Doping-Free Arsenene Heterostructure Metal-Oxide-Semiconductor Field Effect Transistors Enabled by Thickness Modulated Semiconductor to Metal Transition in Arsenene

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Two-dimensional (2-D) materials such as MoS$_2$ and phosphorene provide an ideal platform to realize extremely thin body metal-oxide-semiconductor field effect transistors (MOSFETs) which is highly immune to short channel effects in the ultra-scaled regime. Even with the excellent electrostatic integrity inherent in 2-D system, however, 2-D materials suffer from the lack of efficient doping method which is crucial in MOSFETs technology. Recently, an unusual phase transition from semiconductor to metal driven by the thickness modulation has been predicted in mono-elemental 2-D material arsenene. Utilizing this extraordinary property, we propose doping-free arsenene heterostructure MOSFETs based on the lateral multilayer (metallic source)/monolayer (semiconducting channel)/multilayer (metallic drain) arsenene heterostructure. Metallic multilayer arsenene in the source and drain can serve as electrodes without doping. We investigate the potential performance of arsenene heterostructure MOSFETs through atomistic simulations using density functional theory and nonequilibrium Green's function. The intrinsic upper limit of the on-state current in arsenene heterostructure MOSFETs is estimated by studying the effect of layer number in the source and drain. We comprehensively analyze the competitiveness of arsenene heterostructure MOSFETs through benchmarking with monolayer arsenene homostructure MOSFETs equipped with the highly degenerate doped source and drain, suggesting superior performance of heterostructure MOSFETs over homostructure MOSFETs.

Over the past few years, gapped two-dimensional (2-D) materials represented by MoS$_2$ and phosphorene have been widely explored for the future device applications. Such 2-D materials are attractive candidates especially for the channel materials in the aggressively scaled metal-oxide-semiconductor field effect transistors (MOSFETs) since their extreme thinness allows almost ideal electrostatic control over the channel, making them robust to short channel effects. However, one of the critical issues hindering 2-D materials from the adoption in the MOSFETs technology is the challenge to dope 2-D materials effectively to n- or p-type, which lies in the heart of MOSFETs technology. In the conventional silicon MOSFETs, highly doped source and drain electrodes are achieved by the substitutional impurity atoms. However, applying such a technique to 2-D material MOSFETs is challenging because of the atomically thin nature of 2-D materials. In search for an alternative doping strategy, charge transfer doping using deposition of molecules or adatoms on 2-D materials has been suggested by theoretical calculations and experimentally demonstrated. Such doping methods using surface modification by molecules or adatoms can locally alter the conductivity of 2-D materials but the long-term stability in the ambient atmosphere still remains as an issue.

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Recently, new mono-elemental 2-D materials, namely arsenene and antimonene made of As and Sb, respectively, have emerged as promising materials\textsuperscript{13–17}. It is theoretically predicted that monolayer of arsenene and antimonene has a sizable band gap\textsuperscript{13–15} and comparable or superior mobility than MoS\textsubscript{2} and phosphorene\textsuperscript{17}. \textit{Ab-initio} quantum transport simulations reveal that monolayer of arsenene and antimonene are competitive channel materials for the deeply scaled MOSFETs\textsuperscript{17,18}. In experiments, a few layers of arsenene and antimonene have been successfully synthesized on various substrates\textsuperscript{19–22}. Moreover, even with the concern about the oxidation of antimonene under the ambient condition predicted by the \textit{ab-initio} calculation\textsuperscript{23}, high stability upon the exposure to air and water has been experimentally proven\textsuperscript{20–22,24}. Other than the excellent mobility and stability, the most unique property of arsenene and antimonene is the abrupt switching of electronic properties from semiconducting to metallic depending on the thickness of layers. Monolayer of arsenene and antimonene has a band gap larger than 1 eV while the band gap is entirely closed in multilayer due to the strong interaction between different layers\textsuperscript{13,16}. This thickness modulated phase transition which has never been observed in the other materials before can provide revolutionary solutions to the critical challenges such as the low on-state current issue in tunneling field-effect transistors (TFETs) as discussed in our previous work\textsuperscript{25}. In the present work, we focus on the possibility to utilize this property to resolve the doping issue in 2-D materials MOSFETs. We can consider using multilayer for the source and drain electrodes and semiconducting monolayer for the channel in MOSFETs. Then, this heterostructure MOSFETs will not require any doping processes to form electrodes since multilayer itself is already highly conductive. Similar concept of device has been demonstrated in the other 2-D material MOSFETs based on MoS\textsubscript{2} which also exhibits the phase transition triggered by a crystal structure change from trigonal prismatic \textit{2H} to octahedral \textit{1T}\textsuperscript{26}. Ref.\textsuperscript{26} reports substantial performance improvement in MoS\textsubscript{2} MOSFETs using metallic \textit{1T} phase MoS\textsubscript{2} for the source and drain regions. However, the stability of \textit{1T} phase electrodes under the high-performance device operation is still unknown\textsuperscript{26} and forming local \textit{1T} phase MoS\textsubscript{2} in the source and drain needs an additional chemical process. For arsenene and antimonene, the metallic nature of multilayer is energetically stable since it originates from the interlayer coupling and the multilayer/multilayer heterostucture can be obtained by selective etching, a well-known approach to control the nanostructure thickness. Atomic layer etching\textsuperscript{27–29}, recently demonstrated for other 2-D materials such as graphene\textsuperscript{26} and MoS\textsubscript{2}\textsuperscript{28,29}, may be applicable to arsenene and antimonene to control the layer number accurately.

