Controlled Growth of a Large-Size 2D Selenium Nanosheet and Its Electronic and Optoelectronic Applications

Jingkai Qin,†,§,# Gang Qiu,†,# Jie Jian,‡ Hong Zhou,† Lingming Yang,† Adam Charnas,† Dmitry Y. Zemlyanov,† Cheng-Yan Xu,§ Xianfan Xu,∥ Wenzhuo Wu,⊥ Haiyan Wang,†,‡ and Peide D. Ye*†,‡

†School of Electrical and Computer Engineering, Purdue University, West Lafayette, Indiana 47907, United States
‡School of Materials Science and Engineering, Purdue University, West Lafayette, Indiana 47907, United States
§School of Materials Science and Engineering, Harbin Institute of Technology, Harbin 150001, People’s Republic of China
∥School of Mechanical Engineering, Purdue University, West Lafayette, Indiana 47907, United States
⊥School of Industry Engineering, Purdue University, West Lafayette, Indiana 47907, United States

ABSTRACT: Selenium has attracted intensive attention as a promising material candidate for future optoelectronic applications. However, selenium has a strong tendency to grow into nanowire forms due to its anisotropic atomic structure, which has largely hindered the exploration of its potential applications. In this work, using a physical vapor deposition method, we have demonstrated the synthesis of large-size, high-quality 2D selenium nanosheets, the minimum thickness of which could be as thin as 5 nm. The Se nanosheet exhibits a strong in-plane anisotropic property, which is determined by angle-resolved Raman spectroscopy. Back-gating field-effect transistors based on a Se nanosheet exhibit p-type transport behaviors with on-state current density around 20 mA/mm at $V_{ds} = 3$ V. Four-terminal field-effect devices are also fabricated to evaluate the intrinsic hole mobility of the selenium nanosheet, and the value is determined to be 0.26 cm$^2$ V$^{-1}$ s$^{-1}$ at 300 K. The selenium nanosheet phototransistors show an excellent photoresponsivity of up to 263 A/W, with a rise time of 0.1 s and fall time of 0.12 s. These results suggest that crystal selenium as a 2D form of a 1D van der Waals solid opens up the possibility to explore device applications.

KEYWORDS: selenium, 1D crystal structure, 2D nanosheet, electrical transport, photoresponse

Tetragonal selenium (t-Se) is a one-dimensional elemental semiconductor material, in which Se atoms are covalently connected in a spiral chain along the c-axis with two adjacent atoms rotated by 120°. As shown in Figure 1a and b, all these atomic chains are stacked together in their radical direction by weak van der Waals interactions to form a hexagonal lattice structure. t-Se has attracted extensive attention due to its interesting properties such as high photoconductivity,1-5 high piezoelectricity,6-8 thermoelectricity,9 and nonlinear optical responses.10,11 The high photoconductivity makes it a promising candidate for applications in high-efficiency solar cells and optoelectronic devices.12,13 Recent reports have shown that Se nanosheet photodetectors exhibit a high responsivity of 100 mA/W at 620 nm light illumination with an ultrashort rise/decay time (1.4/7.8 ms).14 A variety of Se-based nanostructures have been obtained by different synthesis methods, such as a hydrothermal solution process and microwave-assisted synthesis in ionic liquids.14,15 Compared with solution-based techniques, which always introduce impurities due to the complex chemical processes involved, a vapor phase approach should be more efficient to obtain high-quality products.16,17 Due to the anisotropic chain-like crystal structure, Se tends to form 1D structures, such as nanowires, nanotubes, and nanobelts. These 1D structures with a high ratio of edges to bulk always exhibit poor electrical transport behaviors due to localized states and contacts, and the issue of electrical noise becomes critical with decreasing size, as described by Hooge’s rule.18,19 To expand its applications in high-performance electronic and optoelectronic devices, it would be useful to expand Se nanostructures to a two-dimensional form.

