Photo-Modulation of Charge Transport in all-semiconducting 2D-1D van der Waals Heterostructures with Suppressed Persistent Photoconductivity Effect

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Abstract

Van der Waals heterostructures (VDWHs), obtained via the controlled assembly of two-dimensional atomically thin crystals, exhibit unique physico-chemical properties, rendering them prototypical building blocks to explore new physics and for applications in optoelectronics. As the emerging alternatives to graphene, monolayer transition metal dichalcogenides and bottom-up synthesized graphene nanoribbons (GNRs) are promising candidates for overcoming the shortcomings of graphene, such as the absence of a bandgap in its electronic structure, which is essential in optoelectronics. Herein, VDWHs comprising GNRs onto monolayer MoS$_2$ are fabricated. Field-effect transistors (FETs) based on such VDWHs show an efficient suppression of the persistent photoconductivity typical of MoS$_2$, resulting from the interfacial charge transfer process. The MoS$_2$-GNR FETs exhibit drastically reduced hysteresis and more stable behavior in the transfer characteristics, which is a prerequisite for the further photo-modulation of charge transport behavior within the MoS$_2$-GNR VDWHs. The physisorption of photochromic molecules onto the MoS$_2$-GNR VDWHs enables reversible light-driven control over charge transport. In particular, the drain current of the MoS$_2$-GNR FET can be photo-modulated by 52%, without displaying significant fatigue over at least 10 cycles. Moreover, four distinguishable output current levels can be achieved, demonstrating the great potential of MoS$_2$-GNR VDWHs for multilevel memory devices.
Graphene, the archetypal two-dimensional (2D) crystal, has been the target of both extensive fundamental studies and numerous technological advancements during the last 15 years.\[1\]

Graphene is an extremely promising material for future generation of flexible/wearable high speed optoelectronics by taking advantage of its extraordinary physical properties, including its ultra-high charge carrier mobility at room temperature, its outstanding mechanical robustness and its high light transmittance in the visible range.\[2\] However, the absence of a bandgap in its electronic structure does not enable graphene to be switched off in field-effect transistors (FETs) based logic circuits.\[1, 3\] The most promising and straightforward strategy for opening the bandgap in graphene relies on mastering quantum confinement effect via the synthesis of graphene nanoribbons (GNRs), which are narrow strips of graphene with atomically precise topology.\[4\] The electronic properties of GNRs are critically dependent on their width and edge structure, thus they can be tuned via the ad-hoc design of the synthetic precursor.\[5\]

Recent ground-breaking studies of on-surface synthesis both under ultra-high-vacuum (UHV)\[6\] and by chemical vapor deposition (CVD)\[7\] demonstrated the versatility of such bottom-up approach to generate a wide library of GNRs exhibiting excellent photo-conductivities while maintaining adjustable electronic bandgaps. Such features make these systems highly promising for future optoelectronics based on low-dimensional nanomaterials\[3b, 8\] (e.g. 0D, 1D, and 2D etc.).

The alternative strategy which has been pursued to overcome the lack of bandgap in graphene widens the focus to a broadest variety of 2D layered materials exhibiting diverse chemical structure and composition. The resulting electronic properties can range from insulators, semiconductors to conductors.\[3b, 9\] Among them, monolayer 2D transition-metal dichalcogenides (TMDCs), whose bandgap can be finely tuned, represent promising alternatives to graphene for applications in nanophotonics, nanoelectronics, nanoscale sensing and actuation, in the post-silicon era.\[10\] So far, MoS\(_2\) is the most extensively studied TMDC because of its robustness and high charge carrier mobility.\[3a, 9a, 11\] However, a major
drawback of MoS$_2$ and related hybrid structures for applications in optoelectronics is its persistent photoconductivity (PPC) effect,$^{[12]}$ which is defined as the light-induced enhancement in conductivity that persists for a long period after termination of photoexcitation,$^{[13]}$ i.e. the device cannot be totally switched off in the dark. Suppressing the PPC effect is therefore of utmost importance to leverage the fundamental features of MoS$_2$ for various technological applications.$^{[14]}$