Herein, potential performance of doping-free arsenene heterostructure MOSFETs is investigated via density functional theory (DFT) and nonequilibrium Green’s function (NEGF) calculations. We focus on only arsenene since band structures of arsenene and antimonene are quite alike and hence almost the same level of device performances is expected as discussed in monolayer arsenene and antimonene MOSFETs\textsuperscript{37}.

Results and Discussion

We first study band structures of monolayer and multilayer arsenene with the optimized geometric structures. As in Fig. 1(a), monolayer arsenene is in the buckled hexagonal honeycomb lattice with the optimized in-plane lattice constant \(a = 3.553 \text{ Å}\) and the buckling height between two sublattices \(d = 1.428 \text{ Å}\) in good agreement with the previous studies\textsuperscript{15,17,30–33}. In the left of Fig. 1(b), band structures for monolayer arsenene is plotted along the high symmetric points \(\Gamma\)–M–K in the hexagonal \textit{1st} Brillouin Zone (BZ). Similar to the other DFT calculations\textsuperscript{17,30–33}, we observe conduction band minimum (CBM) occurring along the line between \(\Gamma\) and \(M\) points and valence band maximum (VBM) at \(\Gamma\) point with the indirect band gap \(E_{\text{g}} = 1.516\text{ eV}\). There are several possible choices for exchange correlation potentials and pseudopotentials in the Perdew–Burke–Ernzerhof (PBE) level which yield a somewhat smaller or larger band gap than 1.516 eV as seen in Fig. S1. We consider the combination of exchange correlation potential and pseudopotential resulting in the band gap value 1.516 eV since it is roughly the average. Our band gap value is also close to the previously reported ones (1.48\textsuperscript{27}, 1.50\textsuperscript{90}, 1.51\textsuperscript{31}, 1.6\textsuperscript{25}, 1.635\textsuperscript{33} and 1.76\textsuperscript{15} eV) by PBE functionals while smaller compared with the ones (2.0\textsuperscript{32}, 2.2\textsuperscript{31} eV) by HSE functionals. Typically, DFT with PBE functionals underestimate the band gap. For several other 2-D materials including MoS\textsubscript{2}, however, band gaps predicted by PBE functionals agree well with the experimental measurements\textsuperscript{34}. Moreover, a significant band gap reduction by increasing temperature from 0 K to room temperature in the ambient condition is predicted by the DFT calculation using HSE functionals for monolayer antimonene which shares the similar crystal structure and band structures with monolayer arsenene\textsuperscript{35}. Therefore, since we explore device performances at room temperature, our band gap value 1.516 eV is valid for the following device simulations. As shown in the energy contour plot for the lowest CB in the right of Fig. 1(b), we observe six-fold degenerate CB valleys with highly anisotropic effective masses. A longitudinal effective mass from \(\Gamma\) to \(M\) and a corresponding transverse effective mass is estimated to \(m_{\text{c}} = 0.461 \times m_{e}\) and \(m_{\text{\perp}} = 0.174 \times m_{e}\) respectively. Such anisotropic effective masses usually lead to the strong dependency of device performances on the transport direction. For arsenene, however, we suppose a negligible effect of the transport direction owing to the degenerate CB valleys located between \(\Gamma\) and \(M\) points in the hexagonal \textit{1st} BZ. As discussed in the anisotropic 2-D material HfS\textsubscript{2}, MOSFETs\textsuperscript{36}, aligning one valley in the light effective mass direction is always accompanied by the alignment of the other valleys in the heavy effective mass direction. Therefore, current increase or decrease in one valley is compensated by current decrease or increase in the other valleys, which results in little change in total current. So, we focus on the transport only in the armchair direction. Band structures of multilayers (bilayer, trilayer, quadlayer and hexalayer) are also studied with the optimized geometry. We perform geometry optimizations for bilayer, trilayer, quadlayer and hexalayer, respectively, with the fixed in-plane lattice constant \(a = 3.553 \text{ Å}\) obtained from the monolayer calculation. As presented by the band structure of bilayer arsenene in the left of Fig. 1(c), stacking one more layer turns semiconducting monolayer into metallic bilayer by entirely closing the band gap. More bands are induced around the Fermi level in trilayer (in the middle of Fig. 1(c)), quadlayer (in the right of Fig. 1(c)) and hexalayer (not shown) arsenene.

