ film surface by spin coating. After being$ prebaked for 65 seconds at 100 degree, the samples are exposed to ultraviolet light for 20 seconds. Submicron ultraviolet mask aligner is applied in this step. After removing the unexposed photoresist in developer solution and rinsed in deionized water, patterned samples are obtained. To enhance the adhesion of photoresist, all the samples are post baked for 5 minutes at 110 degree.

The $K_2Cr_2O_7$ -HCl (H₂SO₄) etchants are prepared by intermixing 9 v/v% $K_2Cr_2O_7$ and 48 v/v% HCl (98 v/v% H_2SO_4) solutions in volume proportions of $1:1, 1:2$, and $1:3$ (thereafter denoted by $1 : n$ K₂Cr₂O₇–HCl (H₂SO₄), $n = 1, 2, 3$), respectively at room temperature. As in Table 1, the acquired acidities, as represent by H^+ concentration ([H $^+$]) in the solutions, are calculated to be 6 mol $\text{L}^{-1},$ 8 mol $\text{L}^{-1},$ and 9 mol L^{-1} for the 1 : 1, 1 : 2, and 1 : 3 $K_2Cr_2O_7$ –HCl etchants, respectively. For the 1 : 1 to 1 : 3 $K_2Cr_2O_7-H_2SO_4$ etchants, $[H^+]$ values are 9 mol L^{-1} , 12 mol L^{-1} , and 13.5 mol L^{-1} , respectively. The $\mathrm{Bi}_2\mathrm{Se}_3$ films are etched in these aqueous acidic solutions for given time and no stirring is carried out to isolate the influence from hydromechanical effect. After etching, the samples are rinsed in acetone until the photoresist can be stripped. To investigate the dissolution of $Bi₂Se₃$, ascertain the character of corresponding reactions and determine limiting stages of the process, various methods are involved in the experiments. The thickness of the etched films is measured by a surface profilometer (Dektak150, Veeco). Optical microscopy (Olympus BX51), atomic force microscopy (AFM), and scanning electron microscopy (SEM) are employed to investigate the surface morphologies of the obtained patterns. Surface structural and compositional details of etched samples are further studied by Raman and electron dispersive spectroscopy (EDX), respectively. The chemical identification and binding energy study of the etched $Bi₂Se₃$ films are conducted by X-Ray Photoelectron Spectroscopy (XPS, Omicron). Paper

Teste 1 If concerned on 23 Go-O-HCl(1/502) etchants

ECO-O-HCl(1/2017.²¹

ECO-O-HCl(1/2020)

ECO-O-HCl(1/2020)

ECO-O-HCl(1/2020)

ECO-O-HCl(1/2020)

ECO-O-HCl(1/2020)

ECO-O-HCl(1/2020)

ECO-O-HCl(1/2020)

ECO-O

3. Results and discussion

Fig. 1(a) shows a photograph of a $Bi₂Se₃$ film etched by using 1:2 K₂Cr₂O₇-HCl solution. A grid pattern of Bi₂Se₃ with designed 8 μ m line width and 50 \times 50 μ m² windows is left on the Si substrate after thoroughly etching. Fig. $1(b)$ exhibits Raman curves taken from the $Bi₂Se₃$ grid and the etched window area of the sample, respectively. Three strong peaks locating at 71 cm^{-1} , 131 cm^{-1} and 173 cm^{-1} are attributed to the A_{1g}^1 , E_g^2 , and A_{1g}^2 transitions of pure Bi_2Se_3 respectively,^{24,25}

Fig. 1 (a) An optical photograph of a 1 : 2 $K_2Cr_2O_7$ –HCl solution etched $Bi₂Se₃$ film with rectangular grid pattern. (b) Raman curves taken from the grid line (red) and window region (black) on (a), respectively. (c) Cross-sectional SEM image indicates the side-wall structure of the etched $Bi₂Se₃$ film in (a).