Received: July 7, 2017
Accepted: September 26, 2017
Published: September 26, 2017
Recently, tellurium (Te), which possesses the same crystal structure as Se, has been successfully synthesized in a 2D form using a substrate-free solution process, and 2D tellurium (termed as tellurene) flakes exhibit strong in-plane anisotropic properties and high carrier mobility. Inspired by this success, using a physical vapor deposition (PVD) method, we successfully obtained highly crystalline Se nanosheets with interesting zigzag edge structure, with a minimum thickness of Se of 5 nm. Scanning transmission electron microscopy (STEM) and angle-resolved Raman spectroscopy confirm that the single-crystal Se nanosheet has an oriented growth direction along ⟨12̅10⟩, which is very different from previous studies. To the best of our knowledge, no works have been reported on the synthesis of such large-size 2D ultrathin Se nanosheets. Back-gated field-effect transistors (FETs) based on our 2D PVD-grown Se nanosheet exhibit p-type transport behaviors with on-state current density around 20 mA/mm at $V_{ds} = 3$ V. Four-terminal field-effect devices are also fabricated to evaluate the intrinsic hole mobility of the Se nanosheet, and the value is determined to be 0.26 cm$^2$ V$^{-1}$ s$^{-1}$ at 300 K. Se nanosheet phototransistors are found to have an excellent photoresponsivity (263 A/W) with a 0.10 s rise time and a 0.12 s fall time. This work demonstrates that the crystal selenium nanosheet, which is built up by 1D van der Waals material selenium, has very interesting properties for further exploration of device applications as a type of 2D semiconductor material.

RESULTS AND DISCUSSION

The Se nanosheets were obtained using the PVD method, and the typical growth setup used is illustrated in Figure 1c. Se powder was placed in a quartz glass tube as the precursor. A plan-view SEM image of Se nanosheets as-grown on a Si substrate is shown in Figure 1d, in which inclined nanosheets (as indicated by arrows) with a density of more than 50% were uniformly distributed on the surface of polycrystalline Se films. Figure 1e shows the representative morphology of the Se nanosheets at high magnification in SEM. Most of the nanosheets exhibit a saw-like structure with zigzag edges on the single narrow side. The average width of the nanosheets is about 8 μm, and the maximum length of the nanosheets could reach up to 50 μm. In the SEM images, the nanosheets are almost electron transparent, suggesting that they are very thin.

Figure 1. PVD-grown large-area Se nanosheets and material characterization. (a, b) Atomic structure of selenium. (c) Schematic diagram of the PVD method. (d) Low-magnification SEM image of as-prepared selenium nanosheets on a Si(111) substrate. (e) Enlarged view of typical Se nanosheets with saw-like structure and (f) feather-like twin structure.
Figure 3. STEM characterization of a Se nanosheet. (a) Bright-field TEM image of Se nanosheets with saw-like structure. (b) HAADF-STEM image and (c) corresponding SAED pattern of Se nanosheets (boxed area in (a)). (d, e) Bright-field TEM image of feather-like Se nanosheets with different magnification. (f) SAED pattern obtained at the coherent crystal boundary (boxed area in (e)), with common spots showing the mutual twin boundary of the \{0104\} plane.

Figure 4. Angle-resolved Raman spectra for few-layer Se nanosheets. (a) Raman spectrum of a Se nanosheet with a thickness of 15 nm. The inset shows the atomic vibration patterns of E\textsubscript{1}, E\textsubscript{2}, and A\textsubscript{1} phonon modes in selenium. (b) Raman spectra evolution with angles between crystal orientation and incident laser polarization. (c, d) Polar figures of Raman intensity corresponding to E\textsubscript{2} and A\textsubscript{1} modes located at 233 and 237 cm\textsuperscript{-1}. 
Se(0), respectively. The Se 3d oxidized peak located 59.9 eV cannot be detected, indicating that the Se nanosheets are poorly crystallized with good quality. Figure 2c and d show the height images with different magnifications, and the surface is atomically flat with a surface roughness \( R_m \) of 190 pm, even less than that of the SiO\(_2\) surface (about 200 pm), indicating that the Se nanosheets are highly crystallized with good quality. Figure 2c and d show the height profiles of the saw-like and twin structure selenium nanosheets, with an average thickness of 15 nm. It should be noted that the twin structure could be clearly detected according to contrast difference by optical microscopy, while the corresponding AFM image shows a uniform height profile. This result suggests that the bicrystalline Se nanosheet divided from the middle has two separate single-crystal grains with different orientations. XPS analysis was also conducted to determine the quality of Se nanosheets (Figure S2). The strong peaks located at 55.5 and 54.3 eV correspond to Se 3d\(_{3/2}\) and 3d\(_{5/2}\) binding energy of Se\(^{0}\), respectively. The Se 3d oxidized peak located 59.9 eV cannot be detected, indicating that the Se nanosheets are elemental crystals without obvious oxidation.\(^{23,24}\)

