Unusually efficient photocurrent extraction in monolayer van der Waals heterostructure by tunnelling through discretized barriers

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Two-dimensional layered transition-metal dichalcogenides have attracted considerable interest for their unique layer-number-dependent properties. In particular, vertical integration of these two-dimensional crystals to form van der Waals heterostructures can open up a new dimension for the design of functional electronic and optoelectronic devices. Here we report the layer-number-dependent photocurrent generation in graphene/MoS\textsubscript{2}/graphene heterostructures by creating a device with two distinct regions containing one-layer and seven-layer MoS\textsubscript{2} to exclude other extrinsic factors. Photoresponse studies reveal that photoresponsivity in one-layer MoS\textsubscript{2} is surprisingly higher than that in seven-layer MoS\textsubscript{2} by seven times. Spectral-dependent studies further show that the internal quantum efficiency in one-layer MoS\textsubscript{2} can reach a maximum of 65\%\textsuperscript{,} far higher than the 7\% in seven-layer MoS\textsubscript{2}. Our theoretical modelling shows that asymmetric potential barriers in the top and bottom interfaces of the graphene/one-layer MoS\textsubscript{2}/graphene heterojunction enable asymmetric carrier tunnelling, to generate usually high photoresponsivity in one-layer MoS\textsubscript{2} device.
The two-dimensional (2D) layered materials such as graphene and semiconducting transition-metal dichalcogenides (TMDs) can exhibit peculiar electronic properties depending on their exact composition, thickness and geometry\(^1\). Furthermore, the broad possibility to combine different materials in van der Waals heterostructures (vdWHs) can create a new paradigm in materials science with unprecedented flexibility to integrate highly disparate materials and enable unique functions, as exemplified by the recent demonstration of vertical tunnelling transistors and vertical field-effect transistors for ultra-thin and flexible devices\(^2\)–\(^13\). Graphene/multi-layer (ML)-TMD/graphene stack has also been shown to function as a unique photodiode for photocurrent generation or photodetection\(^7\)–\(^12\). With a direct band gap in monolayer TMD and ultra-fast carrier transfer (~1 ps) rate\(^14\), vdWHs with ultra-thin TMDs have attracted considerable interest for photovoltaic applications\(^15\)–\(^18\) and ultra-fast photodetection\(^19\). However, classical charge transport theory for bulk semiconductors was often employed to interpret the electron transport in ultra-thin vdWHs even though it is well expected to exhibit completely different photocurrent generation characteristics\(^7\)–\(^12\). Notably, an atomically thin graphene/WSe\(_2\)/MoS\(_2\)/graphene \(p–n\) heterojunction has been investigated for photocurrent generation. However, Shockley–Read–Hall recombination and Langevin recombination in the WSe\(_2\)/MoS\(_2\) \(p–n\) junction could compromise the carrier extraction performance\(^1\). The unique characteristics of ultrathin vdWHs are insufficiently explored to date. In particular, a systematic investigation of layer-number-dependent studies is lacking due to complications from highly variable nature of the van der Waals interfaces and extrinsic factors in creating vdWH devices.

Here we report a layer-number-dependent photocurrent generation in graphene/MoS\(_2\)/graphene vdWHs by creating a device with two distinct regions containing one-layer (1L) and seven-layer (7L)-MoS\(_2\) to exclude extrinsic device factors. Significantly, we discover a surprisingly higher responsivity and internal quantum efficiency (IQE) by going from ML MoS\(_2\) device to a monolayer device. The discretized electrostatic potential barriers were introduced to interpret the photocarrier tunneling and extracting in ultrathin vdWHs.