which indicates a good crystallinity of the un-etched part of $Bi₂Se₃$ film. While there are only two peaks of $Si(111)$ (299.4) cm^{-1} and 520 cm^{-1}) found on the Raman spectrum from the etched area, indicating a complete removal of $Bi₂Se₃$ without any deposits left on the Si substrate after chemical reaction. A cross-sectional SEM image of the etched $Bi₂Se₃$ pattern further manifests the details at the edge of the etched $Bi₂Se₃ film$. As shown in Fig. 1(c), the side wall of the etched groove is straight and perpendicular to the horizontal surface of $Bi₂Se₃ film which$ unambiguously depicts the superiority of the acid $K_2Cr_2O_7$ solution in obtaining mesa-shape microstructures of $Bi₂Se₃$.

For better describing the dissolving properties of $Bi₂Se₃$ in acid K₂Cr₂O₇ solutions, the etching depths (θ) of the Bi₂Se₃ film are measured at different time intervals (t) . A typical θ –t curve of $Bi₂Se₃$ in 1 : 2 K₂Cr₂O₇–HCl etchant is shown in Fig. 2(a), and a constant etching rate of \sim 120 nm min⁻¹ is calculated from its linear fitting, indicating a rate-controlled etching reaction. It must be pointed out that the etching almost ceases at depth of 350 nm which is possibly due to the consumption of $[H^+]$ in the solution. Also noted from the θ –t curve is the reaction delay of $Bi₂Se₃$ during the initial 15 seconds before the balanced etching rate can be reached. Such behavior is also observed in all other reactions. By carefully excluding any extrinsic contamination on sample surfaces, we suppose the initial dissolving of $Bi₂Se₃$ is hindered by a thick native oxidized layer formed on each sample surface which has suffered long-time exposure in air. The induction period varies a lot for different $Bi₂Se₃$ samples, despite of the same acidity and temperature conditions are adopted in those reactions, which is ascribed to the reaction dependence on different chemical compositions and/or thicknesses of the oxidized layers on $Bi₂Se₃$ samples, and would not be discussed here. After the induction period, the reaction accelerates and the dissolving process of $Bi₂Se₃$ exhibits clear dependence on both the reaction temperatures and acidities of the etchants. Fig. 2(b) shows the quantitative $Bi₂Se₃$ etching rate dependence on acidities of $K_2Cr_2O_7$ –HCl and temperatures (20 °C to 45 °C). In the $[H^+]$: 6 mol L^{-1} etchant, the chemical reaction is weak that rather slight dissolution of the film can be detected even after the solution temperature is elevated from 20 °C to 45 °C (actually, the etching rate is only increased from

Fig. 2 (a) Time-dependent etching depth evolution of Bi₂Se₃ in 1 : 2 K₂Cr₂O₇–HCl solution. (b) Temperature-dependent etching rates of Bi₂Se₃ in 1 : 1, 1 : 2, and 1 : 3 K₂Cr₂O₇-HCl solutions, respectively. Inset in (b) is a typical optical microscopic picture taken from a Bi₂Se₃ film with square grid pattern immersed in 1 : 3 K₂Cr₂O₇-HCl for 5 seconds. The arrow indicates an over-eroded site on the pattern.