The TMDCs’ properties can be “corrected” and optimized via the atomically controlled formation of van der Waals heterostructures (VDWHs),$^{[15]}$ by taking full advantage of the versatility of interfacing low-dimensional nanostructures without the constraints of crystal lattice matching.$^{[3b]}$ Hitherto, VDWHs have been primarily fabricated by superimposing two different types of 2D materials.$^{[15b]}$ Little has been done on forming VDWHs by superimposing nanostructures with different dimensionalities. In particular, the combination of 2D with 1D systems has been attempted to combine 2D graphene flakes with 1D dielectrics or conductors.$^{[16]}$ All-semiconducting 2D-1D VDWHs have not been properly addressed for applications in optoelectronics. The large library of recently developed 1D semiconductors, in particular based on suitably designed GNRs, open new avenues for explorations in the field of VDWHs.$^{[17]}$ Moreover, the modulation of the charge transport behavior of the VDWHs using different external stimuli is highly appealing for imparting them new functions. This should ultimately allow diverse applications based on novel stimuli responsive systems.$^{[18]}$

Herein, we fabricate a VDWHs by depositing 1D GNRs onto 2D MoS$_2$, to achieve an effective suppression of the PPC effect of MoS$_2$ as a result of interfacial charge transfer process. The hysteresis of the MoS$_2$-GNR FETs decreases significantly as compared to MoS$_2$ based FETs, and the transfer curves become more stable upon switching between 530 nm light irradiation and dark conditions. The reduction of the PPC effect is also evidenced in dynamic photoresponse analysis: the dark current value after illumination in FET based on
MoS$_2$-GNR VDH displays only 3% enhancement, being about one order of magnitude lower than the case of pristine MoS$_2$ FET. Such a minimal enhancement represents a prerequisite to monitor distinguishable current signal changes determined by light inputs, e.g., when photochromic molecular systems are integrated in the device. The molecular functionalization of MoS$_2$-GNR VDH is achieved through physisorption rather than chemisorption. Although chemisorption of reactive molecules typically leads to more robust hybrid systems as compared to physisorption, the former could be detrimental for the overall electronic properties of MoS$_2$ nanosheets, thereby irreversibly affecting the electrical characteristics of MoS$_2$-GNR VDH. Therefore, the less invasive physisorption approach is employed. A physisorbed photochromic spiropyran monolayer enabled a reversible remote-control of the charge transport in MoS$_2$-GNR FET, by exploiting the change in dipole moment between the two photo-isomers states acting as light activated gate. The photo-modulation of the source-drain current in MoS$_2$-GNR FET can be as high as 52%, which is durable without showing significant fatigue over at least 10 cycles. Moreover, upon varying the light irradiation period, four distinguishable output current levels could be achieved accordingly, which demonstrates the great potential of functionalized MoS$_2$-GNR VDH as prototypical active component in multilevel memory devices.

