

Hybrid dual-channel phototransistor based on 1D t-Se and 2D ReS₂ mixed-dimensional heterostructures

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ABSTRACT

The combination of mixed-dimensional semiconducting materials can provide additional freedom to construct integrated nanoscale electronic and optoelectronic devices with diverse functionalities. In this work, we report a high-performance dual-channel phototransistor based on one-dimensional (1D)/two-dimensional (2D) trigonal selenium (t-Se)/ReS₂ heterostructures grown by chemical vapor deposition. The injection and separation efficiency of photogenerated electron–hole pairs can be greatly improved due to the high-quality interfacial contact between t-Se nanobelts and ReS₂ films. Compared with bare ReS₂ film devices, the dual-channel phototransistor based on t-Se/ReS₂ heterostructure exhibits considerable enhancement with the responsivity (R) and detectivity (D^*) up to 98 A·W⁻¹ and 6 × 10¹⁰ Jones at 400 nm illumination with an intensity of 1.7 mW·cm⁻², respectively. Besides, the response time can also be reduced by three times of magnitude to less than 50 ms due to the type-II band alignment at the interface. This study opens up a promising avenue for high-performance photodetectors by constructing mixed-dimensional heterostructures.

KEYWORDS

van der Waals heterostructures, ReS₂, trigonal selenium (t-Se) nanobelt, phototransistor

1 Introduction

The van der Waals (vdW) heterostructures based on combination of two-dimensional (2D) layered materials exhibit unique physical properties such as gate-tunability [1, 2], fast charge separation and strong interlayer coupling effect [3–6]. Besides, a number of nonlayered materials with different dimensionalities can be also integrated with 2D layered materials by vdW forces, thereby creating mixed-dimensional vdW heterostructures [7–10]. For example, quadrate PbS nanoplates could be grown on the edge of monolayer MoS₂ to form a hybrid structure, and the device based on the structure exhibits a high photoresponsivity up to 4.5 × 10⁴ A·W⁻¹ and fast response time less than 7.8 ms [11]. The photodetector based on CdS/MoS₂ heterostructure shows broader wavelength response and 50-fold improvement in photoresponsivity compared with bare MoS₂ devices [12]. Such 2D+nD ($n = 0, 1$ and 3) combinations have greatly broaden the applications of 2D layered materials in optoelectronic [6, 13, 14], electrochemical [11, 15, 16] and biochemical fields [17–19].

Trigonal selenium (t-Se), as typical one-dimensional (1D) elemental semiconducting material, has a lot of interesting physical properties such as high photoconductivity [20–22], high piezoelectricity [23], and nonlinear optical responses [24, 25]. It possesses a highly asymmetric crystallographic structure, where Se atoms are covalently bonded into helical chains along [001] crystallographic direction, while neighboring chains are stacked together by vdW force to form a hexagonal structure. Specially, the suitable bandgap of 1.7 eV and

low dark current makes t-Se widely applied in the UV-to-visible light detect [26]. Thus, it would be of great interest if t-Se could be combined into layered materials to modulate their optoelectronic properties. However, it is still a challenge for t-Se nanostructures to be integrated with 2D layers due to the large difference of lattice parameters [27, 28]. As a unique member of 2D transition metal dichalcogenides (TMDCs), ReS₂ has proven to be highly sensitive to applied light illumination [29]. In addition, it possesses unique distorted 1T' structure (Fig. S1 in the Electronic Supplementary Material (ESM)), which would introduce abundant Re atoms protruding out of the in-plane direction [30]. These protruding Re atoms could act as binding sites for S atoms absorption and result into the vertical growth of ReS₂ by chemical vapor deposition [30–32]. Thus, we expect that such dangling-bond-abundant surface would make ReS₂ a promising platform for subsequent heterogeneous growth of t-Se nanostructures. Very recently, we demonstrated that t-Se nanoplates could be epitaxially grown on monolayer ReS₂ films to form a vertical heterostructure, which exhibits ultrahigh detectivity of up to 8 × 10¹² Jones and fast response time of less than 10 ms [33]. However, the isolated small Se nanoplates cannot form a continuous conduction channel for carrier transport, which greatly limits its further applications.