For the device simulation, double-gate (DG) arsenene heterostructure MOSFETs with the intrinsic semiconducting monolayer as channel and the metallic multilayer as source and drain are constructed. We consider four
different multilayers (bilayer, trilayer, quadlayer and hexalayer) for the doping-free metallic source and drain which form Schottky junctions at the monolayer/multilayer heterointerfaces. DG quadlayer/monolayer/quadlayer arsenene heterostructure MOSFETs is schematically shown in Fig. 1(d). We choose the HfO2 (κ = 25) gate oxide thickness $T_{OX}$ is 3 nm. Gate length $L_G$ is ~7 nm. Four multilayers (bilayer, trilayer, quadlayer and hexalayer) are considered to construct multilayer/monolayer/multilayer heterostructures.

Figure 1. (a) Top and side views of monolayer arsenene with primitive unit cell. (b) Band structures of monolayer arsenene and energy contour plots of the lowest CB in monolayer arsenene. Effective masses along the longitudinal and transverse directions for one CB valley are indicated. (c) Band structures of bilayer, trilayer and quadlayer arsenene along the high symmetric paths in the 1st BZ. The Fermi level is set to zero. (d) Schematic view of DG quadlayer/monolayer/quadlayer arsenene heterostructure MOSFETs. Top and bottom HfO2 ($\kappa = 25$) gate oxides thickness $T_{OX}$ is 3 nm. Gate length $L_G$ is ~7 nm. Four multilayers (bilayer, trilayer, quadlayer and hexalayer) are considered to construct multilayer/monolayer/multilayer heterostructures.

Periodic boundary conditions are assumed in the device width direction ($y$-direction). We carry out the density of states (DOS) analysis on the semiconducting monolayer/metallic multilayer arsenene heterojunction to understand Schottky barrier (SB) which is critical to the device performance of our proposed MOSFETs. Figure 2(a) shows the supercell for the DOS calculation which is composed of ~6 nm monolayer and ~4 nm multilayer. In the repeated supercell, monolayer/multilayer heterojunctions are formed in the armchair direction ($x$-direction), the transport direction in device simulations, while either monolayer or multilayer is maintained in the zigzag direction ($y$-direction). In the $z$-direction perpendicular to the layer plane, a vacuum region is added on top of the layer to block any interactions between adjacent supercells. At the monolayer/multilayer junction, unsaturated arsenic atoms on the surface of multilayer edges are passivated with hydrogen atoms to prevent the emergence of dangling bond states. Figure 2(b) presents total DOS from the entire
supercell for monolayer/bilayer/multilayer (green), monolayer/trilayer/multilayer (blue), monolayer/quadlayer/multilayer (red) and monolayer/hexalayer/multilayer (violet) heterostructures, respectively. To determine the SB height, we plot the projected DOS (PDOS) only from the monolayer in the monolayer/multilayer/monolayer heterostructure together with total DOS in Fig. 2(b). The SB heights $\Phi_{SB}$ for electrons in multilayer are estimated from the relative positions of CBM with respect to the Fermi level at 0 eV as indicated by arrows in Fig. 2(b). $\Phi_{SB}$ obtained for the monolayer/bilayer heterojunction is the largest (0.808 eV) among the four heterojunctions. Upon the increase of layer number, $\Phi_{SB}$ decreases down to 0.682 eV in the monolayer/trilayer and then remains almost same in the monolayer/quadlayer and the monolayer/hexalayer. Total DOS plots represent a monotonic increase of DOS around the Fermi level by the addition of layers, which is consistent with the band structure plots in Fig. 1(d). From the DOS analysis, we may expect more current conduction by adding layers in the source and drain which reduces $\Phi_{SB}$ and introduces more DOS near the Fermi level and hence increases the tunneling probability and the number of propagating modes at the monolayer/multilayer Schottky junction.