In order to assemble the VDHs based on MoS$_2$-GNR, monolayer MoS$_2$ is first isolated via micro-mechanical exfoliation from bulk MoS$_2$ crystal, followed by its transfer onto a SiO$_2$/Si substrate. GNRs with an armchair type edge structure and 7 carbon atoms in width (7-AGNR) are grown on Au/SiO$_2$ substrates under ultrahigh vacuum conditions, starting from 10,10'-dibromo-9,9'-bianthryl, which undergoes polymerization and cyclodehydrogenation at 200 °C and 400 °C, respectively.$^{[6b, 7d]}$ The 7-AGNR film is then transferred on top of the MoS$_2$ chip by a well-established poly(methyl methacrylate) (PMMA) supporting layer assisted lift-off and film transfer approach,$^{[7b]}$ yielding the VDH architecture portrayed in Figure 1a. Since the electronic properties of MoS$_2$ are highly
sensitive to the influence of its surrounding environment due to the unique atomically thin 2D structure, Raman and photoluminescence (PL) spectroscopy are employed to prove the formation of MoS\textsubscript{2}-GNR VDWH, as well as to monitor the electronic interactions between MoS\textsubscript{2} and 7-AGNR. The Raman spectra of the pristine MoS\textsubscript{2} flake (measured under 532 nm laser, Figure 1b) displays two characteristic first-order phonon modes, i.e. the in-plane oscillation E\textsubscript{2g}^1 mode and the out-of-plane oscillation A\textsubscript{1g} mode. The peaks which are located at 384.8 cm\textsuperscript{-1} and 403.9 cm\textsuperscript{-1}, thus having a ~19 cm\textsuperscript{-1} wavenumber difference, confirm the monolayer nature of the employed MoS\textsubscript{2} flake.\textsuperscript{[19]} It has been proved that for the monolayer MoS\textsubscript{2}, the A\textsubscript{1g} mode is more sensitive to the charge carrier doping level due to a stronger electron-phonon coupling, whereas the E\textsubscript{2g}^1 mode is more sensitive to strain due to the breaking of symmetry. Raman spectroscopy has also emerged as a powerful technique to study the vibrational modes for the sp\textsuperscript{2}-hybridized carbon nanomaterials. When a 7-AGNR film is placed on the top of the same monolayer MoS\textsubscript{2} flake, forming a network of 7-AGNRs covering the surface of MoS\textsubscript{2} to realize the VDWH architecture, four peaks located between 1608 cm\textsuperscript{-1} and 1219 cm\textsuperscript{-1} appear, which can be assigned to G, D, and edge C-H peaks typical of graphene, respectively.\textsuperscript{[7b]} More importantly, the sharp peak located at 395 cm\textsuperscript{-1} is the characteristic width-specific radial breathing-like mode (RBLM) peak, proving the structure of 7-AGNR. Since the RBLM is resonantly excited for photon energies near the lowest optical transitions of 7-AGNR, green laser (532 nm, 2.33 eV) is thus employed to just slightly exceed the optical gap of 7-AGNR (2.1 eV). Meanwhile, as the RBLM frequency varies strongly with GNR width, the sharp peak at 395 cm\textsuperscript{-1} corresponding to a width of 0.74 nm further illustrates the high uniformity in the width at the atomic level of bottom-up on-surface synthesized 7-AGNR. It is worth noting that the Raman spectroscopy is performed based on the 7-AGNR sample transferred from Au onto SiO\textsubscript{2} substrate with MoS\textsubscript{2} flakes. These Raman peaks are in excellent agreement with the calculated values via density functional theory, thereby demonstrating the high quality of the obtained 7-AGNR sample as well as the preserved
structural integrity of the film after the transfer process.\(^{[6b]}\) Moreover, the blue shift of the A\(_{1g}\) phonon mode of MoS\(_2\) by \(\sim 1 \text{ cm}^{-1}\) suggests the p-type doping effect of MoS\(_2\) by the addition of 7-AGNR.\(^{[20]}\) PL spectra of both pristine MoS\(_2\) and MoS\(_2\)-GNR VDWH are shown in Figure 1c, with the low-energy (A) exciton varies significantly over the spectral range of 1.8-2.0 eV. Since the A-exciton of monolayer MoS\(_2\) includes both neutral excitons (X) and trions (negatively charged excitons, X\(^-\)), the trion spectral weight can be correlated to the doping level of MoS\(_2\).\(^{[21]}\) The fitted peaks suggest that the spectral weight of the trion in the MoS\(_2\)-GNR VDWH is significantly lower than those in pristine MoS\(_2\). This feature results from the typical intrinsic n-type doping of MoS\(_2\) as well as the possible p-type doping effect of MoS\(_2\) by 7-AGNR, in accordance with the Raman results.\(^{[14]}\) Such interfacial charge transfer regime within MoS\(_2\)-GNR VDWH under light irradiation might be beneficial for the overall optoelectronic properties.\(^{[22]}\)

