Rhombohedrally stacked MoS₂ has been shown to exhibit spontaneous polarization down to the bilayer limit and can sustain a strong depolarization field when sandwiched between graphene. Such a field gives rise to a spontaneous photovoltaic effect without needing any p-n junction. In this work, we show that the photovoltaic effect has an external quantum efficiency of 10% for devices with only two atomic layers of MoS₂ at low temperatures, and identify a picosecond-fast photocurrent response, which translates to an intrinsic device bandwidth at ~100-GHz level. To this end, we have developed a nondegenerate pump-probe photocurrent spectroscopy technique to deconvolute the thermal and charge-transfer processes, thus successfully revealing the multicomponent nature of the photocurrent dynamics. The fast component approaches the limit of the charge-transfer speed at the graphene-MoS₂ interface. The remarkable efficiency and ultrafast photoresponse in the graphene-3R-MoS₂ devices support the use of ferroelectric van der Waals materials for future high-performance optoelectronic applications.

**INTRODUCTION**

When two layers of transition metal dichalcogenides (TMDs) are stacked in parallel, both the inversion symmetry and mirror symmetry are spontaneously broken, leading to an electric polarization along the out-of-plane direction (1). Under an electric field larger than the coercive field, the polarization direction can be flipped as one layer slides relative to the neighboring layer, which has been termed sliding ferroelectricity (2–5). While the artificially stacked bilayers (BLs) have limited domain sizes due to twist misalignment, homogeneous polarization in a scale as large as the entire flake has been observed in BLs directly exfoliated from a 3R-MoS₂ bulk crystal (6), making it ideal for certain optoelectronic applications. In particular, when 3R-MoS₂ is sandwiched between graphene (7), the spontaneous polarization gives rise to a photovoltaic effect, known as spontaneous photovoltaic effect (8), where photoexcited carriers in MoS₂ transfer asymmetrically to the graphene under a largely unscreened depolarization field. An external quantum efficiency (EQE) of 16% has been observed in a device with 10-layer 3R-MoS₂.

The high EQE in the graphene/3R-MoS₂/graphene heterostructures suggests a potentially ultrafast photocurrent dynamics comparable to the charge-transfer process at the MoS₂-graphene interface, which lasts approximately 1 ps (9–13). Such an ultrafast photocurrent, if confirmed, can potentially be leveraged for high-speed optical communications. Although the MoS₂ photocarrier dynamics generally can be measured by ultrafast techniques such as optical pump-probe spectroscopy (14), it remains a challenge to probe the intrinsic dynamics of the aforementioned photovoltaic effect, due to the existence of multiple photocurrent contributions. Because the spontaneous polarization naturally induces image charges of opposite polarities at the two electrodes, both the bolometric and the photo-thermoelectric (PTE) effects exist in the device (15–20). In addition, the asymmetric tunneling barrier is also known for generating a photocurrent without external bias (21, 22). Here, we disentangle these electronic and thermal contributions and measure their corresponding response times by performing ultrafast photocurrent autocorrelation and nondegenerate pump-probe photocurrent measurements. We conclude that the photocurrent has approximately 2-ps fast dynamics for devices of various thicknesses and a slow dynamics that lasts for 25 ps or more.

Ultrafast optical techniques are powerful tools for probing the photocarrier dynamics (9, 14, 23). Photocurrent autocorrelation technique has been previously used to study the photocurrent dynamics in two-dimensional (2D) material–based photodetectors (24–28). In this approach, two equal-power laser pulses are split from a single femtosecond pulse and illuminate the heterostructure to generate a photocurrent (Fig. 1A). When these two pulses coincide on the device, a saturation in the photocurrent can be observed due to either electronic or thermal mechanisms. For example, electronic saturation may arise from absorption saturation and increased interactions between carriers, whereas thermal saturation may be induced by the nonlinear temperature increase under illumination (29–32). By scanning the delay between these two pulses, a characteristic time associated with the saturation mechanism can be measured. When multiple mechanisms contribute to the photocurrent, the intrinsic photocurrent time response can be obscured.