In this work, we demonstrated that t-Se nanobelts can be successfully grown on continuous few-layer ReS₂ films, thereby creating a novel 1D/2D mixed-dimensional heterostructures for photodetection applications. The t-Se nanobelts grow following a wire-by-wire self-assembly mode due to the low surface diffusion energy of ReS₂.

Dual-channel field-effect transistors (FETs) were fabricated based on such heterostructure, where both Se nanobelts and ReS₂ could act as transport channel. The mixed-dimensional device exhibits a significant enhancement of photoresponse compared with ReS₂-based devices. In addition, due to the efficient injection and separation of photo-excited electron–hole pairs at interface, the response time of the device is greatly reduced by three orders of magnitude to less than 45 ms for rising time and 15 ms for decay time.

2 Experimental

2.1 Growth of ReS₂ and Se/ReS₂ heterostructures

ReS₂ films was grown on mica substrate by chemical vapor deposition (CVD) method. Al₂O₃ boat containing ReO₃ powders was put in the center of the quartz tube, and freshly cleaved fluorophlogopite mica was placed about 1 cm away along the downstream direction. Sulfur powder was placed in another Al₂O₃ boat located upstream direction. The furnace was heated to 600 °C and maintained for 10 min with 40 sccm of Ar. Two types of ReS₂ films were used as substrates for subsequent growth of t-Se nanobelts: few-layer ReS₂ with thickness of ~ 3.1 nm and monolayer ReS₂ (Fig. S2 in the ESM). Figure S3 in the ESM schematically illustrates the two-step growth process of Se/ReS₂ heterostructures. Mica substrate with as-grown ReS₂ films was loaded into a quartz ampoule with Se powder on the other side. The ampoule was vacuumed until pressure lower than 10⁻³ Torr, and then heated at 300 °C for 30 min.

2.2 Characterization

The products were characterized by optical microscopy (Lecia DM4500P), scanning electron microscope (SEM, Hitachi SU-800), atomic force microscope (AFM, Bruker Dimension ICON-PT) and transmission electron microscope (TEM, FEI Talos F200x, acceleration voltage: 200 kV). Raman spectra were recorded using Raman spectrometer (Horiba HR800) with a 633 nm laser. The system was calibrated using Raman peak of Si at 521 cm⁻¹.

2.3 Device fabrication

After the Se/ReS₂ hybrid films were transferred onto SiO₂/Si substrate, the selected overlapped area was first patterned into squares with length of 15 μm by etching with O₂ plasma combined with photolithography process. Then, Ti/Au electrodes with thicknesses of 30/100 nm were fabricated by standard electron beam lithography (EBL) and high-vacuum e-beam evaporation. The width and length of the transport channel were designed to be 10 and 3 μm, respectively. Both the ReS₂ films and Se nanobelts could be contacted with electrodes to form a dual-channel structure.

2.4 Electrical and optoelectrical measurements

Electrical and optoelectrical measurements of the fabricated devices were implemented in a semiconductor instrument (Keithley 4200 SCS) accompanied with a Lakeshore probe station under room temperature and atmospheric pressure. Laser source with different wavelengths calibrated by UV-enhanced silicon photodiode was taken to provide power-tunable illumination. The illumination power to phototransistors was calibrated using power meter (Thorlabs PM100A).