Transfer characteristics ($I_{DS}$-$V_{GS}$) of arsenene heterostructure MOSFETs with the bilayer (green), trilayer (blue), quadlayer (red) and hexalayer (violet) source and drain, respectively, at $V_{DS} = V_{DD} = 0.7$ V are shown in Fig. 3(a) in logarithmic and linear scales. For the fair comparison, metal gate work functions of each device are adjusted to have the same off-state ($V_{gs} = V_{gs} = 0.7$ V and $V_{gs} = 0$ V) current $I_{OFF} = 0.1 \mu A/\mu m$ according to the ITRS requirements for HP devices. For all devices, irrespective of the number of layers in the source and drain, decent subthreshold slope ($SS \approx 74$ mV/dec) is observed in ~7 nm gate length devices. As $V_{GS}$ increases, transfer characteristics start to exhibit a dependency on the layer number of multilayer. At the on-state ($V_{DS} = V_{gs} = V_{DD} = 0.7$ V), heterostructure MOSFETs with the bilayer source and drain offers the smallest on-state current $I_{ON} \approx 297 \mu A/\mu m$.

With the increase of arsenene layer number, $I_{ON}$ continuously improves up to ~642 and ~903 $\mu A/\mu m$ with trilayer and quadlayer, respectively, which is consistent with our expectation from the previous DOS analysis. However, further $I_{ON}$ enhancement is not achievable by stacking more than four layers in the source and drain. Almost the same level of $I_{ON} \approx 882 \mu A/\mu m$ is obtained with the hexalayer source and drain, which suggests an upper limit of $I_{ON}$ in our proposed heterostructure MOSFETs at $V_{DD} = 0.7$ V.

For a deeper understanding of $I_{ON}$ dependency on the thickness of multilayer in the source and drain, we show the local DOS (LDOS) at the on-state with the corresponding energy resolved current density (black line) for each heterostructure MOSFETs in Fig. 4(a). $E_{FS}$ and $E_{FD}$ (black dashed lines) located at 0.35 and $-0.35$ eV in $y$-axis denote the Fermi levels in the source and drain, respectively. From LDOS plots in Fig. 4(a), we confirm the metallic source and drain and the semiconducting channel with a band gap $\sim 1.516$ eV. The energy range in the channel region with bright color represents the absence of states while no such range exists in the source and drain regions which are composed of metallic multilayers. Color of the metallic source and drain regions becomes darker as the layer number increases due to the states originating from the additional layers, which is also predicted from our DOS analysis in Fig. 2(b). At the source/channel junction, the existence of SB is apparent.
and the width of SB is narrow due to the large $V_{GS} = 0.7\,\text{V}$ at the on-state, thereby allowing a substantial tunneling current through SB. From bilayer to trilayer, the peak current density indicated by a white arrow significantly boosts up mainly owing to the reduced $\Phi_{SB}$ as discussed previously with DOS in Fig. 2(b). With quadlayer, even higher peak current density is achieved while no more enhancement with the hexalayer source and drain. To understand this $I_{ON}$ saturation upon the increase of layer number in the source and drain, the transmission and the transmission eigenstates at the on-state are explored for heterostructure MOSFETs with the hexalayer source and drain. We focus on the transmission at $E = 0.15\,\text{eV}$ (marked with a white star in Fig. 4(a)) since the primary current conduction is restricted within the upper half of the Fermi window ($-0.35 \sim 0.35\,\text{eV}$). Figure 4(b) shows the transverse momentum $k_y$ resolved transmission, exhibiting the maximum transmission occurring around the half of maximum and minimum values of $k_y$, respectively. The transmission eigenstate corresponding to the maximum transmission in Fig. 4(b) is visualized at different isovalues of 0.2, 0.3 and 0.4, respectively, with the atomistic configuration of the simulated device in Fig. 4(c). At the iso-value of 0.4 (in the top of Fig. 4(c)), the wave function in the source is localized mostly within the third and fourth layers. As approaching the source/channel junction, the wave function gradually moves down to the first and second layers and tunnels through SB and then eventually emerges again near the middle of channel. For the lower isovalues of 0.3 and 0.2 shown in the middle and bottom of Fig. 4(c), respectively, the wave function still resides in the bottom four layers, which suggests the limited contribution of top two layers to the conducting eigenstate. So even if more states become available within the Fermi window with the hexalayer source and drain, only states mostly localized within the bottom four layers could pass through the multilayer source/monolayer channel junction to reach the drain. Therefore, multilayer thicker than quadlayer can conduct almost the same level of current as quadlayer even with the higher DOS in the source and drain.