**RESULTS AND DISCUSSION**

We first perform the photocurrent autocorrelation measurement on a heterostructure composed of both BL and four-layer (4L) 3R-MoS₂. The optical image of the heterostructure is shown in fig. 1.
From the current-voltage characteristics shown in Fig. 1B, a spontaneous photovoltaic behavior is clearly observed. The zero-bias photocurrent mapping shows a finite photoresponse in BL and 4L regions, and the photocurrent distribution agrees with the MoS$_2$ thickness in the graphene overlapped area (Fig. 1C; photocurrent maps for other devices can be found in fig. S2). This photocurrent grows sublinearly with optical power (fig. S3), giving rise to a negative autocorrelation signal. A representative autocorrelation result is shown in Fig. 1D. A dip of the photocurrent strength is observed at zero delay, with symmetric recovery dynamics at both positive and negative delays, as expected for conventional photocurrent autocorrelation signals (24–28).

More quantitative analysis of the autocorrelation signal in the BL area is presented in Fig. 2. With a fluence of 140 µJ cm$^{-2}$ for a spot size of approximately 1 µm in diameter at 770 nm (50-fs pulse width) for each pulse, the saturation dip at zero time delay is approximately 15% of the steady-state signal. The recovery time is fitted to be 17 ps, more than one order of magnitude slower than the charge-transfer time at the MoS$_2$-graphene interface (10, 12), suggesting that the saturation mechanism is likely not electronic. If such a slowing is caused by the charge transfer at the MoS$_2$-MoS$_2$ interface, a previous study suggests that an external electric field can speed up the process when the TMD is not monolayer (24). However, when we apply a bias voltage from −0.5 to 0.5 V across the device, no substantial change is observed in the recovery time (Fig. 2A, inset; the raw data are shown in fig. S4).

Instead, we suggest that this photocurrent saturation originates from photo-thermal effects. As shown in Fig. 2C, the photoresponse of our device has a surprisingly strong temperature dependence. When the heterostructure is heated by 10 K above room temperature (RT), the photocurrent drops by more than 25% (Fig. 2C, inset). More notably, the photoresponsivity increases by more than one order of magnitude after cooling the heterostructure from RT to 3 K (Fig. 2C, see also in another device in fig. S5, full temperature range in fig. S6A). At the A-exciton resonance, the EQE approaches 10%, which is remarkable for an atomically thin device. In addition, we also observe a pronounced temperature dependence in the total resistance of the device (fig. S6B), which motivates us to develop a shunt-resistance model. In this model, within a few tens of kelvin above RT, which is relevant to our pump-probe photocurrent measurement, only the shunt resistance is expected to change markedly due to its tunneling current origin (33). By modeling the measured photocurrent to be proportional to the shunt resistance, which exponentially decreases with increasing temperature, we are able to fit the temperature dependence of the photocurrent (Fig. 2C, inset, details in the Supplementary Materials). Other factors that can contribute to this strong temperature dependence include the variation in spontaneous polarization (34, 35) and...
change in photocarrier recombination time or exciton linewidth, which affects the absorption of the heterostructure. As a result, when an ultrafast laser pulse is absorbed by the device, the transient temperature of the illuminated region quickly rises, which decreases the shunt resistance and subsequently decreases the photocurrent generated by the following pulse. In this case, the saturation recovery time corresponds to the device heat dissipation time. At low temperatures, the photocurrent becomes less temperature dependent (fig. S6), which is consistent with the smaller autocorrelation signal observed at 3 K, possibly due to the internal quantum efficiency limit.

We confirm the photo-thermal origin of the saturation by measuring the photocurrent autocorrelation at cryogenic temperature (Fig. 2B). After most phonon modes are frozen at low temperature, the thermal conductivity of the 2D materials decreases substantially (36, 37), which should lead to a longer heat dissipation time. Experimentally, we find a recovery time that is approximately three times longer (55 ps) than that at RT, when the heterostructure is cooled to 3 K. All these evidences suggest that the dynamics observed in the autocorrelation measurements is limited by the device heat dissipation.

In addition to this long recovery time, we also observe a transient middle peak in the autocorrelation measurement at 3 K. Exponential fitting of that small peak suggests a time constant of approximately 4 ps. Similar peaks have been observed in the 4L region (fig. S7A) and on other BL devices of varying amplitudes up to RT (fig. S7B). We interpret this feature as a result of the electronic temperature saturation in the graphene electrodes. As discussed in our previous work (7), the top and bottom graphene electrodes are naturally doped with charges of opposite polarities in these devices, which consequently acquire opposite Seebeck coefficients. Upon laser heating, a PTE current in a direction opposite to the photovoltaic current is generated, which reduces the net photocurrent. In the autocorrelation experiment, the two subsequent laser pulses with a small delay time saturate the graphene electronic temperature, which decreases the PTE effect and therefore increases the overall photocurrent. The characteristic time of the middle peak also agrees with graphene electronic temperature relaxation through superradiation (27, 28, 38).

