Photothermoelectric and photovoltaic effects both present in MoS$_2$

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As a finite-energy-bandgap alternative to graphene, semiconducting molybdenum disulfide (MoS$_2$) has recently attracted extensive interest for energy and sensor applications. In particular for broad-spectral photodetectors, multilayer MoS$_2$ is more appealing than its monolayer counterpart. However, little is understood regarding the physics underlying the photoresponse of multilayer MoS$_2$. Here, we employ scanning photocurrent microscopy to identify the nature of photocurrent generated in multilayer MoS$_2$ transistors. The generation and transport of photocurrent in multilayer MoS$_2$ are found to differ from those in other low-dimensional materials that only contribute with either photovoltaic effect (PVE) or photothermoelectric effect (PTE). In multilayer MoS$_2$, the PVE at the MoS$_2$-metal interface dominates in the accumulation regime whereas the hot-carrier-assisted PTE prevails in the depletion regime. Besides, the anomalously large Seebeck coefficient observed in multilayer MoS$_2$, which has also been reported by others, is caused by hot photo-excited carriers that are not in thermal equilibrium with the MoS$_2$ lattice.

Two-dimensional materials have lately attracted tremendous attention owing to their unique structural and physical properties\cite{1}. Understanding the basic physical phenomena as well as their intimate interplays in this new family of materials is essential for further exploitation of the materials towards electronic, photonic, and energy applications. Although it is largely graphene’s tribute that has sparked the recent zeal in two-dimensional materials and their potential applications, the intrinsic shortcoming without a bandgap hampers its application in electronics\cite{2}. For this reason, attention has been turned to other two-dimensional semiconducting transition metal dichalcogenides (TMDs) for their potential role in nanoelectronics, spintronics and optoelectronics\cite{3,4}. Among them, molybdenum disulfide (MoS$_2$), as a representative layered TMD, has a layer-dependent bandgap with a crossover from an indirect bandgap of 1.3 eV in its multilayer form to a direct bandgap of 1.8 eV when thinned to a monolayer\cite{5,6}. Both single- and multi-layer MoS$_2$ have been demonstrated to be suitable for electronic devices, such as field-effect transistors (FETs)\cite{7,8}, photodetectors\cite{9,10}, gas sensors\cite{11}, etc. However, multilayer MoS$_2$ is of greater potential than the monolayer counterpart from the fabrication viewpoint as well as the interesting physical properties. First, the electronic properties of multilayer MoS$_2$ manifested by an indirect bandgap are relatively insensitive to layer thickness. Second, multilayer MoS$_2$ is expected to carry higher drive current than monolayer MoS$_2$ due to higher carrier mobility arising from interlayer screening effect\cite{12} and the 3-fold higher density of states at the conduction band minimum\cite{8,13}.

With respect to optical properties, the subnanometer thickness of monolayer MoS$_2$ yields a low absorption cross-section hence results in less photo-generated carriers, as compared to that of multilayer MoS$_2$. Although fast photodetection has been achieved in monolayer-MoS$_2$ transistors\cite{14}, the photosresponsivity is expected to be lower than that of multilayer-MoS$_2$ devices\cite{15}, due to its direct bandgap and resultant shorter photocarrier recombination lifetime ($\sim 70$ ps)\cite{15}. For multilayer MoS$_2$, intervalley scattering is predominant over direct electron-hole (e-h) recombination when incident light energy is larger than direct bandgap at K point, thus generating microsecond-order photocarrier lifetime of indirect recombination\cite{16}. This was further proven by quenching of photoluminescence in multilayer MoS$_2$\cite{6}. The long carrier lifetime would have a positive effect on photosresponsivity. There is therefore greater potential with multilayer MoS$_2$ than monolayer MoS$_2$ for applications in broad-spectral photodetectors.