Transmission electron microscopy (TEM) was employed to identify the microstructure and growth directions of the Se nanosheets. The nanosheets could be directly transferred onto Cu grids. The low-magnification bright-field TEM image (Figure 3a) shows a 2D saw-like structure, similar to what was observed in the optical microscopy image. The high-angle annular dark field STEM (HAADF-STEM) image could provide Z-contrast with atomic lateral resolution \( Z = \text{atomic number} \). As shown in Figure 3b, Se helical atomic chains could be clearly resolved with a fringe spacing of 0001 lattice planes of about 5.0 Å. The corresponding selected area electron diffraction (SAED) image, which was obtained along the \([10\overline{1}0]\) zone axis of an individual nanosheet, exhibits a set of 2-fold and rotational symmetry pattern, indicating the Se nanosheets are highly crystalline. Combined with the HAADF-STEM image and SAED pattern, we can determine that the Se nanosheets have first grown along the \([0001]\) direction into nanoribbons, and then these nanoribbons expand in parallel along the \([12\overline{1}0]\) direction, leading to the formation of a large-area nanosheet with zigzag edges. TEM analysis was also conducted to determine the crystal structure of the feather-like Se nanosheets with twin structure. As the bright-field TEM images with different magnification show (Figure 3e and f), the angle of two twinned grains is about 124°. By carefully correlating the SAED pattern obtained from the boundary area (Figure 3g), it could be found that \([0\overline{1}04]\) diffraction spots are perpendicular to the twin boundary, indicating that the twin boundary is a \([0\overline{1}04]\) twin. EDX and EELS also confirmed that the synthesized nanosheet is an elemental Se crystal (Figure S3).

The in-plane anisotropy of the Se nanosheet is investigated by angle-resolved Raman spectroscopy at room temperature. The angle between the laser polarization direction and \([0001]\) helical chain direction of the nanosheet is defined as \( \theta \), which could be tuned by rotating the sample in steps of 15° during measurement. Figure 4a shows the typical Raman spectrum of a 15 nm thick nanosheet with an angle \( \theta \) of 45°. Consistent with the previous observations in bulk selenium,\(^{25,26}\) three active Raman photon modes are clearly observed. Raman peaks located at \( \sim 233 \) and \( 237 \) cm\(^{-1}\) are related to the \( E_2 \) and \( A_1 \) modes, respectively. Meanwhile, one degenerate \( E_1 \) mode caused by \( a \)-axis rotation is also identified. Figure 4b depicts the evolution of the Raman spectrum as the sample is rotated in steps of 30° from \( -90° \) to \( 90° \), and a clear intensity change could be observed. The peak intensities of different modes are extracted by fitting with a Gauss function and plotted into the corresponding polar figures (Figure 4c,d). It should be noted that the degenerate \( E_2 \) mode is hard to extract due to the relatively weak intensity. Both the \( E_2 \) and \( A_1 \) modes exhibit significant intensity change with the polarization angle. The \( A_1 \) mode is a maximum at an angle of \( \sim 90° \), which corresponds to the direction vertical to the Se chains, while \( E_2 \) has a maximum value at an angle of \( \sim 45° \). Our results suggest that the in-plane anisotropic properties of Se nanosheets could be easily identified by Raman spectroscopy. It could also be confirmed that the Se nanosheets are grown along the \( [1 \overline{2} 1 0] \) direction, which matches well the STEM results.