To cast light onto the optoelectronic properties of the MoS\(_2\)-GNR VDWHs, they have been integrated as the channel material into the back-gated FET device (schematic device geometry shown in Figure 1a), with top source-drain electrodes (60 nm thick gold) produced by a standard photolithography process. Because of the limited lateral size of the obtained monolayer MoS\(_2\) flakes exfoliated \(\text{via}\) the scotch tape method we opted to use the two-point probe geometry rather than four-point probe. The former geometry is widely employed to study the photoconductivity in MoS\(_2\) devices.\(^{[3a, 9a, 11, 23]}\) As a control experiment, a back-gated FET based on pristine MoS\(_2\) is also fabricated. Figure 2a and b shows the transfer electrical characteristics (\(I_{ds}-V_{g}\)) of the FETs based on pristine MoS\(_2\) and MoS\(_2\)-GNR VDWH measured under dark condition, plotted in both linear and logarithmic scale, while the source-drain voltage \(V_{ds}\) is kept at 1 V. The hysteresis on such transfer curves is investigated by recording consecutive forward and reverse sweeps. The FET based on pristine MoS\(_2\) reveals a significant hysteresis width of 5.2 V, which is a common feature attributed to the presence of trap states either on the MoS\(_2\) surface (possible charge transfer from/to absorbed molecules)
or at the MoS$_2$/SiO$_2$ interface.\textsuperscript{[3a, 24]} By comparison, the FET based on MoS$_2$-GNR VDWH displays a drastically smaller hysteresis width of 1.4 V. Such an improvement is directly correlated to the better interface between MoS$_2$ and 7-AGNR and possible encapsulation effects of 7-AGNR film, which is characterized by a severe reduction of trap states. In order to determine the possible PPC effect of pristine MoS$_2$, the linear $I_D$-$V_G$ characteristics of the FET are also measured under both dark and 530 nm illumination conditions, by performing over 50 cycles of such dark-illumination tests to gain insight into device stability toward light irradiation. As shown in Figure 2c, the dark current rises significantly with increased cycling number. From the 2$^{nd}$ cycle, the shape of the dark current curve begins to bend toward the high current direction, under the same $V_g$ and $V_{ds}$, the current value enhances, thus the resistance decreases and conductivity increases. Such light-induced enhancement in conductivity persisting for a long period after the termination of illumination behavior is in accordance with the definition of PPC effect.\textsuperscript{[3a, 13]} This pronounced PPC effect would influence the determination of the photocurrent value due to instability of the dark current, and is therefore detrimental to further applications of MoS$_2$ FET in optoelectronics. Conversely, the $I_{ds}$-$V_g$ characteristics of FET based on MoS$_2$-GNR VDWH are much more stable when undergoing 50 cycles of such dark-illumination tests than the case of pristine MoS$_2$ FET, as shown in Figure 2d. The dark current varies only in a small range, which demonstrates the suppressed PPC effect by the addition of 7-AGNR to MoS$_2$. Such phenomenon is attributed to the charge transfer regime within the interface of MoS$_2$-GNR VDWH, \textit{i.e.} the photogenerated electrons of MoS$_2$ are more favorable to be transferred into 7-AGNR, rather than filling the trap states on the MoS$_2$ surface or getting localized at the MoS$_2$/SiO$_2$ interface.

A dynamic photoresponse study of $I_{ds}$ at 0 V of gate voltage was also performed to better compare the PPC effect of the two FETs based on pristine MoS$_2$ and MoS$_2$-GNR VDWH. Figure 3 displays the comparison of the dynamic current variation under dark and
530 nm illumination conditions. When illumination is on, the \( I_{ds} \) in both two cases exhibits a sudden increase, which can be attributed to the photogenerated charge carriers. Once the light is turned off, the current of a pristine MoS\(_2\) FET is not fully recovered to the original value, being significantly higher than the adjacent dark current level with 36% enhancement. This is unambiguous evidence of the PPC effect. In contrast, the dark current value after illumination in FET based on MoS\(_2\)-GNR VDWH displays only 3% enhancement, being about one order of magnitude lower than the case of pristine MoS\(_2\) FET. The successful suppression of the PPC effect represents a milestone for further functionalization of the 1D-2D VDWH with light sensitive molecules.