Investigations of multilayer MoS$_2$ and other TMD semiconductors are scarce and insufficient, compared to their monolayer counterparts, especially on optoelectronic properties. The origin of the observed photosresponsivity in MoS$_2$ is still under debate. Most studies suggest that photocurrent generated in the MoS$_2$ phototransistors results from photovoltaic effect (PVE)\cite{9,10,17,18}, similarly to what has been found with other conventional
semiconductors, e.g. Si, GaAs and GaN. Another photocurrent mechanism, i.e. photothermoelectric effect (PTE), is less discussed in the literature, in part due to the use of a broad-beam illumination experimental setup. Recent studies on graphene\textsuperscript{19,20} and carbon nanotubes\textsuperscript{21,22} indicate that PTE may play an important role in their photoresponse. As the amplitude of PTE relies on the temperature difference, multilayer MoS\textsubscript{2} with a well-defined bulk, and therefore a reduced thermal sinking effect via the substrate, is expected to yield a higher temperature increase under illumination\textsuperscript{23}. Because of the complex nature of MoS\textsubscript{2}, there is need to locally probe its photoresponse in order to allow the mechanisms of charge transport and collection in multilayer MoS\textsubscript{2} to be unequivocally unveiled and, in particular, to discern PTE from PVE.

In order to elucidate the nature of photoresponse in thick MoS\textsubscript{2} multilayers, we have employed scanning photocurrent microscopy (SPCM) in combination with ordinary current-voltage characterization to characterize the spatial dependence of photoresponse under various gate bias ($V_g$) conditions. At the metal-semiconductor interface, PVE is found to be predominant in the accumulation regime while thermal diffusion of photo-excited hot electrons is overwhelming and dictates the drifting current by the built-in electric field in the depletion regime.

**Results**

**Device structure and characterization.** The schematic representation of a back-gate MoS\textsubscript{2} transistor used in our work is shown in Fig. 1a while its optical microscope image is given in Fig. 1b. Isolated MoS\textsubscript{2} flakes on a SiO\textsubscript{2}/Si substrate were exfoliated from a bulk MoS\textsubscript{2} crystal using a conventional mechanical exfoliation technique\textsuperscript{24}. The sample preparation and device fabrication are detailed in Methods. The thickness of the MoS\textsubscript{2} flake in the device in Fig. 1b, as measured by atomic force microscopy (AFM), is approximately 65 nm (Fig. 1c) and its Raman spectrum in Fig. 1d shows two typical peaks (E\textsubscript{2g} and A\textsubscript{1g}) with a large separation of 25 cm\textsuperscript{-1}, confirming that the multilayer nature of the MoS\textsubscript{2} flake\textsuperscript{25}.

**Photoelectrical stability of MoS\textsubscript{2} transistor.** Electrical characterization of our MoS\textsubscript{2} FETs was performed in ambient environment. Similar to other low-dimensional devices without surface protection\textsuperscript{26–28}, MoS\textsubscript{2} FETs are highly sensitive to extrinsic effects, especially adsorption of H\textsubscript{2}O/O\textsubscript{2} from air\textsuperscript{29,30}. These effects can induce current hysteresis in transfer characteristics and electrical stress instability of MoS\textsubscript{2} FETs when measured in conventional DC mode (Supplementary Fig. S1). The electrical instability can lead to persistent photoconductivity when switching on and off the laser illumination and therefore, a reduction of photoresponsivity with switching time (Supplementary Figs. S3 and S4). Therefore, stability of the device under electrical stress is a prerequisite for practical optoelectronic applications of MoS\textsubscript{2}. Recently, electrical characterization with $V_g$ pulses of alternating polarities i.e. AP mode, has been reported to distinctively reduce the device instability compared to using DC mode\textsuperscript{28,31,32}. As shown in Supplementary Fig. S1 and Fig. 2a, the hysteresis of our MoS\textsubscript{2} FETs is almost completely eliminated and stable electrical characteristics are obtained using the AP method. This method leads indeed to stable photoswitching behaviors when applying multiple illuminations (Fig. 2b and Supplementary Fig. S4), which can only be achieved in vacuum if MoS\textsubscript{2} FETs are measured using the DC method\textsuperscript{33}. Since the AP method is not based on any structural or chemical modification of the devices, as opposed to other measures employed to improve the device stability\textsuperscript{26,29,34}, it is of great potential in leading to more repeatable and clearly interpretable photo-sensing signals. Therefore, all our photoelectrical measurements were performed in AP mode in order to avoid additional stressing of the devices.