3 Results and discussion

The Se/ReS₂ heterostructure was first characterized by optical microscopy, as shown in Fig. 1(a). Obviously, t-Se nanobelts with gold color could be distinguished from the distinct difference in contrast. According to the AFM topography image (Fig. 1(b)), spindle-like t-Se nanobelts with diameter up to 2 μm are assembled

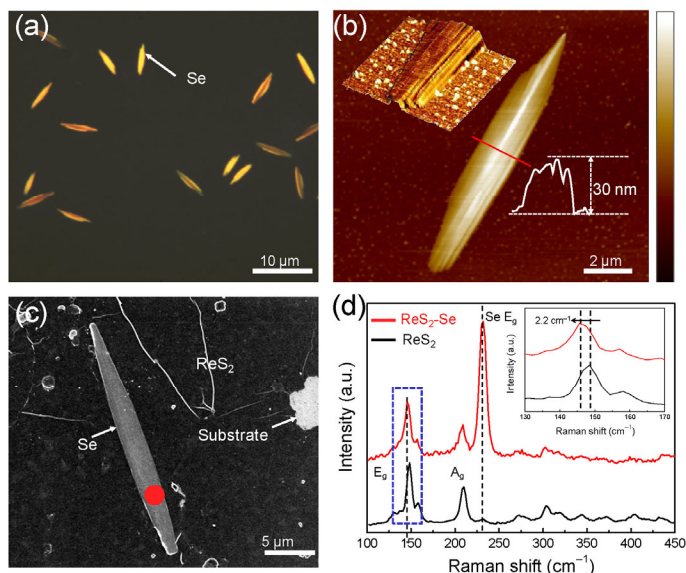


Figure 1 Characterization of Se/ReS₂ mixed-dimensional heterostructures. (a) Optical microscope image of spindle-like t-Se nanobelts on ReS₂ films. (b) AFM height profile of t-Se nanobelts. Inset shows the enlarged three-dimensional (3D) view. (c) SEM image of Se/ReS₂ heterostructure. (d) Comparison of Raman spectra obtained from Se/ReS₂ heterostructure, where the red line is collected from Se/ReS₂ hybrid area (red circle in (c)), and blue line from the bare ReS₂ film.

by parallel narrow nanowires, and they have average height of ~ 30 nm around the center with the length of up to 10 μm. Se nanobelts showing similar morphology could also be grown on monolayer ReS₂ films (Fig. S4 in the ESM). However, these t-Se nanobelts have smaller lateral size (average width < 500 nm) and larger thickness (> 100 nm) compared with that grown on few-layer ReS₂. It is expected to arise from the higher atomic diffusion energy of monolayer ReS₂ surface without defects, which will be discussed later. In the vapor deposition process, the substrate temperature plays an important role in the morphology of the products, which is decided by the location of the substrate in the quartz ampoule [34–36]. In this work, the optimum growth temperature was determined to be 180 °C for growing t-Se nanobelts. Higher deposition temperature would result into the thick t-Se bulks, while lower one gave less dense and small t-Se nanobelts as shown in Fig. S5 in the ESM.

Figure 1(d) displays the Raman spectra of Se/ReS₂ heterostructure and ReS₂ with 633 nm laser excitation at room temperature. Two dominant peaks of ReS₂ located at 150 and 213 cm⁻¹ could be assigned to in-plane E_g and out-of-plane A_g vibration modes, respectively. Remarkably, the active Raman E_g mode of t-Se located at 233 cm⁻¹, could also be clearly identified in the Raman spectrum collected from the Se/ReS₂ hybrid area (marked with the red spot in Fig. 1(c)) [21]. It should be mentioned that the E_g mode of ReS₂ in the Se/ReS₂ hybrid films exhibits distinct shift of 2.2 cm⁻¹ towards low frequency (inset in Fig. 1(d)). Compared with ReS₂, the conductance level of p-type t-Se located at a higher position, and the efficient electron (hole) transfer between them would lead to the reduction of carrier density in ReS₂, resulting into the weaken of electron–phonon coupling effect and Raman shift in ReS₂ after t-Se growth [36–39].