### Results

**Photocurrent generation in graphene/1L-MoS\(_2\)/graphene stack.** Figure 1 shows the schematic illustration of graphene/1L-MoS\(_2\)/graphene heterostructure device on Si/SiO\(_2\) substrate. A focused laser was used to generate electron-hole pairs in the MoS\(_2\) layer, which can be separated by the asymmetric potential between the top graphene (Gr\(_T\))/MoS\(_2\) and the bottom graphene (Gr\(_B\))/MoS\(_2\) junction to produce photocurrent (Fig. 1a). Figure 1b shows an optical image of a typical graphene/1L-MoS\(_2\)/graphene vdWH device. The 8 \(\mu\)m strip of Gr\(_B\) is located below 1L-MoS\(_2\) flake (inside the dotted line) and the Gr\(_T\) layer is located directly above the MoS\(_2\) flake (inside the solid line), to overlap with MoS\(_2\) flake and the Gr\(_B\). The number of layers of MoS\(_2\) was then confirmed by Raman spectra measurements (Supplementary Fig. 1) and atomic force microscopic measurement. Figure 2b display the photoluminescence (PL) mapping image of 1L-MoS\(_2\) and 7L-MoS\(_2\) portion in Fig. 2a under an excitation wavelength of 532 nm (2.33 eV). The monolayer-MoS\(_2\) exhibited a much stronger PL compared with 7L-MoS\(_2\), which is consistent with the direct band gap nature of 1L-MoS\(_2\) (band gap ~1.82 eV)\(^1\)–\(^4\) and the indirect band gap nature of 7L-MoS\(_2\) (band gap ~1.3 eV)\(^1\)–\(^6\).

Two prominent PL peaks can be identified at 630 nm (1.96 eV) and 532 nm (2.33 eV). The monolayer-MoS\(_2\) exhibited a much stronger PL compared with 7L-MoS\(_2\), which is consistent with the direct band gap nature of 1L-MoS\(_2\) (band gap ~1.82 eV)\(^1\)–\(^4\) and the indirect band gap nature of 7L-MoS\(_2\) (band gap ~1.3 eV)\(^1\)–\(^6\).

The photocurrent map of the entire device clearly demonstrates that 1L-MoS\(_2\) region exhibited much more pronounced photocurrent than the 7L-MoS\(_2\) region (Fig. 2c). The current–voltage \((I_d–V_{ds})\) data obtained in the dark (black line, Fig. 3a) and under 514 nm laser irradiation (red line for 1L-MoS\(_2\) region and blue line for 7L-MoS\(_2\) region; Fig. 3a) show a clear photocurrent response in the vdWH. The open-circuit voltage and short-circuit current obtained in the 1L-MoS\(_2\) region are 110 mV and 80 \(\mu\)A, respectively. This is a rather surprising and counter-intuitive discovery, considering that the 7L-MoS\(_2\) should have significantly higher optical absorption than 1L-MoS\(_2\) (Fig. 3b and Supplementary Note 1).

The enhanced photoresponsivity of 1L-MoS\(_2\) may be attributed to two potential factors: higher photocarrier generation rate in the monolayer MoS\(_2\) and higher photocarrier extraction collection efficiency. We will discuss the influence of each factor below, first by modelling the transition probability and subsequently by

\[
I_{\text{DT}} = \frac{A_{\text{eff}} \sqrt{m_p q d^2 V_{ds}}}{h \epsilon^2} \exp \left[ -4 \sqrt{m_p q d^2 V_{ds}} \right]
\]  

where \(A_{\text{eff}}, \phi_B, q, m_p, d, h\) and \(\epsilon\) are effective contact area, barrier height, electron charge, free electron mass, effective electron mass, barrier width (MoS\(_2\) thickness) and Plank’s constant, respectively.\(^1\)

Based on the photocurrent response and input laser power, we can determine the photoresponsivity (\(A W^{-1}\)) of the device. The photoresponsivity of 1L-MoS\(_2\) increases with decreasing the laser power could be attributed partly to absorption saturation in MoS\(_2\) and partly to the screening of a built-in electric field by the excited electrons in the conduction band of MoS\(_2\) (ref. 7). Importantly, the photoresponsivity of 1L-MoS\(_2\) increases with increasing the laser wavelength and a maximum photoresponsivity of 68 mA W\(^{-1}\) in 1L-MoS\(_2\) vdWHs was achieved at the wavelength of 633 nm and the laser power of 100 nW (Fig. 1e), which exceeds that of previous studies on graphene/ML-MoS\(_2\)/graphene vdWHs device (~30 mA W\(^{-1}\) without plasmonic enhancement)\(^7\)–\(^12\). This performance difference is attributed to the number of layers difference between this work (monolayer MoS\(_2\)) and previous reports (5 to 30 nm MoS\(_2\)), which will be further discussed by comparison of external quantum efficiency (EQE) and IQE between 1L-MoS\(_2\) and 7L-MoS\(_2\) vdWHs in Fig. 2gh.
comparing the internal and external quantum efficiencies. These two lines of reasoning will show that at peak IQE (633 nm and 1.96 eV), the second factor (that is, the improved collection) dominates in our device.