**SPCM characterization of MoS\textsubscript{2} transistor.** To identify the mechanism of photocurrent generation, SPCM was conducted at zero source-drain bias ($V_d = 0$ V) with a well-focused laser beam.

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Figure 1 | Device structure and characterization. (a) Schematic and (b) Optical image of the field-effect transistor based on multilayer MoS\textsubscript{2}. (c) AFM line scan along the dashed line across the boundary of the MoS\textsubscript{2} flake in the inset. Inset: high-resolution AFM image of the MoS\textsubscript{2} multilayer on SiO\textsubscript{2}/Si substrate. (d) Raman spectrum for the MoS\textsubscript{2} multilayer on SiO\textsubscript{2}/Si substrate. The frequency separation between E\textsubscript{2g} and A\textsubscript{1g} peaks is 25 cm\textsuperscript{-1}. 

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scanning over the device area. The short-circuit photocurrent and Raman spectra were recorded simultaneously as a function of laser position. To avoid possible photo-damage to the MoS$_2$ during experiment, the step size was set to be 1 µm. Figure 3a shows the photocurrent image measured at $V_g = 15$ V, where two photocurrent extrema of opposite polarities are observed at the electrode edges. As expected from conventional PVE, the opposite built-in electric fields from space charges at the source/drain contacts create an asymmetrical photocurrent characteristic. Such PVE-induced photocurrents localized near electrodes have been widely reported in devices based on silicon nanowires and carbon nanotubes. However, it is remarkable to observe a significant photocurrent when the laser spot is centered inside the electrodes. The distance from the electrode edge is $\sim$10 times larger than the laser spot size in order to make sure that the MoS$_2$ channel is not illuminated. Besides, the light penetration depth is smaller than the thickness for the metal electrodes at the wavelengths used. The absence of Raman signals from the MoS$_2$ flake under the Ti/Au electrodes in Fig. 3b implies, therefore, that the observed photocurrent inside the electrodes could not arise from PVE at the MoS$_2$-metal interface under the electrodes. Another photocurrent mechanism than PVE should be at work.

In fact, the absorption of laser photo energy by the metal electrodes in this manner could result in an inhomogeneous local temperature distribution. When a temperature gradient is generated by laser heating across a junction comprising two materials of different Seebeck coefficients, a thermal voltage driving a current through the device is established via thermoelectric effects. The polarity of thermoelectric current relies on the majority carriers (i.e. electrons or holes) and reverses sign in between the source and drain electrodes. As shown in Fig. 3a, the photocurrent image in the electrode area coincides with the optical surface topography of the MoS$_2$ flake underneath the metal electrodes. This suggests that the PTE mechanism cannot be ignored in the photocurrent contribution. And the PTE-induced photocurrent is also expected to be intensified at the electrode edges where light can be absorbed by both materials and the heat dissipation is lower at the electrode edge than inside the metal electrodes.