Because the photo-thermal effect of the laser pulse obscures the electronic response speed in autocorrelation measurements, an alternative method is needed to separate the thermal and electronic contributions. Here, we develop a nondegenerate pump-probe photocurrent spectroscopy technique that permits the determination of the intrinsic speed of our devices (Fig. 3A). First, we use a strong sub-bandgap infrared pulse (1030 nm, 50 fs, 350 μJ cm\(^{-2}\); IR-pulse) to heat the graphene electrodes, which subsequently heats the MoS\(_2\) region. Second, we use a weak visible pulse (670 nm, 50 fs, 2 μJ cm\(^{-2}\); VIS-pulse) that is resonant with the A exciton of MoS\(_2\) to generate a photocurrent pulse but without generating sufficient heat.

At negative delays when the VIS-pulse arrives earlier than the IR-pulse, despite the fact that the IR-pulse generates a finite photocurrent, the photo-thermal saturation effect should be negligible because little heat is deposited by the VIS-pulse (details in the Supplementary Materials). The photocurrent should decrease considerably immediately after the zero delay because the substantial heating induced by the IR-pulse would reduce the photocurrent generated by the VIS-pulse, in accordance with the results from our autocorrelation experiments. At a large positive delay, the photocurrent should be restored to the steady state as the heat is fully dissipated. Therefore, as we scan the time delay, we expect to see an asymmetric pump-probe photocurrent signal similar to photocarrier dynamics from an optical pump-probe measurement, which exhibits a quick rise followed by an exponential decrease in signal. Because the

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**Fig. 2. Temperature-dependent measurements in the BL region.** Autocorrelation signals of the BL region measured at (A) RT and (B) 3 K. The dotted lines are from experimental measurements, and black lines are exponential fittings to extract the time constants. Inset of (A): photocurrent time constant extracted from bias voltage–dependent autocorrelation measurements. The dashed line indicates a photocurrent time constant, \(\tau\), of 17 ps. (C) Energy-dependent photoresponsivity of the BL region measured at RT and 3 K. A and B peaks correspond to the A-exciton and B-exciton absorption features in BL 3R-MoS\(_2\). The average power is kept at around 0.5 μW at all wavelengths to avoid saturation. Inset: photocurrent of the BL region at temperatures above RT with 532-nm CW laser excitation. The dots are experimental data, and the solid line is a fit based on the shunt-resistance model.
The pump-probe photocurrent dynamics is approximately a cross-correlation between the photocurrent and device temperature dynamics, the decay of the signal is likely limited by the heat dissipation, while the rise of the signal should be determined by the photocurrent decay and lattice heating.

Experimentally, a highly asymmetric temporal response is observed in the pump-probe photocurrent measurement (Fig. 3B). The pump-probe photocurrent signal captures the photocurrent change due to thermal saturation. As expected, we find a very sharp drop of the photocurrent around the zero delay. The drop lasts for approximately 4 ps (Fig. 3B, inset), which gives an upper bound for the characteristic time of the transient photocurrent. It is followed by a 20-ps slow recovery dynamics similar to that observed in the autocorrelation experiment, corresponding to the cooling of the device. These pump-probe photocurrent results confirm the existence of a picosecond photocurrent response. In addition, a small photocurrent saturation is observed ahead of the sharp drop before zero delay, suggesting that multiple dynamics could be associated with the photocurrent. The delayed IR-pulse can affect the photocurrent generated earlier by the VIS-pulse if the photocurrent lasts long enough to overlap with the heating pulse. With the fast photocurrent component determined from the quick drop, the saturation long before the zero delay indicates the existence of a slow photocurrent component. In addition, after symmetrizing the pump-probe photocurrent signal by adding itself to the time-reversed copy, the result shows similar dynamics as the photocurrent autocorrelation signal, except that the autocorrelation is affected by the PTE effect near zero delay (details in the Supplementary Materials). Since the signal in the pump-probe photocurrent experiment is much larger at positive delay than that at negative delay, we confirm that the slow decay in the autocorrelation signal is dominated by the photo-thermal saturation effect.