In general, the spectral dependence of the photocarrier generation rate can be estimated from first principles, using the Fermi golden rule. The probability of transition from a valence band state ($v$) to a conduction band state ($c$) is thus given (in 2D) by

$$W(\omega) = k_2(\omega) \frac{m_e^2 \omega^2}{2\pi q^2 \hbar}$$

where $k_2$ is the imaginary part of the dielectric function, $\hbar$ is the plank constant, $\omega$ is the frequency of the incident radiation, $m_e$ and $q$ are the electron mass and charge. In MoS$_2$, $k_2$ diverges because the conduction and valence bands run nearly parallel in some regions of the reciprocal space (that is, band nesting). This can be noticed in Fig. 3c where we compare the imaginary part of the dielectric function, obtained from relativistic first-principles calculations, for 1L and thick multilayer (bulk) MoS$_2$. Thus, $W$ increases with the excitation energy up to the energy of the first peak, which is at ~2.4 eV for the bulk and 2.8 eV for the monolayer, respectively. The effect of band-nesting is more pronounced in monolayer, because this system is closer to the 2D limit and is
Figure 3 | Quantum efficiency between monolayer and ML MoS₂ heterostructures. (a) $I_d-V_{ds}$ characteristics of the device under laser illumination of 514 nm laser (red line for 1L-MoS₂, blue line for 7L-MoS₂) and dark (black line). (b) Absorption spectra of a 7L-MoS₂ flake (red line) and 1L-MoS₂ flake (black line). (c) Calculated imaginary part of the dielectric function ($\varepsilon_2$) of 1L- and bulk MoS₂ (as a model for thick ML). The effective thickness for the 1L-MoS₂ was taken to be the inter-layer separation in ML-MoS₂. (d) EQE of 1L-MoS₂ stack and 7L-MoS₂ stack at a laser power of 100 µW. (e) Calculated density of states of 1L-MoS₂ and ML-MoS₂.

Photocarrier extraction mechanism. Photocarrier transport in bulk MoS₂ is driven by diffusion and drift process, which is dictated by different band bending at GrT and GrB MoS₂ interface (originated from the difference of GrT and GrB doping densities)⁷,¹² (Supplementary Fig. 3a). The photo-carrier extraction in atomically thin vdWHs is predicted to be completely different from that in bulk heterojunctions. Unlike bulk MoS₂ having a continuous band bending (Supplementary Fig. 3a), atomically thin vdWH has discrete energy states in the vertical direction (Supplementary Fig. 3b). The classical semiconductor charge transport model—the charge drifts downhill following the band slope—cannot describe photocurrent generation phenomena via such discrete energy states in vdW heterojunctions. To this end, we introduce a tunnelling transport model to describe the photocarrier tunnelling through electrostatic potential barriers formed at the atomically thin vdW heterojunctions. Electrostatic potential barriers of 1L-MoS₂ heterostructures are calculated by density functional theory (DFT; Supplementary Fig. 4 and Supplementary Note 2). Large electrostatic potential barriers are discretely constructed between GrT/GrB and MoS₂ interface. The barrier heights at the top and bottom junctions are symmetric because of symmetric environmental conditions imposed in the DFT calculations. In real device, the different environmental conditions of GrT and GrB result in asymmetric doping between GrT and GrB. Doping characteristics of GrT and GrB were confirmed by 2D peak shift in Raman spectrum of another graphene/1L-MoS₂/graphene heterostructure device (Supplementary Fig. 5 and Supplementary Note 3). With $n$- or $p$-type doping, the 2D Raman band exhibits a red or blue shift, respectively. In our device, the GrT is more $p$-type doped than GrB. Such doping difference generates internal field, leading to asymmetric barrier height between top and bottom junctions (Supplementary Fig. 6). In this way, the photocarrier transport is described by the tunnelling model.