10 nm min^{-1} to a few tens of nanometers per minute as temperature ramps up from 20 $^{\circ} \mathrm{C}$ to 45 $^{\circ} \mathrm{C}$). While after the $[\mathrm{H}^+]$ of the $K_2Cr_2O_7$ etchants is increased larger than 8 mol L^{-1} , the chemical reaction is obviously activated and the etching rates yield rapid increase at elevated temperatures. As judged from the measured rate curves depicted in Fig. 2(b), the dissolution of $Bi₂Se₃$ in 1 : 3 K₂Cr₂O₇–HCl is nearly three times faster than it in $1:2 K₂Cr₂O₇$ -HCl etchant at room temperature. To quantitatively describe the chemical reaction intensity of $Bi₂Se₃$ in the oxidizing acids with different $[H^+]$, we calculate the activation energies of etching reactions of $Bi₂Se₃$ in 1 : 1, 1 : 2, and 1 : 3 K2Cr2O7–HCl solutions (the corresponding $[H^+]$ is 6 mol L^{-1} , 8 mol L^{-1} , and 9 mol L^{-1} , respectively) from the plots using Arrhenius equation for these curves shown in Fig. 2(b), and the calculated activation energies are 0.88 eV, 0.74 eV, and 0.47 eV, respectively. It thus strongly manifests that the chemical reaction barrier of $Bi₂Se₃$ in the oxidizing acids decreases dramatically as the acidity increases in the etching solutions. However, such an intense reaction leads to severely uneven etching of the patterned film. As shown in inset of Fig. $2(b)$ is an optical microscopic picture of slightly etched Bi_2Se_3 by 1 : 3 $K_2Cr_2O_7$ - HCl solution at room temperature, on which it can be clearly figured out that the areas near the grid corners are attacked much faster than the other regions. As the etching proceeds, undesired "W"-shape depth profiles of the film gradually evolve and part of the $Bi₂Se₃$ grid pattern gets eventually dissolved.

Almost the same etching behaviors of Bi_2Se_3 in $K_2Cr_2O_7$ - $\rm H_2SO_4$ etchants are observed. The 1 : 2 in $\rm K_2Cr_2O_7$ – $\rm H_2SO_4$ ([H⁺]: 12 mol L^{-1}) etchant also yields the same etching rate of \sim 120 nm min⁻¹ as obtained in the 1 : 2 in K₂Cr₂O₇-HCl and can produce "U" shape etching profiles of the $Bi₂Se₃$ films. Anyway, the electronic characteristics of electrodes and heterojunctions based on the etched $Bi₂Se₃$ subject on the microscopic surface properties, it's therefore necessary to analyze the morphologies and stoichiometry of the $Bi₂Se₃$ surfaces etched by these two kinds of etchants. As illustrated in Fig. 3(a) is a SEM image of an as-deposited $Bi₂Se₃$ thin film. High dense growth spirals surrounded by straight steps are observed in the as-grown $Bi₂Se₃$ thin films (highlighted by dashed triangles in Fig. 3(a)). As measured by AFM, the root mean square (RMS) roughness of such surface is \sim 3 nm (in area of 5 \times 5 μ m²), as shown in Table 2. After being immersed in $1:2 K_2Cr_2O_7-H_2SO_4$ solution for 30

Fig. 3 SEM images of an as-deposited Bi₂Se₃ surface (a) and Bi₂Se₃ surfaces etched in 1 : 2 K₂Cr₂O₇ - H₂SO₄ etchant (b) and 1 : 2 K₂Cr₂O₇ - HCl etchant (c), respectively.

seconds which corresponds to an etching depth of ~ 60 nm, these pyramid morphologies completely vanish and a surface decorated with large amounts of irregular-shaped islands sized from \sim 20 nm to \sim 300 nm comes out as exhibited in Fig. 3(b). The surface morphology does not improve too much (RMS roughness \sim 2.5 nm) as compared to the as-deposited Bi₂Se₃ film. On such etched surface, a Se : Bi mole ratio of \sim 1.60 is detected by EDX measurement. The Se : Bi mole ratio doesn't change significantly after a further 30 seconds etching period indicates a stoichiometric dissolution of $Bi₂Se₃$ in the 1 : 2 in $K_2Cr_2O_7-H_2SO_4$ etchant. Fig. 3(c) shows a SEM image taken from a Bi_2Se_3 film etched in 1 : 2 in $K_2Cr_2O_7$ –HCl solution at \sim 60 nm depth too. A much smoother surface (RMS roughness \sim 1.5 nm) is obtained for Bi₂Se₃ in this etchant. Not identical to the case of Bi_2Se_3 in 1 : 2 in $K_2Cr_2O_7-H_2SO_4$, Se : Bi mole ratio on the 1 : 2 in $K_2Cr_2O_7$ –HCl etched surface is \sim 1.1 as measured by EDX, implying a Se-dissolving preferred reaction kinetics. As etching proceeds to \sim 270 nm depth, only Bi signal from the film can be detected by EDX indicating a full depletion of Se. It should be noted that the good epitaxial character with twodimensional surface terrace-step structures of initial $Bi₂Se₃$ films disappears after being etched in both solutions, which strongly implies that both etching processes would cause amorphous surface parts of the $Bi₂Se₃$ films. It's thus also reasonable to attribute the nano-sized islands with irregular Paper

Task 2 3:54:18 in first 805 consistents and 5:2 Birmie sais between disputes are commonly as article in the same of th

shapes (as shown in Fig. 3(b) and (c)) to be amorphous $Bi₂Se₃$ deposits after etching.