Hybrid Se/ReS₂ films is transferred onto Cu grid for TEM characterization. The t-Se nanobelt exhibits darker contrast compared with ReS₂ membrane, showing a typical spindle-like structure around the ending area (Fig. 2(a)). High-resolution TEM (HRTEM) image was obtained at the edge of Se/ReS₂ heterostructure marked with red rectangular in Fig. 2(a), where distinct lattice fringes belonging to t-Se can be clearly identified. By combining fast Fourier transform (FFT) pattern with HRTEM (Fig. 2(c)), we can conclude that the single-crystalline t-Se nanobelts are grown along the [001] crystallographic direction. Indeed, the few-layer ReS₂ films are

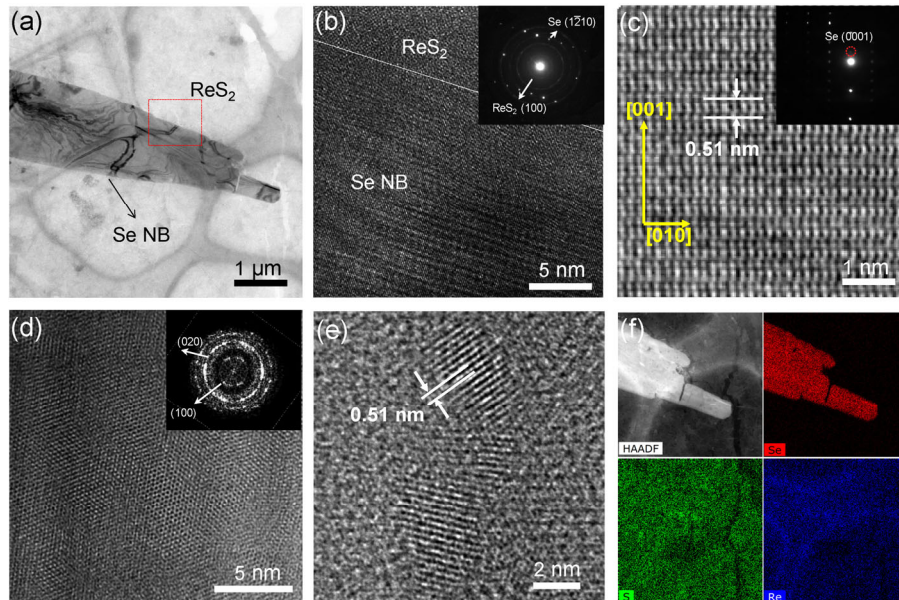


Figure 2 TEM characterization of Se/ReS₂ heterostructures. (a) Bright-field TEM image of Se/ReS₂ heterostructures. (b) HRTEM image of the Se/ReS₂ heterostructures at the edge area (marked with red rectangle in (a)). (c) HAADF-STEM image of t-Se nanobelts. Inset image shows the SAED pattern. (d) HRTEM image of few-layer ReS₂ films. Inset image shows the corresponding FET pattern. (e) HRTEM image of t-Se NCs in the initial stage of growth process. (f) EDS elemental mapping of the heterostructure at the ending area.

polycrystal stacked with random-orientated small flakes, which can be demonstrated by the dim diffraction ring as the FFT pattern shows (Fig. 2(d)). It should be noted that highly single-crystallized zero-dimensional (0D) t-Se nanocrystals (NCs) with average diameter of 2 nm were also observed in HRTEM image (Fig. 2(e)). The lattice spacing is estimated to be 0.51 nm, which can be assigned to (001) planes of t-Se. These 0D t-Se nanocrystals are expected to be the products at the initial stage during growth process. Chemical compositions of such hybrid structure are also confirmed by energy dispersive spectrometer (EDS) as shown in Fig. 2(f).

Great interfacial contact of heterojunction would facilitate the energy transfer process between different materials, which would help us to modulate its physical properties more effectively [8, 36, 40]. In this work, AFM scratching method was introduced to evaluate the contact quality of Se nanobelt and ReS₂ films as illustrated in Fig. S6 in the ESM. Remarkably, after t-Se nanobelt being scratched away, the underlying ReS₂ films could be totally lifted off, exposing bare mica substrate. The depth around the pit is about 3.2 nm, which is well in accordance with the thickness of few-layer ReS₂ films, and the corresponding phase image exhibits distinct contrast difference in the exposing area (Figs. S6(c) and S6(d) in the ESM). Besides, Re and S elements can be clearly detected from the EDS mapping image collected from scratched t-Se nanobelts (Fig. S7 in the ESM), indicating that the ReS₂ films are tightly attached on the bottom of t-Se nanobelts. These results strongly demonstrated that the above t-Se nanobelts could contact

tightly with underlying ReS₂ films, forming good heterointerface connection. The scratching technique was also applied on t-Se nanobelts grown on monolayer ReS₂ films. Similarly, the underneath ReS₂ is totally lifted off after AFM scratching (Fig. S8 in the ESM).