Further insight is gained by the IQE, which can be obtained by dividing the EQE by the optical absorbance of 7L- and 1L-MoS₂ (Fig. 3e). Importantly, the maximum IQE in 1L-MoS₂ can reach up to 65%, far exceeding to IQE in 7L-MoS₂ (~7%). It is also noted that the IQE in 1L-MoS₂ region steadily increases when the photon energy is reduced to approach the bandgap of 1L-MoS₂ (1.82 eV (ref. 15)). The highest IQE reached ~65% at 633 nm (1.96 eV), very close to the optical absorption edge of 1L-MoS₂ (1.82 eV). It is noteworthy that in this region $k_2$ is nearly independent on the thickness. According to the theoretical calculations, 1L-MoS₂ has a much larger density of states near the band edges, at the valence band top due to the heavier hole masses, and near the conduction band edge, due to the presence of the 2D van-Hove singularities (Fig. 3f), increasing the probability of transfer to the graphene electrodes.
Furthermore, the different dielectric environment at the Gr\textsubscript{T} and Gr\textsubscript{B}/MoS\textsubscript{2} interface can significantly alter the electrostatic potential. Figure 4a shows the schematic illustration of the dielectric environment of the device. The Gr\textsubscript{B} is sandwiched between SiO\textsubscript{2} substrate and 1L-MoS\textsubscript{2}, and the Gr\textsubscript{T} between 1L-MoS\textsubscript{2} and air. The dielectric constant of Gr\textsubscript{B} (4.2) and Gr\textsubscript{T} (2.7) are determined by the standard approximation of $\varepsilon_{\text{GrB}} = (\varepsilon_{1L-MoS_2} + \varepsilon_{\text{SiO}_2})/2 = 4.2$ and $\varepsilon_{\text{GrT}} = (\varepsilon_{1L-MoS_2} + \varepsilon_{\text{air}})/2 = 2.7$, respectively\textsuperscript{25}.\textsuperscript{25}

The calculated electrostatic potentials with these dielectric constants are shown in Fig. 4b,c (calculation is shown in Supplementary Note 3). Green coloured areas are the potential energy barriers. $E_\text{F}$ is the Fermi level of graphene. CB, VB and $E_\text{F}$ are conduction band, valence band and band gap of MoS\textsubscript{2}, respectively.

For partially confined at the $T\text{B}$ (Fig. 4d–ii), the barrier at the bottom junction is higher than the kinetic energy of excited electron. The excited electrons transfer to Gr\textsubscript{T} over the barrier with a tunnelling probability of $(2k_{\text{MoS}_2}/\hbar^2) \times \exp(-2\sqrt{2m^*/\hbar^2}a)$, where $k_{\text{MoS}_2} = \sqrt{2m^*(E - E_\text{F})/\hbar^2}$, $a$ is energy barrier width, $m^*$ is electron effective mass, $U$ is electrostatic barrier, $E$ is kinetic energy of electron and $\hbar$ is the Planck’s constant. The top junction allows for higher tunnelling probability due to smaller barrier height compared with the bottom junction. Consequently, electrons can transfer to the Gr\textsubscript{T} with positive $I$. The photocarrier extraction rate ($I$) is not altered appreciably in response to the energy of excited electrons (or absorbed photon energy), because effective barrier height $(U-E)$ of the bottom and top junction changes by the similar magnitude (Fig. 4e–i).

For fully confined at $T\text{B}$ and $T\text{T}$ (Fig. 4d–i), the excited electrons in MoS\textsubscript{2} can be transported to Gr\textsubscript{T} and Gr\textsubscript{B} by tunnelling through both sides of the potential barriers. The tunnelling probabilities through $T\text{B}$ or $T\text{T}$ is defined to exp$(-2\sqrt{2m^*/\hbar^2}a)$ (ref. 26), where $a$ is energy barrier width, $m^*$ is electron effective mass, $U$ is electrostatic barrier, $E$ is kinetic energy of electron and $\hbar$ is the Planck’s constant. The photocarrier extraction rate ($I$) decreases in proportion to electron excitation energy (Fig. 4e–ii).
**MoS₂ layer-dependent electrostatic potentials.** The electrostatic potentials were calculated in terms of the number of MoS₂ layers (Fig. 5). The region of fully confined between \( T \) and \( T_B \) case is expanded in response to the number of MoS₂ layers due to the increased barrier heights at both sides. On the other hand, the region of partially confined at the \( T_B \) case is shrunken when the number of MoS₂ layers increases due to the reduction of barrier height difference at both junctions. The region of no confinement case is also shrunken by increased barrier heights at both sides. Electrostatic potential is saturated over the 7L-MoS₂ and behaves as bulk MoS₂. The photocarrier extraction rates (\( I \)) were calculated in terms of the electrostatic potentials.