Trigonal selenium possesses a highly anisotropic crystal structure. Under thermodynamic equilibrium conditions, it tends to grow into 1D structures along the \([0001]\) direction. However, the products in our experiments exhibit a 2D structure with very small thickness, which is very different from the previous bulk Se. Time-dependent experiments were conducted to explore the growth mechanism of Se nanosheets. Figure 5a shows the typical morphology of products when the growth temperature just reaches 160 °C. Amorphous nanospheres with diameters ranging from 10 to 30 μm appeared on the Si substrate surface. With the temperature rising up to 210...
amorphous nanospheres start to crystallize and form polycrystalline films, with numerous short nanorods protruding out of the surface (Figure 5b and Figure S4). Based on previous studies, heat treatment could accelerate the transformation of amorphous selenium to crystalline selenium, and Imura et al. have successfully prepared polycrystalline films consisting of nanoparticles by in situ thermal treatment for photodetector applications. The preferable growth of 1D Se nanorods is kinetically favored, since the binding energy of Se atoms along the c-axis is much higher than that along the ⟨1210⟩ directions. Besides, these 1D nanorods possess high chemical activity especially at the vertexes and ridges, which could play the role of roots for the subsequent nanosheet growth. Interestingly, as the reaction continues, the growth along the c-axis is partly hindered, and the nanorods tend to expand in parallel along the ⟨1210⟩ direction to form 2D thin nanosheets as shown in Figure 5c. Under the same magnitude of supersaturation, the growth of Se nanosheet is mainly driven by surface energy as well as effective activation sites count. In the initial crystallization, 1D growth of the nanorod is dominant due to the distinct surface energy difference. However, after the nanorod has grown to a certain length, Se(g) molecules would seldom fall within the end cavity of a nanorod due to the large free path at such high temperatures. It is more likely that they would impinge on the nanorod surface and diffuse from site to site until they encounter a defect or void of the crystal; thus the growth along the ⟨1210⟩ direction would be enhanced. Actually, at the lateral side of the Si substrate, where the downstream Se(g) molecules have greater odds of encountering a (0001) facet of the nanorod, the nanorod would continue to grow along the ⟨0001⟩ direction into the nanowire with lengths up to 200 μm (Figure S5). Figure 5d illustrates the products’ morphology when the growth duration is up to 60 min; the nanorods have completely turned into long, saw-like ultrathin nanosheets.

\(^{10}\)C, amorphous nanospheres start to crystallize and form polycrystalline films, with numerous short nanorods protruding out of the surface (Figure 5b and Figure S4). Based on previous studies, heat treatment could accelerate the transformation of amorphous selenium to crystalline selenium, and Imura et al. have successfully prepared polycrystalline films consisting of nanoparticles by in situ thermal treatment for photodetector applications. The preferable growth of 1D Se nanorods is kinetically favored, since the binding energy of Se atoms along the c-axis is much higher than that along the ⟨1210⟩ directions. Besides, these 1D nanorods possess high chemical activity especially at the vertexes and ridges, which could play the role of roots for the subsequent nanosheet growth.

Interestingly, as the reaction continues, the growth along the c-axis is partly hindered, and the nanorods tend to expand in parallel along the ⟨1210⟩ direction to form 2D thin nanosheets as shown in Figure 5c. Under the same magnitude of supersaturation, the growth of Se nanosheet is mainly driven by surface energy as well as effective activation sites count. In the initial crystallization, 1D growth of the nanorod is dominant due to the distinct surface energy difference. However, after the nanorod has grown to a certain length, Se(g) molecules would seldom fall within the end cavity of a nanorod due to the large free path at such high temperatures. It is more likely that they would impinge on the nanorod surface and diffuse from site to site until they encounter a defect or void of the crystal; thus the growth along the ⟨1210⟩ direction would be enhanced. Actually, at the lateral side of the Si substrate, where the downstream Se(g) molecules have greater odds of encountering a (0001) facet of the nanorod, the nanorod would continue to grow along the ⟨0001⟩ direction into the nanowire with lengths up to 200 μm (Figure S5). Figure 5d illustrates the products’ morphology when the growth duration is up to 60 min; the nanorods have completely turned into long, saw-like ultrathin nanosheets.