The ideal interface contact between ReS₂ and Se could be attributed to abundant intrinsic binding sites on ReS₂ surface. As we mentioned before, ReS₂ is a distinct 2D layered material with distorted 1T' structure, and the ultra-low symmetry would introduce lots of protruding Re atoms out of the ReS₂ (001) planes [30]. These unbounded Re atoms with high chemical activity would act seeds for subsequent Se growth, resulting into the covalent bonding of Re and Se atoms at the interface. Besides, these intrinsic protruding Re atoms could also lower the energy barrier for atomic nucleation and migration and facilitate the in-plane growth of t-Se nanobelts. A possible growth mechanism was proposed to explain such nucleation mode as shown in Fig. 3. Due to the highly anisotropic crystal structure of t-Se, and it tends to grow along [001] direction into 1D nanowire [27]. At the first stage, Se atoms would be absorbed on ReS₂ surface and chemically bonded to the protruding Re atoms, resulting into the small clusters (Fig. 3(e)). Due to the low atomic migration energy barrier, these small t-Se clusters adjacent to the ReS₂ surface are likely expanded laterally along in-plane direction into thin nanobelts, while the vertical growth can be partly inhibited due to the higher migration barrier introduced by vdW interaction between Se helical chains [41, 42]. It is worth noting that few-layer ReS₂ films could provide much more active sites compared with

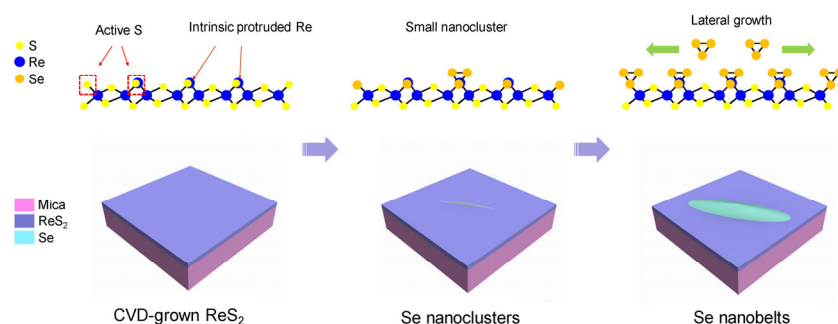


Figure 3 Schematic illustration of the growth mechanism of Se/ReS₂ heterostructures.

monolayer ReS₂ due to the smaller grain size and larger edge-to-area ratio, resulting in the formation of t-Se nanobelts with relatively small thickness and large width.