**Calculated photocarrier extraction rate.** The carrier lifetime in direct gap (1L-MoS₂) is shorter than that of indirect gap (7L-MoS₂), which should be taken into account in recombination process during carrier drift. We have measured the carrier lifetime in direct gap (1L-MoS₂) and indirect gap (7L-MoS₂) based on transmission change in pump–probe measurement (Fig. 6a) and then calculated the normalized excited carriers (\( \delta n_e \)) by initial excited carriers (\( \delta n_0 \)) for 1L-MoS₂ (direct gap, red line) and ML-MoS₂ (indirect gap, black line) along with carrier drift distance (nm; Fig. 6b and Supplementary Note 5). The normalized excited carrier density was then multiplied with photocarrier extraction rate (\( I \)) to clarify recombination difference in ML-MoS₂. The 1L-MoS₂ shows ultra-fast carrier transfer (~1 ps) rate into graphene layer, much shorter than carrier lifetime in 1L-MoS₂ (60 ps), resulting in highly efficient transfer of photoexcited carriers in MoS₂ into graphene before recombination. Therefore, we multiplied photocarrier extraction rate (\( I \)) in 1L-MoS₂ by 100% of normalized excited carrier density. Figure 6c clearly reveals that the photocarrier extraction rate (\( I \)) reaches around 65% when the number of MoS₂ layers is reduced from 7L-MoS₂ to 1L-MoS₂ and the photon energy approaches the bandgap (1.82 eV). The calculated \( I \) are in good agreements with measured IQE of 1L-MoS₂ (black dashed line in Fig. 6d) and 7L-MoS₂ heterostructures (red dashed line in Fig. 6d).

**I-V characteristics.** Previous studies have demonstrated an external gate voltage can be used to effectively modulate the
Fermi energy of graphene and thus the potential offset and the driving force for carrier separation in graphene/MoS2/graphene devices\(^7,12\). We have also investigated gate voltage dependency of dark current (Supplementary Fig. 8) and photocurrent generation (Fig. 7) in the 1L-MoS2 and ML MoS2 heterostructures. Overall, the 1L-MoS2 device showed considerably smaller modulation of short circuit current \((I_{sc})\) and open circuit voltage \((V_{oc})\) by an external gate voltage (Fig. 7a) than that of ML-MoS2 (Fig. 7b). To understand the weak gate modulation in graphene/1L-MoS2/graphene device, the band diagram of the 1L-MoS2 and ML-MoS2 devices at the positive and negative gate voltage are calculated (Supplementary Fig. 9 and Supplementary Note 6) and schematically illustrated in Fig. 7c,d. It is noted that the electrostatic screening of the GrB layer is dependent on its carrier concentration, which can be varied by applied gate voltage. This factor is considered in the simulation and schematics. In general, the gate voltage can more efficiently modulate the GrB \(E_F\) than GrT \(E_F\) due to the partial screening of the gate field by the GrB layer and sandwiched MoS2 layers. The larger gate modulation of the \(T_B\) than \(T_C\) can create additional potential offset or band slope that leads to change in \(V_{oc}\). In the ML-MoS2 device, the gate electric field primarily modulates the Fermi level \((E_F)\) of the GrB due to the relatively strong electrostatic screening by ML-MoS2 (Fig. 7d), leading to a large modulation of the band slope in ML-MoS2 and thus a substantial modulation of the \(V_{oc}\) and photocurrent. In contrast for the monolayer device, the gate electric field can effectively modulate both bottom and GrT \(E_F\) because of much weaker screening effect by the ultra-thin 1L-MoS2 (Fig. 7c). As a result, both top and bottom potential barrier are modulated together in the graphene/1L-MoS2/graphene device, leading to much smaller dependence of potential offset and photocarrier extraction rate on the gate voltage.