Aiming at tuning the charge transport through the VDWHs, therefore enabling multifunctional control over the output current signals, the interfacing of 2D materials and hybrids thereof with photo-responsive molecular systems provides an efficient and versatile approach to achieve different charge carrier doping levels, ultimately leading to a reversible modulation of the electronic properties of the VDWHs.\(^{[10a, 18c, 25]}\) As a proof-of-concept, we have physisorbed a top layer of photochromic spiropyran (SP-C18) molecules and we explored the photo-modulation of the charge transport through MoS\(_2\)-GNR VDWH (Figure 4a). The initial device without SP-C18 molecules exhibits unipolar n-type behavior with pristine electron mobility of 21.4 cm\(^2\) V\(^{-1}\) s\(^{-1}\). Figure 4b shows that the presence of SP-C18 ad-molecules induces an increase of drain current \( I_{ds} \) and a negative shift of threshold voltage (\( \Delta V_{th} = -1.9 \) V), indicating a minor n-type doping effect by the SP-C18 molecules. After UV irradiation of the whole FET channel, the closed ring SP-C18 isomer is converted to the opening merocyanine (MC-C18) form, accompanied by a large variation of the molecular dipole moment, which induces a major increase of current and a continuous downshift of threshold voltage \( \Delta V_{th} = -4.1 \) V (Figure 4c). Therefore, the MC-C18 molecules further strengthen the n-type doping effect of MoS\(_2\)-GNR VDWH, with a significantly enhanced carrier concentration of \( 9.7 \times 10^{11} \) cm\(^{-2}\). The estimated sheet carrier density amounts to \( 3.8 \times 10^{12} \) cm\(^{-1}\).
and 4.8 x 10^{12} \text{ cm}^{-2} \text{ for MoS}_2\text{-GNR FET with SP-C18 molecules before and after UV irradiation.}^{[26]} \text{ The current modulation induced by the SP-C18/MC-C18 photo-switching is calculated to be around 52\%. By subsequent visible light irradiation, the reverse process takes place and all these electrical properties restore to the original values. Therefore, by programming the molecules into different states, the FET devices exhibit different electrical transport behavior. In total 10 illumination cycles are launched to investigate the stability of such light-induced current modulation. In Figure 4d, we plot the normalized current values obtained from the transfer curves at } V_g = 30 \text{ V, which proves a good stability and reversibility of our device system.}