FETs based on Se/ReS₂ heterostructure were fabricated using Ti/Au as contact on SiO₂/Si substrate. Figure 4(a) shows the optical microscopy image of the device. Both of the ReS₂ films and t-Se nanobelt could contact with device electrodes, thus forming a back-gate transistor with dual conductive channels. The thickness of t-Se nanobelt and few-layer ReS₂ films are about 41 and 3.1 nm, respectively. FETs based on ReS₂ films were also fabricated for comparison. As shown in Fig. S9 in the ESM, the transistor based on ReS₂ films shows distinct n-type transport behavior with high on/off ratio of 10⁵, while the hybrid Se/ReS₂ device exhibits a degenerate n-type transport characteristic (Figs. 4(b) and 4(c)). The off-state current in Se/ReS₂ hybrid transistor is increased by three orders of magnitude as negative gate voltage (V_g) is applied, suggesting the transport behavior is dominated by the hole carriers of p-type t-Se nanobelts. On the contrary, the hybrid device exhibits a small on-state current of 3 nA at $V_g = 80$ V with source-drain voltage (V_{ds}) of 5 V, which is only one-tenth of that in bare ReS₂ devices, indicating that the vertical stacked t-Se nanobelt has a distinct p-doping effect on the underlying ReS₂ films. The carrier mobility could be extracted from the transfer curve using the expression $\mu = L/W \times (1/C_g) \times dG/dV_g$, and it is estimated to be 0.19 cm²·V⁻¹·s⁻¹ for the hybrid transistor. This value is much smaller than that of bare ReS₂ devices (about 3.42 cm²·V⁻¹·s⁻¹ at 300 K). The electron (hole) would transfer between ReS₂ and t-Se nanobelt due to the difference of Fermi level [39, 43], which would cause the reduction of carrier concentration and lead to a depletion region at interface. Thus, the on-state current could be considerably reduced under dark condition. The small on-state dark current could greatly decrease the measuring noise and improve

the detectivity of phototransistors [44].

The conduction level (valence level) of trigonal Se and ReS₂ are estimated to be -3.20 eV (-4.92 eV) and -4.25 eV (-6.05 eV), respectively. This difference would lead to the formation of type-II band structure, resulting into the efficient separation of photo-generated electron-hole pairs in Se/ReS₂ heterostructures [39, 43]. The optoelectronic property of Se/ReS₂ mixed-dimensional heterostructure was also investigated using the same devices. Figure 5(a) presents the schematic structure of the hybrid Se/ReS₂ phototransistor, where the above t-Se nanobelts could act as light absorber to enhance the photoresponse of the devices. As shown in Fig. 5(b), by combining the t-Se nanobelts with ReS₂, the photocurrent in Se/ReS₂ device was enhanced by more than 5 times under zero bias voltage as exposed to 400 nm laser illumination. The source-drain current is greatly increased in the whole range of V_g , and it gradually rises up with the increase of power intensity from 1.7 to 7.4 mW·cm⁻² (Fig. 5(c)). Figure 5(d) shows the on/off ratio of the phototransistor under dark and light condition at different bias voltages. Compared with the ReS₂ device, the on/off ratio of Se/ReS₂ device is improved by an order of magnitude and it could remain at 200 as V_g scanning from negative to positive, suggesting good stability of Se/ReS₂ device under gate voltage. The photoresponsivity (R) and photodetectivity (D^*) are two important parameters to evaluate the performance of photodetectors. R could be calculated according to the equation $R = I_{ph}/P_{light}$, where I_{ph} is the generated photocurrent and P_{light} represents the incident power intensity. D^* could be obtained using following equation: $D^* = (RA^{1/2})/(2eI_d)^{1/2}$ where A is the effective area of the detector, e is the value of electron charge, and I_d represents the dark current. The photoresponsivity of Se/ReS₂ hybrid phototransistor is estimated to be 98 A·W⁻¹ at $V_g = 60$ V, which is one order higher than that of ReS₂ device ($R = 8.7$ A·W⁻¹). Besides, the D^* of Se/ReS₂ device is calculated

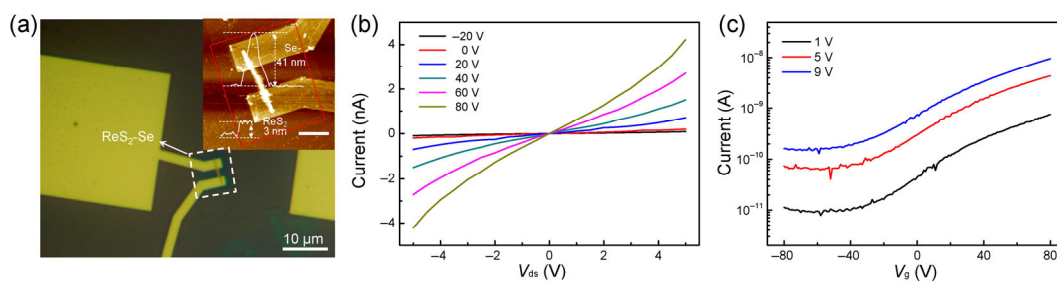


Figure 4 Electrical properties of Se/ReS₂ dual-channel FETs. (a) Optical image of the device. Inset shows the corresponding AFM height profile, and ReS₂ films are marked with red dashed line. Scale bar is 5 μ m. (b) Output curves and (c) transfer curves of the device based on Se/ReS₂ heterostructures.