**Discussion**

In summary, we have reported an unusually efficient photocarrier generation from monolayer MoS2 in graphene/MoS2/graphene vdWHs. Owing to intrinsically slow layer-to-layer charge transport in TMD materials, the carrier mobility in the vertical direction is usually several orders of magnitudes lower than that in lateral direction, which can seriously slow down the charge separation process in the vertical heterostructures, leading to undesired photocarrier recombination in MoS2 before they arrive at the current collector. Importantly, we show that this drawback can be largely overcome by using atomically thin MoS2 to enable an unusually high photoresponsivity up to 68 mA W\(^{-1}\) with 65% of IQE in monolayer MoS2. This was congruent with carrier tunnelling through asymmetric electrostatic potential barriers that exist discretely in vdWHs with monolayer MoS2. Conversely, the symmetric potential barrier in ML MoS2 reduces photocurrent via carrier tunnelling. The proposed quantum mechanical-based tunnelling mechanism provides a new theoretical framework to understand van der Waals interaction in vdWHs and to design the next generation of atomically thin optoelectronics including photodetectors and photovoltaic devices.

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**Figure 6 | Calculated photocarrier extraction rate of vdWHs with various number of MoS2 layers.** (a) Normalized dynamics of differential transmission \((\Delta T/T_0)\) of 1L-MoS2 and 7L-MoS2 for pump fluence (dots) and fitting lines. \(\Delta T\) is the pump-induced probe transmission change and \(T_0\) is the probe transmission without pump excitation for pump fluence. The decay lifetime of 1L-MoS2 \((\tau_{1\text{L-MoS2}})\) and 7L-MoS2 \((\tau_{7\text{L-MoS2}})\) obtained from the fitting. (b) Normalized excited electron carriers \((\Delta n_e)\) by initial excited electron carriers \((\Delta n_0)\) for direct gap MoS2 (black line) and indirect gap MoS2 (red line) along with carrier drift distance (nm). (c) 2D colour plot of photocarrier extraction rate as a function of the number of MoS2 layers and photon excitation energy. (d) Photon energy dependence of IQE (black dots: 1L-MoS2; red dots: 7L-MoS2) and calculated photocarrier extraction rate (black dashed line: 1L-MoS2; red dashed line: 7L-MoS2).
and (MoS$_2$), the IQE in our device is about twice higher. The lower approach27. The GrT electrode was transferred and patterned on the MoS$_2$ flake. The fabrication of the vertical heterostructures was carried out using a photolithography and oxygen plasma etching process. The MoS$_2$ flakes were then transferred onto the graphene strips through a dry transfer (300 nm SiO$_2$) substrate, and patterned into 8 strips as the bottom electrode. The metal electrode, for probe contact and Langevin recombination, which is not present in graphene/ML-MoS$_2$/graphene devices.

It is noteworthy that the 2.5% EQE achieved in our graphene/1L-MoS$_2$/graphene devices are comparable to the 2.4% value achieved in graphene/1L-WSe$_2$/1L-MoS$_2$/graphene p–n heterojunctions. Considering the absorbance in two layers stack (MoS$_2$/WSe$_2$) is approximately twice of that in single layer (MoS$_2$), the IQE in our device is about twice higher. The lower IQE in graphene/1L-WSe$_2$/1L-MoS$_2$/graphene p–n heterojunctions may be partially attributed to Shockley–Read–Hall recombination and Langasite recombination, which is not present in graphene/1L-MoS$_2$/graphene devices.

**Methods**

**The fabrication of the vertical heterostructures.** For the fabrication of the vertical heterostructures of graphene/MoS$_2$/graphene device, the graphene was grown with a chemical vapour deposition process27,28 and transferred onto Si/SiO$_2$ (300 nm SiO$_2$) substrate, and patterned into 8 × 30 μm strips as the bottom electrode using a photolithography and oxygen plasma etching process. The MoS$_2$ flakes were then transferred onto the graphene strips through a dry transfer approach27. The Gr$_B$ electrode was transferred and patterned on the MoS$_2$ flake and Gr$_B$. Directly overlapping graphene area was patterned and etched away, to avoid short circuit between the Gr$_T$ and Gr$_B$. The metal electrode, for probe contact or wire bonding purposes, was patterned on the bottom and Gr$_T$ electrodes by e-beam lithography followed by e-beam deposition of Ti/Au (50/50 nm).