The formation of Se nanosheets could be well explained by the vapor–solid (VS) growth mechanism, and similar growth phenomena have been reported in the preparation of ZnSe nanobelts. It is worth noting that some of the nanosheets exhibit a feather-like structure with mirror twins.
We believe that they were developed from sections of intersected nanorods, which are introduced at the initial crystalline stage (Figure S6). These crossed nanorods could act as growth roots and enable the identical lateral growth on both sides of the nanosheet, leading to the symmetrical structure.\textsuperscript{33,34}

Se nanosheet FETs were fabricated using electron beam lithography (EBL), thermal evaporation, and a lift-off process. Ni/Au (30/100 nm) were selected as metal contacts, which could significantly reduce the contact resistance in p-type FET devices due to the relatively high work function.\textsuperscript{35} Figure 6 shows the electrical characteristics of a typical Se nanosheet FET device with a channel thickness of 16 nm. It shows typical p-type transport behavior with a high current on/off ratio over 10\textsuperscript{6}. The presence of hydrogen and hydroxyl terminations on the nanosheets surface is considered to be the main reason for the p-type conduction of selenium FET, which has been demonstrated in Se nanowire and nanobelts.\textsuperscript{17,36} The Se nanosheet FET presents a relatively low on-state current, with the maximum value around 20 mA/mm at $V_{dd}$ = 3 V. Four-terminal field-effect devices are fabricated to evaluate the intrinsic hole mobility of Se nanosheets, as shown in Figure S7. The mobility could be extracted from the $G$ versus $V_{bg}$ curves in the $\sim$100 < $V_{bg}$ < $\sim$30 V range using the expression $\mu = I_{on}/W \times (1/C_{bg}) \times dG/dV_{bg}$ and the value of hole mobility is expected to be 0.26 cm$^2$ V$^{-1}$ s$^{-1}$ at 300 K. Compared with other p-type semiconductor materials such as tellurene and black phosphorus,\textsuperscript{35,37–39} the Se nanosheet has a low hole mobility similar to those values reported in the literature on bulk selenium or selenium nanobelts.\textsuperscript{40–47} Because of its low mobility and special band structure, it has the highest Seebeck coefficient (+1250 \textmu V K$^{-1}$) among all elements and has the potential for thermoelectric applications. It is worth noting that the Se nanosheets and the transistors exhibit a good stability in ambient conditions (Figure S8). Even after 15 days’ exposure to air, the surface morphology of the Se nanosheet almost does not change, and there is no significant degradation of device performance.

Selenium is known as an excellent material candidate for high-sensitivity optoelectronics. Herein, we also examine the optoelectronic performance of Se nanosheet phototransistors, and a schematic diagram is shown in Figure 7a. Figure 7b shows the $I_{ds}$ curves as a function of $V_{gs}$ at various illumination power densities, exhibiting an obvious gate tunability of the photocurrent response of the Se nanosheet phototransistor. The device exhibits a pronounced photoresponse even at a very low illumination power down to 0.21 mW/cm$^2$, and the on-state photocurrent could reach up to 54 nA (Figure S9a). The enhancement of conduction under illumination conditions confirms that the Se nanosheet could be used for low-noise, high-sensitivity optoelectronic applications.

Photoresponsivity ($R_p$), defined as $R_p = I_{ph}/PS$, is calculated to better evaluate the performance of the phototransistor, where $I_{ph}$ is the generated photocurrent, $P$ is the incident power, and $S$ is the effective illuminated area. $R_p$ is estimated to be 263 A/W at an illumination power of 0.21 mW/cm$^2$ with $V_{dd}$ = 3 V and $V_{bg}$ = $-80$ V. This value is 4 orders of magnitude higher than the previously reported crystalline Se films grown by physical vapor deposition (17 mA/W with a bias of $-10$ V) and among the highest values reported for 2D materials.\textsuperscript{43–47} The responsivity is linearly proportional to the power of the illumination as shown in Figure 7c, suggesting that the photocurrent is mainly determined by the photoexcited carriers.\textsuperscript{48} Time-dependent photoresponse of laser ON and OFF is also measured at room temperature (Figure 7d). The response speed is characterized by a typical rise time of 0.10 s and decay time 0.12 s for $V_{dd}$ = 3 V and $V_{bg}$ = 0 V (taking a 10–90% photocurrent change for the rise times and 90–10% for the fall times). Figure S9b shows the photoresponse after four illumination cycles. The same level of photocurrent and noise with the laser switch demonstrates a stable and repeatable photoresponse of our device.