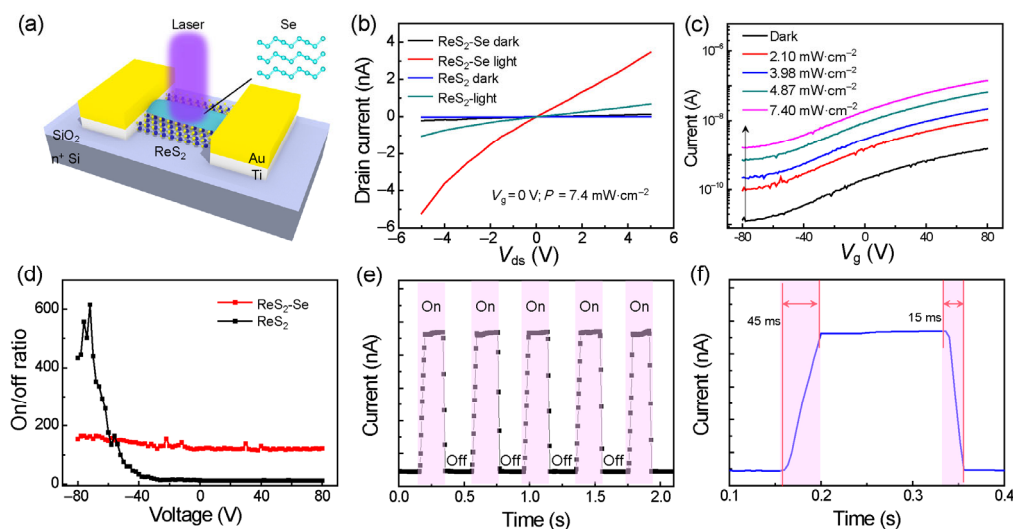


Figure 5 Optoelectronic performance of the Se/ReS₂ phototransistor. (a) Schematic of Se/ReS₂ device. (b) Output curves of ReS₂ and Se/ReS₂ phototransistors. (c) Transfer curves of the device under 400 nm irradiation with different power densities. (d) Bias dependence of the on/off ratios on the two samples. (e) Time-dependent photocurrent measured under chopped light illumination in the hybrid device. (f) A separated response and reset cycle of hybrid devices.

to be 6×10^{10} Jones under the same measuring condition, which is comparable with previous reported MoS₂-PbS and WS₂-Bi₂Te₃ hybrid films [40, 45].

The response speed is another critical parameter for photodetectors. Remarkably, the photoresponse time of Se/ReS₂ hybrid device can be significantly reduced as demonstrated by the time-dependent photocurrent ($I_{ph}-t$) curves in Fig. 5(f). The response and recovery time are estimated to be 45 and 15 ms, respectively, which are almost three orders of magnitude smaller than that of ReS₂ device (15 and 18 s), as shown in Fig. S9 in the ESM. The hybrid device also exhibits an excellent stability without degeneration even after five cycles of alternative illumination (Fig. 5(e)). This excellent response property provides Se/ReS₂ heterostructure a promising application to monitor fast optical signals, and it is also superior than previous reported ReS₂ based phototransistors [13, 32]. Table S1 in the ESM summarizes the important parameters of reported photodetectors based on related mixed-dimensional heterostructures. Obviously, the devices in our work have very high detectivity, great photoresponsivity and relatively fast response.