The temperature rise at the MoS$_2$-Ti/Au interface under laser illumination can be measured by Raman spectroscopy. Since the positions of the Raman-active phonons of MoS$_2$ are sensitive to temperature, the local temperature change induced by laser heating can be quantified. For calibration, the laser power was minimized in order to avoid heating effects in the Raman measurement and the temperature of the whole device was tuned by a temperature controller. The temperature-dependent and laser-power-dependent shifts of the Raman mode $E_{2g}$ are plotted in Supplementary Fig. S5. The linear temperature coefficient for mode $E_{2g}$ is extracted to be $-0.015$ cm$^{-1}$/K in Supplementary Fig. S5c. In combination with the power-dependent coefficient of $-0.74$ cm$^{-1}$/mW (Supplementary Fig. S5d), the temperature rise in the illuminated spot can be acquired. From this measurement, a 100-µW laser illumination at the MoS$_2$-Ti/Au contact could result in a temperature increase by $\sim$5 K. Thicker MoS$_2$ layer could lead to a higher temperature rise due to the reduction of thermal coupling to the substrate.

**Photothermoelectric and photovoltaic effects.** In the following discussion, we focus on two main competing mechanisms of photocurrent generation: PVE and PTE. The energy band diagrams of a MoS$_2$ transistor with a Schottky barrier for various $V_g$ conditions are schematically shown in Fig. 3c. For large positive $V_g$, the conduction band bends downward at the metal/semiconductor interface and the direction of the built-in electric field points from MoS$_2$ to electrode. When $V_g$ moves from positive to negative, the band bending direction changes from downward to upward and the built-in electric field reverses. The polarity of the photocurrent generated by PVE is dependent on the direction of the built-in electric field. However, in the PTE model, the generated photocurrent is equal to thermal voltage divided by device resistance $R$. Here, the thermal voltage is proportional to the local temperature increase in the junction, as well as to the difference in the Seebeck coefficients ($S$) between the MoS$_2$ and the electrode. Herein, $S$ of the metal electrodes is negligible with respect to that of MoS$_2$; the latter is gate-dependent and given by the Mott relation as follows:

$$S = \frac{\pi^2 k_B^2 T}{3q} \frac{1}{R \Delta V_g}$$

where, $k_B$ is the Boltzmann constant, $q$ is elementary charge, $T$ is absolute temperature, and $E_F$ is Fermi energy. The calculated $S$ is plotted in Fig. 3d (black curve) using equation (1) in combination with the transfer characteristic curve of the device in dark. $S$ shows a non-monotonic variation with gate bias, maximum at the threshold voltage, $V_T$ ($\sim$10 V) and reversing sign as the majority carrier changes from electron to hole when $V_g < -15$ V. At $V_g > 0$ V, both PVE and PTE contributions have the same sign, and therefore yield a larger photocurrent, as illustrated in Fig. 3c. However, in the regime where the built-in electric field just switches the sign, the two contributions can have opposite signs and the photocurrent can then have either sign depending on their relative strengths.

In order to gain more insights into the photocurrent generation mechanisms, we performed line-scan across the contact edge at a step size of 200 nm as shown by the black dashed line in Fig. 3a at various $V_g$ (Figs. 4a–b). The peak position and peak amplitude of the photocurrents are extracted and plotted in Fig. 4c. The most striking

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feature in the plot is the movement of the peak position at the contact edge as $V_g$ is decreased below $-5$ V. Concurrently, the peak amplitude decreases exponentially with $V_g$ without sign flip. According to Fig. 3d, the thermoelectric photocurrent is expected to be negligible in the depletion regime ($-15 < V_g < -10$ V) since $S$ approaches zero in this region. As a result, PVE should dominate the photocurrent there. When $V_g$ is changed from positive to negative, the built-in electric field reverses and the peak electric field at negative $V_g$ moves away from the contact edge and extends more into the channel region compared to that at positive $V_g$ (Supplementary Fig. S6). At $V_g = -15$ V, the photocurrent peak moves $\sim 0.6$ µm away from the contacts, which is in qualitative agreement with the movement of the simulated peak electric field. If the photocurrent generation is dominated by PVE in the depletion regime, the sign of photocurrent should reverse when ($V_g - V_t$) changes from positive to negative, which is however not observed experimentally. The contradicting polarity of the photocurrent suggests that the effect of the built-in field is less likely to be the dominant mechanism to determine the direction of the photocurrent in the depletion regime.