To have a quantitative understanding of the photocurrent dynamics, a phenomenological model is developed to model the joint response of the device temperature evolution (thermal pulse) and transient photocurrent (photocurrent pulse). In this model, each VIS-pulse generates two concurrent photocurrent pulses in the device with independent decay constants \( I(t) = I_1 e^{-t/\tau_{1}} + I_2 e^{-t/\tau_{2}} \). Upon overlapping with the thermal pulse, the instantaneous photocurrent is reduced to \( \alpha I \), where \( \alpha \) is a dimensionless saturation factor and its temperature dependence is defined in the Supplementary Materials. As the device temperature is a function of time, \( \alpha \) is consequently time dependent. After calculating the evolution of the transient lattice temperature of the device excited by an IR-pulse at a delay of \( \Delta \tau \) through a two-temperature model (Supplementary Materials), we can obtain the average photocurrent through the cross-correlation relation \( T(\Delta \tau) = \frac{1}{t_{\text{rep}}} \int_{0}^{t_{\text{rep}}} \alpha(t) I(t+\Delta \tau) dt \), where \( I \) is the average photocurrent and \( t_{\text{rep}} \) is the repetition time of the laser pulse, which is much longer than the photocurrent response time. The calculated \( I \) fits our experimental results well (Fig. 3B), with the transient photocurrent profile \( I(t) \) presented in Fig. 3C. The fast decay in the early stage has a time constant \( t_1 \) of approximately 2 ps, which is similar to the graphene/TMD charge-transfer time. The measured time constant is nearly independent of the power of IR-pulse or VIS-pulse. This characteristic time of the transient photocurrent approaches that of the charge transfer at the graphene-TMD interface [9–13].
suggesting a likely faster interlayer charge-transfer process at the MoS₂ interface, similar to those observed in TMD heterostructures (9). On the other hand, the slow component has a time constant $t_2$ of approximately 25 ps. It was previously found that the two-component photocurrent can originate from defect-related processes (30, 32, 39), which agrees with the dynamics we observe.

To further confirm the electronic and thermal contributions in the photocurrent dynamics, a comparison is made between the BL and 4L regions with the pump-probe photocurrent measurements. Similar to the BL region, an asymmetric time dependence before and after zero delay as well as a sharp drop in photocurrent around zero delay are observed in the 4L region (Fig. 4A, isolated curve in fig. S8), indicating a similarly fast dynamics at the graphene-MoS₂ interface. Nevertheless, the pump-probe photocurrent signal before zero delay is much more prominent in the 4L than BL region, suggesting a much larger slow component contribution in the thick area. This contrast might result from a larger defect density in 2D or the extra MoS₂-MoS₂ interfaces in the 4L region. An external bias can be applied to tune the slow component of the photocurrent, which is observed before zero delay (Fig. 4A). As fitted by the aforementioned model, the slow component becomes faster when the electric field is along the photocurrent direction and vice versa (Fig. 4B). On the other hand, the recovery time at positive delay is dominated by the device cooling dynamics and is therefore largely bias independent. These results further confirm the thermal saturation nature and the existence of a two-component photocurrent response in the 3R-MoS₂ photovoltaic device.

In conclusion, we have observed a large temperature dependence in the spontaneous photovoltaic effect in heterostructures comprising atomically thin 3R-MoS₂ and graphene. At low temperatures, the EQE can reach 10% even in the thinnest area with only two atomic layers of MoS₂. Upon laser heating, such a temperature dependence leads to a strong thermal saturation of the photocurrent, which dominates in the commonly used photocurrent autocorrelation measurement and obscures the photocurrent dynamics. In addition, we have developed a nondegenerate pump-probe photocurrent spectroscopy technique, which can distinguish the electronic and thermal dynamics. With this measurement technique, we find that the transient photocurrent has two contributions with distinct temporal responses. We quantify the fast photocurrent response to be on the picosecond level, which is similar to the charge-transfer process at the graphene-TMD interface, suggesting an intrinsic device bandwidth of hundreds of gigahertz. At RT, the photoresponsivity and speed of our homobilayer device are comparable to those of TMD heterobilayer devices with type II band alignment (9, 40, 41). Our results may stimulate future uses of ferroelectric van der Waals devices in optoelectronic applications that require high performance with low power consumption and built-in memory function, such as in high-speed optical communications and optical computing. Furthermore, the nondegenerate pump-probe photocurrent spectroscopy technique can also be applied to study the interplay between various photo-excitation mechanisms in other types of photosensors.