The enhancement of photoresponse in Se/ReS₂ hybrid device could be well explained by the efficient charge transfer at interface. As mentioned before, both conduction and valence levels of t-Se are located at higher position compared with ReS₂, which would result into a type II band alignment at contact (Fig. 6(b)). As illustrated in Fig. 6(c), the electrons (holes) transfer between t-Se and ReS₂ would cause the depletion of carriers and introduce a built-in field toward ReS₂. Thus, the on-state current could be greatly suppressed under dark condition [7]. Once the light is applied, the photo-generated electrons in t-Se nanobelts can be immediately separated and transferred to ReS₂ driven by the built-in electric field, and photo-excited holes in underlying ReS₂ could also transfer to t-Se nanobelts, which would cause the rising (lowering) of Fermi level in ReS₂ (t-Se nanobelts). The accumulated electrons (holes) in ReS₂ (t-Se nanobelts) could transport through these two channels driven applied electric field of V_{ds} and be captured immediately by the electrodes, resulting in the increase of photocurrent. On the other hand, the modulation of Fermi level would reduce the Schottky barrier at Ti/ReS₂ (Ti/t-Se) interface, which would also facilitate the carrier injection and increase the channel current [8, 10].

As mentioned before, the protruding Re atoms could easily bond with gas molecules from ambient experiment due to the high chemical activity, which would introduce trap states in the band gap and severely deteriorate the response speed [30, 46]. Upon the growth of Se nanobelts, the chemically bonded Se atoms could effectively

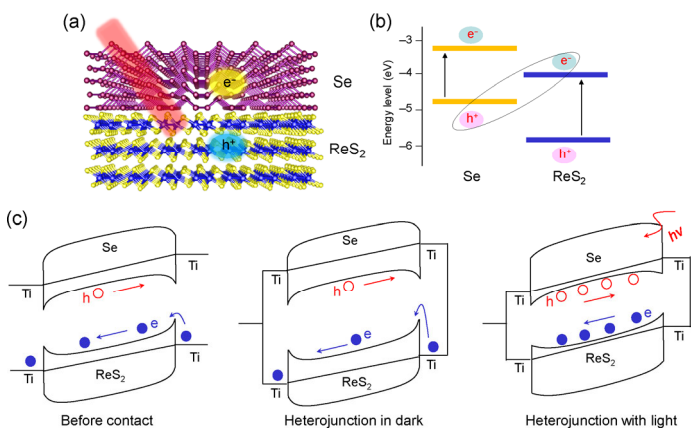


Figure 6 The injection and separation of photo-excited electron–hole pairs at Se/ReS₂ interface. (a) Schematic description for the carrier separation and transport in Se/ReS₂ hybrid structure under light illumination. (b) Band structures of the Se/ReS₂ heterostructure. (c) Schematic diagram of the energy band and carrier transport in Se/ReS₂ heterostructure under dark and light condition.

passivate the surface defects in ReS₂, and the Re–Se bonding at interface can provide a fast transition path for carriers' separation and transfer, leading to the acceleration of response speed. Besides, the potential barrier induced by built-in field at interface would also in turn hinder the carriers transfer back as the light tuned off [9]. Thus, the photo-excited electrons (holes) in ReS₂ (t-Se) could be rapidly recombined, leading to the reduction of recovery time.

4 Conclusions

In summary, we reported a vapor disposition approach to obtain Se/ReS₂ mixed-dimensional heterostructures. Due to low surface diffusion energy, t-Se nanobelts could be easily assembled wire by wire, thereby creating a fascinating 1D–2D heterostructure. The high-quality interface contact between t-Se nanobelts and underlying ReS₂ films could efficiently facilitate the charge transfer and separation at interface. The dual-channel phototransistor based on Se/ReS₂ heterostructure exhibits a considerable enhancement in photoresponsivity and photodetectivity compared with the values obtained from bare ReS₂ device under identical illumination. The response time could be considerably reduced by 3 orders of magnitude to less than 50 ms. These findings open up new opportunities to improve the optoelectronic performance of photodetectors by fabricating mixed-dimensional heterostructures.

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