To evaluate the potential of arsenene heterostructure MOSFETs, it is benchmarked with monolayer arsenene homostructure MOSFETs equipped with the n-type doped source and drain. Two high doping levels of $5 \times 10^{13}$ and $1 \times 10^{14}/\text{cm}^2$ are considered for the source and drain electrodes. Fermi levels corresponding to $5 \times 10^{13}$ and $1 \times 10^{14}/\text{cm}^2$ doping levels are located slightly above and $\sim 100\,\text{meV}$ above CB edge, respectively, thereby leading to the degenerately doped monolayer arsenene. For the heterostructure MOSFETs, we focus on the quadlayer source and drain since the best $I_{ON}$ is obtained with quadlayer. For the performance assessment against the other 2-D material MOSFETs, we also simulate monolayer phosphorene homostructure MOSFETs with the doping levels of $5 \times 10^{13}$ and $1 \times 10^{14}/\text{cm}^2$. We choose phosphorene since phosphorene has been identified as one of the best candidates for 2-D material MOSFETs among the various 2-D materials due to the highly anisotropic CB and VB valleys as in Fig. S2. Transport calculations for arsenene and phosphorene homostructure MOSFETs with doped reservoirs are performed for the same device geometry ($T_{OX} = 3\,\text{nm}$ and $L_G \approx 7\,\text{nm}$) within the same simulation setting (GGA PBE functionals, FHI pseudopotential and DZP basis set) as arsenene heterostructure.

![Figure 3. Transfer characteristics of arsenene heterostructure MOSFETs with the bilayer, trilayer, quadlayer and hexalayer source and drain at $V_{DS} = 0.7\,\text{V}$ in logarithm and linear scales.](https://doi.org/10.1038/s41598-019-40675-7)
MOSFETs. The armchair direction which exhibits the light effective mass is used for phosphorene homostructure MOSFETs to maximize device performances. Figure 5 compares transfer characteristics of arsenene homostructure MOSFETs for the doping levels of $5 \times 10^{13}$ (black diamond) and $1 \times 10^{14}$/cm$^2$ (black square) and arsenene heterostructure MOSFETs (red circle). Transfer characteristics of phosphorene homostructure MOSFETs with $5 \times 10^{13}$ (grey diamond) and $1 \times 10^{14}$/cm$^2$ (grey square) doping levels are also shown. In the log scale plot, arsenene homostructure MOSFETs exhibits degraded subthreshold behaviors with $SS \approx 80$ and 92 mV/dec for the doping levels of $5 \times 10^{13}$ and $1 \times 10^{14}$/cm$^2$, respectively, as compared to the heterostructure MOSFETs ($SS \approx 74$ mV/dec). Even more or similar level of degradation in $SS$(~84 and ~88 mV/dec for $5 \times 10^{13}$ and $1 \times 10^{14}$/cm$^2$ respectively) is observed in phosphorene homostructure MOSFETs. On the other hand, linear scale transfer characteristics present significantly better performances in the $1 \times 10^{14}$/cm$^2$ doped homostructure MOSFETs in the above threshold region. $I_{ON}$ reaches up to ~3800 and ~3950 $\mu$A/μm in arsenene and phosphorene homostructure MOSFETs, respectively, more than 4 times higher $I_{ON}$ in arsenene heterostructure MOSFETs. For the lower doping level ($5 \times 10^{13}$/cm$^2$), current ramps up more rapidly than in the heterostructure MOSFETs with the increase of $V_{GS}$. However, current starts to saturate early around $V_{GS} \approx 0.45$ V due to the limited supply of electrons from the source while the heterostructure MOSFETs exhibits linearly increasing current. As a result, the heterostructure MOSFETs can eventually provide about 11% larger $I_{ON}$ than arsenene and almost comparable $I_{ON}$ to phosphorene homostructure MOSFETs, respectively, suggesting that more than a $5 \times 10^{13}$/cm$^2$ doping is required for arsenene homostructure MOSFETs to outperform arsenene heterostructure MOSFETs. The doping level of $5 \times 10^{13}$/cm$^2$ corresponds to adding one extra electron for every ~36 As atoms in monolayer arsenene, which is extremely challenging even if the conventional substitutional doping strategy becomes applicable to arsenene.