**MATERIALS AND METHODS**

**Sample fabrication**

The 3R-MoS₂ flakes are exfoliated from a bulk crystal (HQ Graphene) on SiO₂/Si substrates. Graphene electrodes are cut into specific shapes with a femtosecond laser. The flake thickness is identified by an optical microscope and reflection contrast spectroscopy. The 3R phase is confirmed by second-harmonic generation measurement (42). Exfoliated hexagonal boron nitride (hBN) is used for encapsulation. All devices are fabricated using dry transfer method under ambient conditions with a homebuilt transfer stage. Electrical contact is achieved by overlapping the graphene with gold electrodes prepatterned by optical lithography on heavily p-doped Si/SiO₂ substrates.

**Photocurrent measurement**

For the autocorrelation measurements, a homebuilt wavelength-tunable optical parametric oscillator (OPO) (76-MHz repetition rate, 50-fs pulse width) is used to illuminate the sample, which sits in a continuous-flow optical cryostat (Oxford Microstat-He), through a long working distance 100× objective lens (Mitutoyo, numerical aperture = 0.5). The signal is detected through a lock-in amplifier phase-locked to a mechanical chopper. In the pump-probe photocurrent measurement, we split some light from the pump laser (light conversion, 1030 nm) of the OPO and then combine it with the OPO beam for collinear measurements. The photocurrent is measured by modulating the VIS-pulse from the OPO. For wavelength-dependent measurements, a supercontinuum white light source (YSL) (~100-ps pulse width, 10 MHz) is wavelength-selected by an acousto-optic tunable filter to illuminate the

![Fig. 4. Pump-probe photocurrent measurement in the 4L region. (A) Bias-dependent pump-probe photocurrent measurement in the 4L region with bias voltages ranging from −0.25 to 0.5 V. The measurements are carried out with an IR-pulse(VIS-pulse) fluence of approximately 350(2) μJ cm⁻². (B) Numerical modeling results with the response time of the slow component varying from 35 to 50 ps.](image-url)
device. For device photocurrent mapping, current-voltage characteristics, and temperature dependence, a 532-nm continuous-wave laser is used.

**Supplementary Materials**

This PDF file includes:

- Figs. S1 to S14

**REFERENCES AND NOTES**

1. J. Sung, Y. Zhou, G. Scuri, V. Zólyomi, T. I. Andersen, H. Yoo, D. S. Wild, A. Y. Joe, R. J. Gelly, H. Heo, S. J. Magorrian, D. Bérué, A. M. M. Valdivia, T. Taniguchi, K. Watanabe, M. D. Lukin, P. Kim, Y. I. Fal’ko, H. Park, Broken mirror symmetry in excitonic response of reconstructed domains in twisted MoS$_2$/MoS$_2$ bilayers. Nat. Nanotechnol. 15, 750–754 (2020).

2. X. Wang, K. Yasuda, Y. Zhang, S. Liu, K. Watanabe, T. Taniguchi, J. Hone, L. Fu, P. Jarillo-Herrero, Interfacial ferroelectricity in rhombohedral-bilayer transition metal dichalcogenides. Nat. Nanotechnol. 17, 367–371 (2022).

3. M. Vizner Stern, Y. Waschitz, W. Cao, I. Nevo, K. Watanabe, T. Taniguchi, E. Sela, M. Urbakh, O. Hod, M. Ben Shalom, Interfacial ferroelectricity by van der Waals sliding. Science 372, 1462–1466 (2022).

4. M. Wu, J. Li, Sliding ferroelectricity in 2D van der Waals materials: Related physics and future opportunities. Proc. Natl. Acad. Sci. U.S.A. 118, e211573118 (2021).

5. A. Weston, E. G. Castanon, V. Enalidiev, F. Ferreira, S. Bhattacharjee, S. Xu, H. Corte-León, Z. Wu, N. Clark, A. Summerfield, T. Hashimoto, Y. Gao, W. Wang, M. Hamer, H. Read, L. Fumagalli, A. V. Kretinin, S. J. Higah, O. Kazakova, A. K. Geim, V. I. Fal’ko, R. Gorbachev, Interfacial ferroelectricity in marginally twisted 2D semiconductors. Nat. Nanotechnol. 17, 390–395 (2022).