It's well known that wet chemical etching proceeds by oxidation of the semiconductor constituents followed by chemical dissolution of the oxides in suitable solvents. For etching covalent chalcogenides such as CdSe, CdTe, and HgCdTe, different dissolving rates between cations and chalcogens are often observed because the hydration of cations is much easier than that of nonfully-oxidized chalcogens on the material surfaces.²⁶–²⁹ As a result, the etched surfaces of chalcogenides are usually cation-deficient. While no cationdeficient surfaces of etched $Bi₂Se₃$ are obtained in our experiments, which suggests quite different reaction kinetics of layered $Bi₂Se₃$ as compared to that of covalent chalcogenides in oxidizing acids etchants. To further understand the chemical reaction kinetics of $Bi₂Se₃$ in oxidizing acids with different components, XPS study is carried out for characterizing the chemical status of $Bi₂Se₃$ surfaces etched in different acid $K_2Cr_2O_7$ solutions.

As shown in Fig. 4 are core level spectrum of Bi and Se elements detected by XPS from the samples etched in 1 : 2 $K_2Cr_2O_7$ -HCl and 1 : 2 $K_2Cr_2O_7$ -H₂SO₄ etchants with etching depth of ~ 60 nm, respectively. The binding energies (BE) obtained in the XPS analysis are standardized for specimen charging using C 1s peak as the reference at 284.8 eV. To precisely determine the peaks' positions, Lorentz–Gauss profiles and Shirley background have been taken for the deconvolution. As indicated in Fig. 4(a) is Bi 4f spectrum from the $K_2Cr_2O_7-H_2SO_4$ etched Bi_2Se_3 sample, in which the intensity contribution for spin–orbit-splitting Bi 4f peaks ($4f_{7/2}$ and $4f_{5/2}$) from either Bi-O bonding^{30,31} or Bi-Se bonding can be clearly distinguished upon deconvolution. The intensities of Bi–O bonding Bi peaks are much stronger than that of Bi–Se bonding Bi peaks, which suggest there exists a bismuth oxide (BiO_x) thin layer deposited on the $Bi₂Se₃$ surface during etching. While for the $K_2Cr_2O_7$ –HCl etched Bi₂Se₃ case, only a pair of Bi 4f peaks located at 158.82 eV and 164.11 eV are observed which can be

Fig. 4 Bi 4f and Se 3d XPS spectra of Bi₂Se₃ films after etched by 1 : 2 K₂Cr₂O₇ - H₂SO₄ etchant [(a) and (c)] and by 1 : 2 K₂Cr₂O₇ - HCl etchant [(b) and (d)], respectively. Experimental data are the open circles and fits are the solid lines.