**Thermoelectric power with two-carrier transport.** Actually, the Mott formulism, which is established on the basis of thermal diffusion of one carrier type, is not valid for two types of carriers involved in the ambipolar transport. With light illumination, e–h pairs are generated when the photon energy exceeds the bandgap. Such excited carriers should contribute not only to the electrical conductivity, but also to the thermoelectric power through changing carrier concentration, which is more significant in the depletion regime. For such a two-carrier process, the thermoelectric power is given by

$$S = -\frac{k_B}{q} \frac{n \mu_n \left( A_n + \ln \frac{N_e}{n} \right) - p \mu_p \left( A_p + \ln \frac{N_v}{p} \right)}{n \mu_n + p \mu_p}$$

where, $n$ and $p$ are electron and hole concentrations, respectively, $\mu_n$ and $\mu_p$ are the corresponding mobilities, and $N_e$ and $N_v$ are the effective density of states in the conduction band and valence band, respectively, and $A_n$ and $A_p$ are thermoelectric constants of electrons and holes. When the concentration of one type of carriers is much higher than that of the opposite type of carriers, such as in the accumulation regime, $S$ is then dominated by the majority carriers and equation (2) is reduced to the Mott relation$^{43}$. If the electrons and the holes are equal in number, for example for photo-excited e–h
pairs in the depletion regime, the effective carrier density of states would play a dominant role in determining the amplitude and sign of thermoelectric power. In multilayer MoS$_2$, the minimum of the conduction band lies in the lower symmetry Q point in the k-space along the $\Gamma$-K line with a 6-fold valley degeneracy while the single valence band maximum is situated at the $\Gamma$ point. This results in an effective density of states in the conduction band almost 6 times that in the valence band by assuming an equal effective mass for electrons and holes. Therefore, PTE in the multilayer MoS$_2$ is still dominated by the photo-excited electrons and both $S$ and conductivity increase with illumination in the depletion regime. The unchanged polarity of photocurrent in the MoS$_2$ FETs, when $V_g$ is swept from positive to negative in Fig. 4b, reveals that PTE is predominant over PVE in the depletion regime. However, this photo-induced $S$ in MoS$_2$ is spatially inhomogeneous along the channel. In the space charge region, the built-in electric field reduces the e–h recombination rate, resulting in a higher $S$ compared to that in the flat-band region. Recent studies reveal that a nonuniform $S$ in the channel could produce an additional PTE photocurrent, which is different from that at the semiconductor-metal junction. This PTE photocurrent, as observed in Fig. 4b, moves with the gate bias in the depletion regime due to the photo-induced $S$ depending on the electric field.

Figure 4 | Gate-dependent photocurrent in MoS$_2$ transistor. (a) Photocurrent ($I_{PC}$) line scans across the contact edge as indicated by the black dashed line in Fig. 3a at $V_g$ from 0 V to 20 V and (b) $V_g$ from 0 V to −20 V. The red arrows in panels (a) and (b) indicate the position of peak photocurrent. (c) Photocurrent peak position and amplitude in panels (a) and (b) varied with $V_g$. (d) The laser-power ($P$) dependency of the photocurrent with laser illuminated at the contact edge for $V_g$ = 15 V and −15 V. The black and red lines are power-law fits with $I_{PC} \propto P^\alpha$.