6. J. Liang, D. Yang, J. Wu, J. I. Dadap, K. Watanabe, T. Taniguchi, Z. Ye, Optically probing the asymmetric interlayer coupling in rhombohedral-stacked MoS$_2$ bilayer. Phys. Rev. X 12, 041005 (2022).

7. D. Yang, J. Wu, B. T. Zhou, J. Liang, T. Ideue, T. Siu, K. M. Awan, K. Watanabe, T. Taniguchi, Y. Iwasa, M. Franz, Z. Ye, Spontaneous-polarization-induced photovoltaic effect in hombohedrally stacked MoS$_2$. Nat. Photon. 16, 469–474 (2022).

8. T. Akamatsu, T. Ideue, L. Zhou, Y. Dong, S. Kitamura, M. Yoshii, D. Yang, M. Ona, Y. Nakagawa, K. Watanabe, T. Taniguchi, J. Launzeno, J. Huang, Z. Ye, T. Morimoto, H. Yuan, Y. Iwasa, A van der Waals interface that creates in-plane polarization and a spontaneous photovoltaic effect. Science 372, 68–72 (2021).

9. X. Hong, J. Kim, S.-F. Shi, Y. Zhang, C. Jin, Y. Sun, S. Tongay, J. Wu, Y. Zhang, F. Wang, Ultrafast charge transfer in atomically thin MoS$_2$/WS$_2$ heterostructures. Nat. Nanotechnol. 9, 682–686 (2014).

10. L. Yuan, T.-F. Chung, A. Kuc, Y. Wan, Y. Xu, Y. P. Chen, T. Heine, L. Huang, Photocarrier generation from interlayer charge-transfer transitions in WS$_2$-graphene heterostructures. Sci. Adv. 4, e1703234 (2018).

11. X. Zhang, D. He, L. Yi, S. Zhao, J. He, Y. Wang, H. Zhao, Electron dynamics in MoS$_2$-graphite heterostructures. Nano Lett. 9, 14533–14537 (2019).

12. D. Luo, J. Tang, X. Shen, F. Ji, J. Yang, S. Weathersby, M. E. Kozina, Z. Chen, J. Xiao, Y. Ye, T. Cao, G. Zhang, X. Wang, A. M. Lindenberg, Twist-angle-dependent ultrafast charge transfer in MoS$_2$/WS$_2$ van der Waals heterostructures. Nano Lett. 21, 8051–8057 (2021).

13. Y. Chen, Y. Li, Y. Zhao, H. Zhou, H. Zhu, Highly efficient hot electron harvesting from graphene before electron-hole thermalization. Sci. Adv. 5, eaax9958 (2019).

14. C. Jin, E. Y. Ma, O. Karni, E. C. Regan, F. Wang, T. F. Heinz, Ultrafast dynamics in van der Waals heterostructures. Nat. Nanotechnol. 13, 994–1003 (2018).

15. J.-Y. Wu, Y. T. Chun, S. Li, T. Zhang, J. Wang, P. K. Shrestha, D. Chu, Broadband MoS$_2$ field-effect phototransistors: Ultrasmall visible-light photoreponse and negative infrared photoresponse. Adv. Mater. 30, 1705880 (2018).

16. W. Zhang, C.-P. Chiu, J.-K. Huang, C.-H. Chen, M.-L. Tsai, Y.-H. Chang, C.-T. Liang, Y.-Z. Chen, Y.-L. Chueh, J.-H. He, M.-Y. Chou, L.-J. Li, Ultrafast gain photodetectors based on atomically thin graphene-MoS$_2$ heterostructures. Sci. Rep. 4, 3826 (2014).

17. M. Buscema, M. Barkeld, V. Zwiller, H. S. van der Zant, G. A. Steele, A. Castellanos-Gomez, Large and tunable photovoltaic effect in single-layer MoS$_2$. Nano Lett. 13, 358–363 (2013).

18. D. J. Groenendijk, M. Buscema, G. A. Steele, S. Michaelis de Vasconcellos, R. Batitschatsch, H. S. van der Zant, A. Castellanos-Gomez, Photovoltaic and photothermoelectric effect in a double-gated WSe$_2$ device. Nano Lett. 14, 5846–5852 (2014).
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