precisely attributed to Bi–Se bonding of pure $Bi₂Se₃$ (Fig. 4(b)).³²⁻³⁴ Peaks referred to Se $3p_{1/2}$ and Se $3p_{3/2}$ can also be figured out in both fitted XPS spectra as exhibited in Fig. $4(a)$ and (b), and the broaden peak profiles depict a mixed valence states of Se on both samples.³⁵ The valence states of Se from $Bi₂Se₃$ films after etched by $K₂Cr₂O₇ - H₂SO₄$ and $K₂Cr₂O₇ - HCl$ are further investigated by analyzing the peak fitted Se 3d $(3d_{5/2})$ and $3d_{3/2}$) spectra as illustrated in Fig. 4(c) and (d), respectively. In either Se 3d curve, the pair of high BE component peaks belong to Se–Se bonding of element Se and the other pair are related to Bi–Se bonding, respectively. And no traces of any selenium oxides can be found on the etched $Bi₂Se₃$ surfaces in our XPS studies.³⁰ Therefore, it's reasonable to determine that there's only a $\text{Se}^{2-} \rightarrow \text{Se}^{0}$ oxidizing reaction happens for the Se element in both etching processes and the free element Se reactants are deposited onto the $Bi₂Se₃$ surfaces. Further, peak intensity contribution from Se–Se bonding dominates in Se 3d spectra of the $K_2Cr_2O_7-H_2SO_4$ etched Bi_2Se_3 sample while it becomes not prominent in $K_2Cr_2O_7$ –HCl etched one, which manifests that there're more Se deposits residual on $K_2Cr_2O_7$ - H_2SO_4 etched Bi_2Se_3 surface than those on $K_2Cr_2O_7$ –HCl etched one. **Exchanges**

The distribution and the Broadback Article and Schop considers by resident control on the consideration of the state are considered to the state of the state are considered to the state of the state of the sta

Based on the XPS observations, it should be feasible to determine that the surface chemistry is predominated by bismuth oxides and element Se all through the etching procedure in $K_2Cr_2O_7-H_2SO_4$ solutions. Therefore the surface reaction kinetics are only subjected by the rate of reactants reaches and leaves the etched surface.³⁶ As to our knowledge, there's only tempered chemical reactions happen between bismuth oxides and concentrated acids (such as H_2SO_4 and HCl) at room temperature. So do them between element Se and concentrated sulphuric acid. As a result, congruently dissolving of Bi and Se will dominate in the etching reaction which is confirmed by our experiment observations.

On the other hand, the valence states of Bi in $Bi₂Se₃$ remains intact during etching in $K_2Cr_2O_7$ -HCl solutions. It's thus suggested that the etching action on the $Bi₂Se₃$ surface is fully triggered by dissolving of Se. As a result, it leads to the observed incongruently etching behaviors of $Bi₂Se₃$ in $K₂Cr₂O₇$ -HCl etchants.

4. Conclusions

To summary, mesa-shaped etching profile of $Bi₂Se₃$ is obtained either in 1 : 2 $K_2Cr_2O_7$ -HCl or 1 : 2 $K_2Cr_2O_7$ -H₂SO₄ aqueous solutions with H $^+$ concentrations of 8 mol L^{-1} and 12 mol L^{-1} , respectively. Dissolving rates of $Bi₂Se₃$ are approximately 120 nm min⁻¹ in both the 1 : 2 K₂Cr₂O₇-H₂SO₄ and 1 : 2 $K_2Cr_2O_7$ -HCl etchants at room temperature. Though either etchant is qualified for preparing desired microscopic patterns of $Bi₂Se₃$, quite distinct dissolving kinetics between these two etchants signicantly lead to different reaction products on surfaces of the etched samples. In $1:2 K_2Cr_2O_7-H_2SO_4$ etching case, the surface stoichiometry of $Bi₂Se₃$ is found intact that the TI surface states would probably being reserved after etching. While the Se-deficient dissolving behavior of $Bi₂Se₃$ noted in the $K_2Cr_2O_7$ -HCl etchants implies surface quality degradation.

Nevertheless, slightly metal-rich surface may benefit a lowresistance ohmic contact formation between $Bi₂Se₃$ and electrode metals that has been argued crucial for improving performance of TE devices.³⁷ Considering chemical treatment of semiconductor surfaces is a convenient way to define the surface properties and manufacture micropatterns, our findings may serve as useful reference for fabricating $Bi₂Se₃$ -based devices with various purpose.

Acknowledgements

This work is supported by the National Natural Science Foundation of China (Grant No. 11104010, 61474014, 51272038, and 51302030) and Open Research Fund Program of the State Key Laboratory of Low-Dimentional Quantum Physics (No. 20120910).