In all device simulations discussed so far, the source and drain metal contact regions are excluded. So, our simulation results are valid for the idealized intrinsic devices ignoring any effects from the metal contact. In real
devices, metal contacts impose an additional resistance which degrades the device performances. This extrinsic resistance $R_{S/D}$ can be characterized by the source and drain resistances $R_S$ and $R_D$ ($R_{S/D} = R_S + R_D$) connected in series with the source and drain, respectively. For more realistic evaluation of the device performances, we add $R_{S/D}$ in the postprocessing step to the intrinsic transfer characteristics obtained from the quantum transport simulations. For the postprocessing step, a set of $I_{DS}$-$V_{GS}$ curves at several different $V_{DS}$ is prepared. Then, new $V_{GS, ext}$ and $V_{DS, ext}$ accounting for the extrinsic metal contact effects are calculated following the equations of $V_{GS, ext} = V_{GS, in} + I_{DS}R_S$ and $V_{DS, ext} = V_{DS, in} + I_{DS}(R_S + R_D)$ where $V_{GS, in}$ and $I_{DS}$ are the original biases and current, respectively, provided by the quantum transport simulations. Once $V_{GS, ext}$ and $V_{DS, ext}$ are computed, we collect the $I_{DS}$-$V_{GS, ext}$ data corresponding to $V_{DS, ext} = 0.7$ V which yields new transfer characteristics at $V_{DS, ext} = 0.7$ V. This method has been adopted in the previous studies providing a good interpretation of experimental data. We postprocess intrinsic transfer characteristics (Fig. 5) of the homostructure MOSFETs for the different doping levels of $5 \times 10^{13}$ and $1 \times 10^{14}$/cm$^2$ and the heterostructure MOSFETs with the quadlayer source and drain and then carry out the performance benchmarking again. Different resistance values for $R_S = R_D = 200, 300, 400$, and 500 $\Omega \cdot \mu$m are considered for the homostructure MOSFETs. We use the lower resistance values for the heterostructure MOSFETs since it is reasonable to expect a smaller resistance from the metal-to-metal interface than from the metal-to-semiconductor interface. Moreover, a recent experimental study on the device in similar concept with our proposed heterostructure MOSFETs reports dramatic improvement in the contact resistance through the phase engineering. The source and drain resistances have a negligible effect on the subthreshold behaviors, but strongly influence the on-state current. With $R_S = R_D = 200$ $\Omega \cdot \mu$m in Fig. 6(a), $I_{ON}$ in the $1 \times 10^{14}$/cm$^2$ doped homostructure MOSFETs substantially decreases from its intrinsic $I_{ON}$ by more than 75% while only 3% degradation with the $5 \times 10^{12}$/cm$^2$ doping. Such low $I_{ON}$ degradation in the $5 \times 10^{13}$/cm$^2$ doped homosource MOSFETs is attributed to the early saturation of current observed in Fig. 5. The source

Figure 5. Transfer characteristics of monolayer arsenene and monolayer phosphorene homostructure MOSFETs for the two different doping levels of the source and drain and quadlayer/monolayer/quadlayer arsenene heterostructure MOSFETs.