**CONCLUSION**

In summary, high-quality 2D Se nanosheets were successfully synthesized by the physical vapor deposition method. The as-synthesized Se nanosheet has a large lateral size up to 30 \textmu m and a minimum thickness of 5 nm. The crystal structure of the 2D Se nanosheet and its growth mechanism have been studied. Back-gated FETs and four-terminal devices based on the Se nanosheet have been demonstrated, and the intrinsic carrier mobility is determined to be 0.26 cm$^2$ V$^{-1}$ s$^{-1}$ at 300 K. The 2D Se nanosheet phototransistor exhibits an excellent photoresponsivity of 263 A/W, although the response time is slow. As a class of 2D materials formed by 1D van der Waals materials, Se nanosheets would have great potential in electronic and optoelectronic applications.

**EXPERIMENTAL METHODS**

**Growth of Selenium Nanosheets.** High-purity Se power (Sigma-Aldrich, 99.99%) was placed at the center of the heating zone in a multizone furnace, with a freshly cleaned Si(111) substrate located about 20–25 cm away from the powders. During the growth process, pure Ar gas was fed with a constant flow rate of 50 sccm. The tube pressure was maintained constant at 100 mbar. The whole reaction process was carried out under a temperature of 210 °C for the source and 100 °C for the substrate, and this temperature was maintained for 60 min. After the reaction was complete, a black-colored needlelike material coated on the Si substrate could be clearly indentified.

**Raman and STEM Measurements.** Raman measurement was performed using a HORIBA LabRAM HR800 Raman spectrometer. The system is equipped with a He–Ne excitation laser of 633 nm wavelength. The system was calibrated with the Raman peak of Si at 520 cm$^{-1}$ before measurement. The incident laser was polarized along the [0001] direction of the selenium nanosheets and illuminated perpendicularly to the nanosheet surface. The polarized laser is parallel to spiral atom chains, and we denote this configuration as 0°. To avoid destroying the sample, the laser power is less than 1 mW. The HAADF-STEM was performed with a FEI Talos F200X equipped with a so-called probe corrector. This microscope was operated with an acceleration voltage of 200 kV.

**Device Fabrication and Characterization.** Selenium nanosheets were transferred onto 300 nm SiO$_2$/Si substrates using the Scotch tape method. Electron beam lithography was used to pattern electrodes, followed by electron beam evaporation of 30 nm Ni and 100 nm Au as metal contacts. The channel length $L_{ch}$ was designed to be 4 \textmu m, while the channel width $W_{ch}$ = 10 \textmu m. Four-terminal field-effect devices are also fabricated to evaluate the intrinsic hole mobility of Se nanosheets, the channel length $L_{ch}$ between two voltage probes is 8.0 \textmu m, and channel width $W_{ch}$ is 4.0 \textmu m. The devices were measured with a probe station connected to a semiconductor characterization system (4200SCS, Keithley) at room temperature. For photodetection, a 637 nm laser source (S1FC637, Perot Benchtop) calibrated by an UV-enhanced silicon photodiode was used to provide power-tunable irradiation.
ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.7b04786.

Schematic diagram of the transfer procedure, XPS, EDX, and EELS spectra of Se nanosheets, SEM image of Se microspheres and nanowires, air stability of Se nanosheets, four-terminal devices, and related measurement, and photoelectric properties of Se nanosheets (PDF)

AUTHOR INFORMATION

Corresponding Author
*E-mail: yep@purdue.edu.

ORCID
Cheng-Yan Xu: 0000-0002-7835-6635
Haian Wang: 0000-0002-7397-1209
Peide D. Ye: 0000-0001-8466-9745

Author Contributions
J. Qin and G. Qiu contributed equally to this work.

Author Contributions
P.D.Y. conceived the idea and supervised the experiments. J.K.Q. performed the growth experiments and analyzed the growth mechanism. H.Z. performed the fabrication and analyzed the experimental data. C.Y.X. and J.K.Q. performed device measurements. J.K.Q. is supported by the Chinese Scholarship Council. The work is in part supported by the AFOSR/NSF.

ACKNOWLEDGMENTS

The authors would like to thank Chao Wang at Harbin Institute of Technology for the discussion on Raman measurements. J.K.Q. is supported by the Chinese Scholarship Council. The work is in part supported by the AFOSR/NSF EFRI 2DARE program, ARO, and SRC.

REFERENCES

(26) Lucovsky, G.; Mooradian, A.; Taylor, W.; Wright, G.; Keezer, R. Identification of the Fundamental Vibrational Modes of Trigonal, α-