References

- 1 C. Wood, Rep. Prog. Phys., 1988, 51, 459–539.
- 2 G. Chen, M. S. Dresselhaus, G. Dresselhaus, J. P. Fleurial and T. Caillat, Int. Mater. Rev., 2003, 48, 45–66.
- 3 D. Hsieh, Y. Xia, D. Qian, L. Wray, F. Meier, J. H. Dil, J. Osterwalder, L. Patthey, A. V. Fedorov, H. Lin, A. Bansil, D. Grauer, Y. S. Hor, R. J. Cava and M. Z. Hasan, Phys. Rev. Lett., 2009, 103, 4.
- 4 Y. Xia, D. Qian, D. Hsieh, L. Wray, A. Pal, H. Lin, A. Bansil, D. Grauer, Y. S. Hor, R. J. Cava and M. Z. Hasan, Nat. Phys., 2009, 5, 398–402.
- 5 H. Zhang, C. X. Liu, X. L. Qi, X. Dai, Z. Fang and S. C. Zhang, Nat. Phys., 2009, 5, 438–442.
- 6 H. L. Peng, K. J. Lai, D. S. Kong, S. Meister, Y. L. Chen, X. L. Qi, S. C. Zhang, Z. X. Shen and Y. Cui, Nat. Mater., 2010, 9, 225–229.
- 7 J. Chen, H. J. Qin, F. Yang, J. Liu, T. Guan, F. M. Qu, G. H. Zhang, J. R. Shi, X. C. Xie, C. L. Yang, K. H. Wu, Y. Q. Li and L. Lu, Phys. Rev. Lett., 2010, 105, 4.
- 8 J. Wang, H. Li, C. Chang, K. He, J. S. Lee, H. Lu, Y. Sun, X. Ma, N. Samarth and S. Shen, Nano Res., 2011, 5, 739–746.
- 9 G. H. Zhang, H. J. Qin, J. Teng, J. D. Guo, Q. L. Guo, X. Dai, Z. Fang and K. H. Wu, Appl. Phys. Lett., 2009, 95, 3.
- 10 H. D. Li, Z. Y. Wang, X. Kan, X. Guo, H. T. He, Z. Wang, J. N. Wang, T. L. Wong, N. Wang and M. H. Xie, New J. Phys., 2010, 12, 11.
- 11 H. D. Li, L. Gao, H. Li, G. Y. Wang, J. Wu, Z. H. Zhou and Z. M. Wang, Appl. Phys. Lett., 2013, 102, 4.
- 12 A. Richardella, D. M. Zhang, J. S. Lee, A. Koser, D. W. Rench, A. L. Yeats, B. B. Buckley, D. D. Awschalom and N. Samarth, Appl. Phys. Lett., 2010, 97, 3.
- 13 X. Guo, Z. J. Xu, H. C. Liu, B. Zhao, X. Q. Dai, H. T. He, J. N. Wang, H. J. Liu, W. K. Ho and M. H. Xie, Appl. Phys. Lett., 2013, 102, 4.
- 14 S. Schreyeck, N. V. Tarakina, G. Karczewski, C. Schumacher, T. Borzenko, C. Brune, H. Buhmann, C. Gould, K. Brunner and L. W. Molenkamp, Appl. Phys. Lett., 2013, 102, 4.
- 15 S. Augustine and E. Mathai, Mater. Res. Bull., 2001, 36, 2251– 2261.
- 16 C. Shafai and M. J. Brett, J. Vac. Sci. Technol., A, 1997, 15, 2798–2801.
- 17 L. M. Goncalves, J. G. Rocha, C. Couto, P. Alpuim, G. Min, D. M. Rowe and J. H. Correia, J. Micromech. Microeng., 2007, 17, S168–S173.
- 18 L. M. Goncalves, C. Couto, P. Alpuim and J. H. Correia, J. Micromech. Microeng., 2008, 18, 1171–1185.
- 19 J. P. Carmo, L. M. Goncalves and J. H. Correia, IEEE Trans. Ind. Electron., 2010, 57, 861–867.
- 20 L. M. Goncalves, P. Alpuim and J. H. Correia, J. Electron. Mater., 2010, 39, 1516–1521.