![Figure 5](https://doi.org/10.1038/s41598-019-40675-7)
resistance reduces the effectively applied $V_{GS}$ by $I_{DS}$ from the intrinsic $V_{GS}$, which results in the significant current decrease for the $1 \times 10^{14}$/cm$^2$ doped homostructure MOSFETs exhibiting linearly increasing transfer characteristics as seen in Fig. 5. For the doping level of $5 \times 10^{13}$/cm$^2$, however, similar level of $I_{ON}$ is still maintained as long as the effective $V_{GS}$ remains above $\sim 0.45$ V due to the current saturation and hence, a limited $I_{ON}$ degradation is observed in Fig. 6(a). The heterostructure MOSFETs is relatively less affected by $R_{S/D}$ in comparison with the $1 \times 10^{14}$/cm$^2$ doped homostructure MOSFETs because the original intrinsic $I_{ON}$ is much smaller (~25%), and the reduced $R_{S/D}$ is added. However, the homostructure MOSFETs with the $1 \times 10^{14}$/cm$^2$ doping can still offer $\sim$33% larger $I_{ON}$ than the heterostructure MOSFETs. In Fig. 6(b), $R_S = R_D = 300$ and $60 \Omega \cdot \mu$m for the homostructure and heterostructure MOSFETs, respectively, degrade transfer characteristics further. The degree of degradation is the most severe in the $1 \times 10^{14}$/cm$^2$ doped homostructure MOSFETs, which makes $I_{ON}$ for all devices within the range of $\sim 10\%$ difference. With $R_S = R_D = 400$ and $80 \Omega \cdot \mu$m in Fig. 6(c), almost same transfer characteristics are predicted for all three devices, but slightly better $I_{ON}$ for the heterostructure MOSFETs ($\sim$10 and $\sim$4% larger than for the $5 \times 10^{13}$ and $1 \times 10^{14}$/cm$^2$ doped MOSFETs, respectively) is expected. The resistance value larger than 400

Figure 6. Transfer characteristics of monolayer arsenene homostructure MOSFETs for the two different doping levels of the source and drain and quadlayer/monolayer/quadlayer arsenene heterostructure MOSFETs. $R_S = R_D = \begin{align*} (a) & \ 40 \text{ and } 200 \Omega \cdot \mu \text{m}, \quad (b) \ 60 \text{ and } 300 \Omega \cdot \mu \text{m}, \quad (c) \ 80 \text{ and } 400 \Omega \cdot \mu \text{m} \quad \text{and} \quad (d) \ 100 \text{ and } 500 \Omega \cdot \mu \text{m} \end{align*}$ are included for arsenene heterostructure MOSFETs and monolayer arsenene homostructure MOSFETs, respectively.
and 80 $\Omega\mu$m enables the heterostructure MOSFETs to further outperform the homostructure MOSFETs with more than ~15% $I_{\text{ON}}$ boost as seen in Fig. 6(d).

As discussed previously, there is an uncertainty in the band gap value of monolayer arsenene originating from the different choices of exchange correlation potentials and pseudopotentials in DFT. Such uncertainty could influence device performances of arsenene heterostructure MOSFETs since the SB height $\Phi_{\text{SB}}$ is affected by the band gap size. In general, a smaller band gap (a smaller $\Phi_{\text{SB}}$) improves $I_{\text{ON}}$, but with the sacrifice of SS, while a larger band gap (a larger $\Phi_{\text{SB}}$) degrades $I_{\text{ON}}$ but leads to the better SS. Since our band gap value is roughly the average, our simulations suggest the average performance which we can expect from arsenene heterostructure MOSFETs. Additionally, we note that our ballistic simulations provide an upper limit of device performances. Inclusion of electron-phonon scattering may degrade actual device performances. However, ref.12 estimates a few tens of nanometer length for the mean free path limited by the longitudinal acoustic phonon in arsenene. Therefore, in the extremely scaled MOSFETs such as our ~7 nm gate length device, electron-phonon scattering would have a limited impact.

Conclusions

We present a rigorous analysis for the potential performance of arsenene heterostructure MOSFETs utilizing the exotic property of arsenene, the thickness modulated semiconductor to metal transition. Such an abrupt modification of electronic structures depending on the number of arsenene layers can provide a solution to the fundamental issues in the realization of high performance 2-D material MOSFETs such as the lack of doping technique and the high contact resistance. Exploiting the thickness controlled phase transition in arsenene, the heterostructure MOSFETs could be realized within the same technological platform without doping simply by constructing heterostructures consisting of different thicknesses of arsenene layers through the top-down selective etching. Through DFT combined with NEGF simulations, we assess the performance of doping-free arsenene heterostructure MOSFETs and discuss the effect of layer number in the metallic source and drain on the device performances, which suggests the upper limit of intrinsic $I_{\text{ON}}$ in the heterostructure MOSFETs. Benchmarking with the monolayer arsenene homostructure MOSFETs equipped with the n-type doped source and drain, the heterostructure MOSFETs can intrinsically offer larger $I_{\text{ON}}$ as well as better SS than the homostructure MOSFETs with the $5 \times 10^{13} \text{cm}^{-2}$ doped source and drain in the ~7 nm gate length device. Including the extrinsic metal contact resistance, the heterostructure MOSFETs can possibly surpass even more highly doped homostructure MOSFETs since the lower contact resistance is expected at the interface between contact metal and multilayer metallic arsenene.