Hot photo-excited carrier inducing giant Seebeck coefficient. Apart from short-circuit photocurrent, open-circuit photovoltage can be a complementary diagnostic parameter to further probe the nature of the photoresponse. As shown in Fig. 3d (red curve), the gate-dependent photovoltage is measured at the contact edge. Identical to the photocurrent in Fig. 4c, the sign of photovoltage is not reversed during the gate sweeping and the maximum of photovoltage is shifted to more negative voltage (−15 V) compared to the gate-dependent $S$. Obviously, such a negative shift can be attributed to the photo-excited carriers. Since the main contribution of photovoltage arises from PTE in the depletion regime, the maximum $S$ can be estimated from the photovoltage because there is only a constant temperature difference in between them. We have already determined that approximately a temperature increase by 5 K is obtained with a 100-μW laser illumination. The 20.18 V peak photovoltage would correspond to an $S$ value of $\Delta S$ = 3.6 × 10$^4$ $\mu$V/K, which is one order of magnitude larger than the calculated maximum $S$ value (black curve in Fig. 3d). This discrepancy in $S$ has also been reported by Buscema et al. Nonetheless, the $S$ value obtained for multilayer MoS$_2$ is much larger than that of graphene$^{19,20}$, carbon nanotubes$^{21,22}$ and other thermoelectric materials$^{46,47}$. In the
estimation of S from the photovoltage data, the temperature increase induced by laser illumination is assumed to be equivalent to the temperature rise causing the Raman shift in the MoS$_2$. However, this Raman-shift thermometry measures the phonon temperature instead of the electron temperature in the MoS$_2$. It is the gradient in the electron, rather than phonon, temperature that ultimately drives the PTE effect. In the course of photo-excitation, the electrons and the phonons do not reach thermal equilibrium$^{48}$. For example, in laser-illuminated graphene layers, electrons are much hotter than the surrounding crystal lattices and form a hot Fermion distribution$^{19,49,50}$. Therefore, the measured phonon temperature is likely to underestimate the temperature of the electrons in the multilayer MoS$_2$, resulting in a false anomalously giant $S$.

Strikingly, when $V_g < -15$ V, the calculated $S$ in Fig. 3d reverses sign due to a weak hole inversion, while the polarity of photovoltage is opposite still staying positive. This suggests that the thermoelectric effect is still dominated by the excited hot electrons according to equation (2). In the strong electron accumulation regime ($V_g > 0$ V), the photovoltage saturates at a relative high value of $-0.12$ V, which results in a ratio of maximum-to-saturation photovoltage to be $-1.5$. This is nearly two orders of magnitude smaller than that of calculated $S$ ($-10$). This indicates that PTE rather than PVE is the dominant effect in the accumulation regime because the high electron concentration induced by gate voltage effectively cools the excited electrons due to the strong electron-electron interaction in reduced-dimension semiconductors$^{45}$.

It has recently been demonstrated that the power-law relationship between photocurrent and incident laser power ($P$), i.e., $I_{ph} \propto P^n$, can be used as an alternative approach to characterizing the dominant mechanism for photocurrent generation$^{2}$: the exponent $n$ is unity for PVE and 2/3 for PTE. The laser-power dependency of the photocurrent at $V_g = 15$ V and $-15$ V is shown in Fig. 4d with the laser focused at the contact edge. The linear relationship at $V_g = 15$ V and the sublinear dependence on the laser power ($n = 0.69$) at $V_g = -15$ V further confirm that PVE dominates in the accumulation regime while PTE dictates in the depletion. This observation strikingly contradicts the common perception that one photocurrent mechanism (PVE or PTE) should dominate in the whole working regime, as is the case for nanotubes$^{12,22,37,51}$ and graphene$^{49,52}$. That PTE rather than PVE dominates in the depletion regime can be explained as follows. The optical absorption of MoS$_2$ creates hot electrons. They can gain sufficient kinetic energy to overcome the potential barrier in the depletion regime. As a result, the hot electrons thermally diffusing along the channel generate the observed photocurrent in the depletion regime.