- 21 J. P. Carmo, M. F. Silva, J. F. Ribeiro, R. F. Wolffenbuttel, P. Alpuim, J. G. Rocha, L. M. Goncalves and J. H. Correia, Microsyst. Technol., 2011, 17, 1283–1291.
- 22 T. Ngai and U. Ghoshal, Wet etching of Bi2Te3 thin films compatible with microelectronic fabrication processes, 26th International Conference on Thermoelectrics, Jeju, Korea, June 2007.
- 23 S. Adachi, H. Kawaguchi and G. Iwane, J. Mater. Sci., 1981, 16, 2449–2456.
- 24 J. Zhang, Z. P. Peng, A. Soni, Y. Y. Zhao, Y. Xiong, B. Peng, J. B. Wang, M. S. Dresselhaus and Q. H. Xiong, Nano Lett., 2011, 11, 2407–2414.
- 25 K. M. F. Shahil, M. Z. Hossain, V. Goyal and A. A. Balandin, J. Appl. Phys., 2012, 111, 8.
- 26 R. Tenne and G. Hodes, Appl. Phys. Lett., 1980, 37, 428–430.
- 27 W. H. Chang, T. Lee and W. M. Lau, J. Appl. Phys., 1990, 68, 4816–4819.
- 28 I. M. Kotina, L. M. Tukhkonen, G. V. Patsekina, A. V. Shchukarev and G. M. Gusinskii, Semicond. Sci. Technol., 1998, 13, 890–894.
- 29 V. Srivastav, R. Pal and H. P. Vyas, Opto-Electron. Rev., 2005, 13, 197–211.
- 30 D. Kong, J. J. Cha, K. Lai, H. Peng, J. G. Analytis, S. Meister, Y. Chen, H. J. Zhang, I. R. Fisher and Z. X. Shen, ACS Nano, 2011, 5, 4698–4703.
- 31 A. J. Green, S. Dey, Y. Q. An, B. O'Brien, S. J. O'Mullane, B. Thiel and A. C. Diebold, 2016, arXiv: cond-mat.mtrl-sci/ 1601.04057.
- 32 V. V. Atuchin, V. A. Golyashov, K. A. Kokh, I. V. Korolkov, A. S. Kozhukhov, V. N. Kruchinin, S. V. Makarenko, L. D. Pokrovsky, I. P. Prosvirin, K. N. Romanyuk and O. E. Tereshchenko, Cryst. Growth Des., 2011, 11, 5507–5514.
- 33 M. T. Edmonds, J. T. Hellerstedt, A. Tadich, A. Schenk, K. M. O'Donnell, J. Tosado, N. P. Butch, P. Syers, J. Paglione and M. S. Fuhrer, J. Phys. Chem. C, 2014, 118, 20413–20419.
- 34 C. Han, J. Yang, C. Yan, Y. Li, F. Y. Liu, L. X. Jiang, J. C. Ye and Y. X. Liu, CrystEngComm, 2014, 16, 2823–2834.
- 35 V. A. Golyashov, K. A. Kokh, S. V. Makarenko, K. N. Romanyuk, I. P. Prosvirin, A. V. Kalinkin, O. E. Tereshchenko, A. S. Kozhukhov, D. V. Sheglov, S. V. Eremeev, S. D. Borisova and E. V. Chulkov, J. Appl. Phys., 2012, 112, 4. Paper

16 C. Shada and M. J. Breth, J. Brokn, C. Common, P. Alpointre, C. Min. A. V. Shahnkare, G. V. Paterskins, 2017. Downloaded Street, 2017. Downloaded on 2017. Downloaded on 2017. Downloaded article. Commons Article.
	- 36 Y. C. Lin, Y. C. Jian and J. H. Jiang, Appl. Surf. Sci., 2008, 254, 2671–2677.
	- 37 M. Hines, J. Lenhardt, M. Lu, L. Jiang and Z. Xiao, J. Vac. Sci. Technol., A, 2012, 30, 041509.