Methods

Electronic band structures, DOS and geometry optimization calculations are carried out with DFT simulation package Atomistix ToolKit (ATK)39,44,45. We employ generalized gradient approximation (GGA) and PBE functionals to represent exchange-correlation potentials with double zeta polarized (DZP) basis set39, Troullier-Martins type norm-conserving pseudopotential set (FHI [z = 5] DZP) is adopted for arsenene. We include Grimme’s DFT-D2 empirical dispersion correction40 to PBE to account for the van der Waals interaction which is known to be important for the accurate description of such layered structures. A $k$–point sampling of $\times 7 \times 1$ for the BZ integration is used with a mesh cut-off energy of 45 Hartree. Geometry optimization is performed until the maximum residual force becomes smaller than 0.01 eV/Å.

Device characteristics are investigated through ballistic quantum transport simulations within NEGF formalism using ATK38,44,45. Same exchange-correlation functionals, pseudopotentials and basis sets adopted for band structure, DOS and the geometry optimization calculations are also used for the atomistic description of arsenene in device simulations. Since periodic boundary conditions are imposed in the device width direction ($y$-direction), we include different values of transverse momentum $k_y$ in the transport calculation. The electrostatic potential of the device for each bias condition is obtained by solving Poisson’s equation self-consistently with the quantum transport equations. After self-consistent electrostatic potential is determined, total current is calculated by summing the transmitted current over all modes with the Fermi function weight.

References

1. Radišćević, B., Radenović, A., Brivio, J., Giacometti, V. & Kis, A. Single-layer MoS$_2$ Transistors. Nat. Nanotechnol. 6(3), 147–150 (2011).
2. Fuhrer, M. S. & Hone, J. Measurement of Mobility in Dual-Gated MoS$_2$ Transistors. Nat. Nanotechnol. 8, 146–147 (2013).
3. Li, L. et al. Black Phosphorus Field-Effect Transistors. Nat. Nanotechnol. 9, 372–377 (2014).
4. Liu, H. et al. Phosphorene: An Unexplored 2D Semiconductor with a High Hole Mobility. ACS Nano 8, 4033–4041 (2014).
5. Yoon, Y., Ganapathi, K. & Salahuddin, S. How Good Can Monolayer MoS$_2$ Transistors Be? Nano Lett. 11, 3768–3773 (2011).
6. Alam, K. & Lake, R. K. Monolayer MoS$_2$ Transistors Beyond the Technology Road Map. IEEE Trans. Electron Devices 59, 3230–3235 (2012).
7. Chang, J., Register, L. F. & Banerjee, S. K. Atomicistic Full-Band Simulations of Monolayer MoS$_2$ Transistors. Appl. Phys. Lett. 103, 223509 (2013).
8. Giovannetti, G. et al. Doping Graphene with Metal Contacts. Phys. Rev. Lett. 101, 026803 (2008).
9. Chang, J., Larentis, S., Tutuc, E., Register, L. F. & Banerjee, S. K. Atomicistic simulation of the electronic states of adatoms in monolayer MoS$_2$. Appl. Phys. Lett. 104, 141603–141603 (2014).
10. Sanders, S. et al. Engineering high charge transfer n-doping of graphene electrodes and its application to organic electronics. Nanoscale 7, 13135–13142 (2015).
11. Fang, H. et al. Degenerate n-Doping of Few-Layer Transition Metal Dichalcogenides by Potassium. Nano Lett. 13, 1991–1995 (2013).
12. Liao, W. et al. Efficient and reliable surface charge transfer doping of black phosphorus via atomic layer deposited MgO toward high performance complementary circuits. Nanoscale 10, 17007–17014 (2018).
13. Zhang, S., Yan, Z., Li, Y., Chen, Z. & Zeng, H. Atomically Thin Arsenene and Antimonene: Semimetal–Semiconductor and Indirect–Direct Band-Gap Transitions. Angew. Chem. 54, 3112–3115 (2015).
14. Akturk, U. U., Ozcelik, V. O. & Ciraci, S. Single-layer crystalline phases of antimony: Antimonenes. Phys. Rev. B: Condens. Matter Mater. Phys. 91, 235446 (2015).