**Microscopic and electrical characterization.** Electrical transport measurements were conducted with a probe station and a computer-controlled analogue to-digital converter at room temperature. The scanning photocurrent measurements were conducted with the same electrical measurement system under a SP2 MP Film confocal microscope coupled with an Ar/ArKr laser (wavelength 458, 476, 488, 496 and 514 nm) and HeNe laser (543, 596 and 633 nm). All optical measurements were carried under ambient conditions at room temperature by using an inverted microscope coupled to a grating spectrometer with a charge-coupled device camera. The optical beams were focused on the sample with a spot diameter of ~1 μm. MoS$_2$ heterostructures were excited with a cw solid-state laser at a wavelength of 405 and 532 nm. A low laser power of ~100 μW was used to avoid heating and PL saturation. The PL of MoS$_2$ samples were calibrated by using rhodamine 6G molecules as the standard. Raman spectroscopy (RM1000 microprobe; Renishaw) was used to characterize the MoS$_2$ and graphene with a wavelength of 514 nm (2.41 eV) and a Rayleigh line rejection filter. As Raman scattering efficiency has usually little layer thickness dependence, the PL spectra normalized by Raman intensity reflects directly the luminescence efficiency16.

**The DFT calculations.** The DFT calculations were performed using the Quantum ESPRESSO code29. Geometry optimizations and total energy calculations are non-relativistic. Bandstructures calculated using a fully relativistic formalism are obtained using the atomic coordinates optimized in a similar non-relativistic calculation. The core electrons were described by norm-conserving, full relativistic pseudopotentials28 with nonlinear core-correction and spin–orbit information. The exchange correlation energy was described by the generalized gradient approximation, in the scheme proposed by Bueke et al30. The energy cutoff was 50 Ry.

The monolayer was modelled using a supercell consisting of 4 × 4 unit cells of the monolayer material, comprising a total number of 48 atoms. The lattice parameter (α) used was optimized for the primitive unit cell. The supercell length along the direction perpendicular to the plane c is taken to be twice the lattice parameter, except for preliminary bandstructure calculations performed using a smaller c, which are identified in the text. The Brillouin zone was sampled using a 2 × 2 × 1 Monkhorst–Pack grid31.

The 7L material was modelled as a bulk crystal, using a 4 × 4 × 1 supercell. The in-plane lattice parameter was taken from the monolayer calculation and the c/a ratio was taken from experimental data32.

**Ultrafast optical pump-probe spectroscopy.** The femtosecond laser source is based on a 250 kHz Ti:sapphire regenerative amplifier (Coherent RegA 9050), which provides a 1.55 eV photon energy, 50 fs pulse width and 6 μJ per laser pulse. The Reg A output pulse is separated into two pulses by a ratio of 7:3. The 70 % laser pulse is used to generate 3.1 eV pump–photon energy by second harmonic generation in a 1 mm-thick beta barium borate crystal. The other 30% laser beam is focused in an ultra-clean sapphire disk to generate a white-light super-continuum, which serves as a probe pulse. Both the pump and the probe pulses are focused on the ML-MoS$_2$ and 7L-MoS$_2$ samples by an objective lens (Mitutoyo M Plan Apo × 10) and the delay between the two pulses is controlled by a mechanical delay stage (Newport M-IMS300PP).

**Data availability.** The data that support the findings of this study are available from the corresponding author upon request.

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Author contributions
X.D. conceived the research and designed the experiment. W.J.Y. performed most of the experiments including device fabrication, characterization, data analysis and theoretical analysis. Q.A.V. and Y.H.L. stacked graphene-monolayer MoS₂-graphene by dry transfer process and performed Raman spectrum. H.O. and M.J. performed PL measurement. H.Z. synthesized the graphene. S.C., J.-Y.K. and H.C. performed optical pump–probe spectroscopy. A.C. and A.H.C-N. performed the DFT calculations. A.Y. performed absorption measurement. X.D., Y.H.L. and W.J.Y. wrote the paper. All authors discussed the results and commented on the manuscript.

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