Discussion

A spatially resolved photoresponse of multilayer-MoS$_2$ transistors has been investigated by using SPCM. When exposed to ambient environment, the transistors are electrically unstable causing undesired hysteresis and persistent photocondoictivity when they are characterized using conventional DC measurement. Such instabilities are completely eliminated by using a novel pulsed-gate-bias method. As a result, a stable and repeatable photoresponse of MoS$_2$ transistors is, for the first time, to our knowledge, achieved in ambient environment. This has allowed us to systematically explore the interplay of thermal, optical and electrical behaviors of multilayer MoS$_2$. Both photovoltaic and photothermoelectric effects are found to be at work in the multilayer-MoS$_2$ transistors, although they take dominant roles in different regimes of device operation. In the accumulation regime, the PVE at the MoS$_2$-metal interface is the dominant mechanism whereas the hot-carrier-assisted PTE prevails in the depletion regime. This behavior is different from that observed in other low-dimensional materials that only contribute with either PVE or PTE. Moreover, the anomalously large Seebeck coefficient in MoS$_2$, which has also been reported by others, is found to be the consequence of an oversimplification by assuming thermal equilibrium of the hot photo-excited carriers with the surrounding phonons. Our findings that shed light on the photoresponse mechanism of multilayer MoS$_2$ may apply to other TMDS as well. They will also be helpful in advancing the art in light harvesting and photodetector applications.

Methods

Exfoliation and characterization of MoS$_2$ thin flakes. Thin MoS$_2$ flakes were peeled off from bulk MoS$_2$ (SPI supplies) by mechanical exfoliation. They were subsequently transferred to a heavily doped p-type Si substrate with a 300-nm-thick thermally grown SiO$_2$. The SiO$_2$/Si substrate was pre-cleaned by sonication in acetone, isopropanol alcohol, and deionized water. The transferred MoS$_2$ flakes were identified using an optical microscope (Keyence digital microscope VHX-600). The thickness of the MoS$_2$ flakes was measured using AFM (Dimension 3100 with Nanoscope IIIa controller, Veeco) operated in tapping mode under ambient conditions. Raman spectra of MoS$_2$ films were acquired by employing micro-Raman spectrometry (Renishaw inVia) in a backscattering configuration with a He-Ne 633 nm laser. The scattered signal was collected through a 100× objective and dispersed by a grating of 1800 lines/mm with a spectral resolution of ~1.0 cm$^{-1}$ and a peak position accuracy of 0.1 cm$^{-1}$. The power of the excitation laser line was kept well below 0.1 mW to avoid sample heating and an exposure time of 30 s was used to improve the signal to-noise ratio.

Device fabrication and electrical characterization. In order to avoid the contamination from photolithography or electron-beam lithography, a 10-μm spacing copper grid was placed on top of the thin MoS$_2$ flakes as a shadow mask for the electrode fabrication. A bilayer stack Ti/Au of 5/50 nm thickness was then deposited by means of electron-beam evaporator as the source and drain electrodes. Heavily doped Si substrate was used as the common back gate for the fabricated MoS$_2$ FETs. Electrical characterization of the devices was carried out in a shielded probe station with Keithley 4200 semiconductor characterization system in the ambient environment.

SPCM characterization. Spatial dependence of photoresponse in the MoS$_2$ transistors was obtained on the Renishaw Raman spectrometer equipped with the same electrical characterization system described above. A laser beam of 633-nm wavelength was focused to a diffraction-limited spot size of about 0.6 μm by using a long working distance 50× objective lens. The incident laser power could be adjusted with the use of neutral density filters. A computer-controlled XYZ translational stage was used to acquire scanning photocurrent image and Raman mapping image simultaneously. When the laser spot was raster-scanned over the device area, the resultant current flow between electrodes and the Raman spectrum for each point in the scan were recorded simultaneously, thus generating spatially resolved images of photocurrent and Raman signal. All scanning photocurrent measurements were performed at room temperature in air.

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Author contributions
Z.-J.Q. conceived the research. Z.-J.Q. and Y.Z. designed the experiments. H.W., X.X. and Z.-J.Q. conceived the research. Z.-J.Q. and Y.Z. designed the experiments. H.W., X.X. and J.Z. conceived the research. Y.Z. performed AFM analysis, Raman spectroscopy, and photocurrent response in graphene. Nano Lett. 7, 114–118 (2012).

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