Figure S1 displays the output characteristics of the MoS$_2$-GNR FET with SP-C18/MC-C18 molecules. It exhibits a linear and symmetric relationship between } I_{ds} \text{ and drain biases } V_{ds}, \text{ proving a low contact resistance with the electrodes. As a control experiment, transfer characteristics are recorded in both the dark and after UV/vis illumination on MoS$_2$-GNR FET device without SP-C18 molecules (Figure S2). It displays no obvious variation, thus proving that the SP-C18 molecules are responsible for the current modulation. To demonstrate the importance of a suppressed PPC effect, another control experiment on pristine MoS$_2$ FET device coated with SP-C18 molecules is performed. As shown in Figure S3, an increase of current can be observed after UV irradiation, but the current can only be partially restored to the original value after irradiation with visible light. This phenomenon is attributed to the fact that the PPC effect hinders the observation of the molecule-induced modulation. The dynamic evolution of current under alternative dark and UV/vis illumination conditions is portrayed in Figure 4e. When UV light is on (red boxed region), the current has a swift increase due to the photocurrent followed by a further yet gradual increase which is resulting from the SP-C18 to MC-C18 switching. The current level is higher than original curves when UV light is off, in accordance with the results obtained in the transfer measurement. Under visible light (blue boxed region), apart from the photocurrent, the current displays a gradual decrease which can be attributed to the MC-C18 to SP-C18
relaxation. The current level returns to the original value after one switching cycle. For the sake of comparison, control experiments on a MoS$_2$-GNR FET device without SP-C18 molecules are performed under the same illumination conditions. As displayed in Figure S4, the current reveals photoresponse under both UV and vis illumination, whereas it does not evidence any significant modulation after the illumination. In Figure 4f, we present a multilevel current obtained by exposing the device to a periodic short time UV illumination. The current level is proportional to the SP-C18/MC-C18 population ratio, therefore by controlling the irradiation time, the device can reach different current levels. At the end of each UV illumination, the current level shows a distinguishable increase compared with adjacent levels and the value remains stable. Upon 4 subsequent UV irradiations the device reaches 4 different current levels and as a result of the final long-time visible light irradiation enables the current to recover its original state, demonstrating its potential as multibit nonvolatile memories.

In conclusion, the fabrication of mixed-dimensional VDWH based on 1D 7-AGNR onto 2D MoS$_2$ represents an efficient strategy to suppress the PPC effect of MoS$_2$ due to the presence of interfacial charge transfer process, as evidenced by a 3-fold reduction of the hysteresis width, and more stable transfer characteristics upon switching between light irradiation and dark conditions. Such a platform is therefore ideal to photo-modulate the charge transport in MoS$_2$-GNR FET when decorated with physisorbed photochromic molecules. The drain current of the MoS$_2$-GNR FET can be photo-modulated by 52%, and four distinguishable output current levels can be achieved, demonstrating the great potential of MoS$_2$-GNR VDWH for multilevel memory devices. These results may pave the way toward ultra-thin multifunctional optoelectronics for logic-gates in the framework of integrating “More than Moore” technologies (analog functions, such as sensor, biochips, etc.) into CMOS digital circuits.
Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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References


Figure 1. Van der Waals heterostructures based on MoS$_2$-GNR. (a) Schematic illustration of the FET device configuration based on MoS$_2$-GNR, with the molecular structures of employed 7-AGNR. (b) Raman and (c) Photoluminescence spectroscopy characterization of MoS$_2$ and MoS$_2$-GNR, respectively.
Figure 2. (a) and (b) Hysteresis test of FETs based on (a) pristine MoS$_2$ and (b) MoS$_2$-GNR VDWH, in both linear and logarithmic scale as indicated by the blue and green arrows, under dark condition. Inset shows the digital pictures of the FET devices. (c) and (d) Transfer characteristics of FETs based on (c) pristine MoS$_2$ and (d) MoS$_2$-GNR VDWH, under dark and 530 nm illumination conditions over 50 cycles.
Figure 3. Comparison of the dynamic photoresponse of source-drain current upon 530 nm illumination (blue box area) between FETs based on pristine MoS$_2$ (red curve) and MoS$_2$-GNR VDWH (black curve), $V_g = 0$ V.
Figure 4. (a) Schematic illustration of the FET device architecture based on MoS$_2$-GNR with SP-C18 molecules. (b) and (c) Transfer evolution of the device before and after SP-C18 molecules deposition and after UV/vis illumination in (b) linear and (c) logarithm scale. (d) Current modulation with SP-C18 and MC-C18 isomers over ten illumination cycles of alternating UV (red boxed region) and visible (blue boxed region) light irradiation. All current values are normalized to the initial value obtained from the MC-C18 isomers. The connecting
lines are used as guides to the eye. (e) Dynamic current variation over alternative dark and UV/vis illumination conditions during four cycles. (f) Dynamic current variation under periodic UV illumination. Four distinct current levels can be achieved by short-time UV illumination and the current can be recovered by subsequent long-time vis illumination.