High Photovoltaic Quantum Efficiency in Ultrathin van der Waals Heterostructures

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ABSTRACT: We report experimental measurements for ultrathin (<15 nm) van der Waals heterostructures exhibiting external quantum efficiencies exceeding 50% and show that these structures can achieve experimental absorbance >90%. By coupling electromagnetic simulations and experimental measurements, we show that pn WSe2/MoS2 heterojunctions with vertical carrier collection can have internal photocarrier collection efficiencies exceeding 70%

Keywords: van der Waals, heterojunction, photovoltaics, quantum efficiency, high efficiency, MoS2, WSe2

Owing to their naturally passivated basal planes and strong light−matter interactions, transition-metal dichalcogenides (TMDs) are of considerable interest as active elements of optoelectronic devices such as light-emitting devices, photodetectors, and photovoltaics.1,2 Ultrathin TMD photovoltaic devices a few atomic layers in thickness have been realized using TMDs such as molybdenum disulfide (MoS2) and tungsten diselenide (WSe2).3−8 Complete absorption of the solar spectrum is a challenge as the thickness is reduced to the ultrathin limit,3−11 whereas efficient carrier collection is challenging in thicker bulk TMD crystals. The active layers in conventional photovoltaics typically range from a few microns in direct gap materials (gallium arsenide) to a hundred microns thick or more in indirect gap materials (silicon).12 Efficient ultrathin and ultralight (<100 g/m2) photovoltaics have long been sought for many applications where weight and flexibility are important design considerations, such as applications in space power systems, internet-of-things devices, as well as portable and flexible electronics.13−15 Conventional photovoltaic materials are mechanically fragile when thinned down to the ultrathin (<10 nm) regime, and interfacial reactions mean that a large fraction of the crystal consists of surface-modified regions rather than intrinsically bulk material. Surface oxides and dangling bonds in ultrathin films often result in increased nonradiative recombination losses, lowering photovoltaic efficiencies. By contrast, TMDs have intrinsically high absorption and their layered crystallographic structures suggest the possibility of achieving intrinsically passive basal planes in high-quality crystals.

Photovoltaics that can approach the Shockley−Queisser limit16,17 have two prerequisites: First, at open circuit, every above-bandgap photon that is absorbed is extracted as an emitted photon at the band-edge of the material, i.e., it has perfect external radiative efficiency. Amani et al. have recently demonstrated that superacid-treated monolayers of MoS2 and WSe2 exhibit internal radiative efficiency >99%,19 suggesting that the condition of very high external radiative efficiency might be satisfied in TMDs. The second prerequisite is that at short circuit, the photovoltaic device must convert every incident above-bandgap photon into an extracted electron, i.e., it has external quantum efficiency (EQE) approaching unity.

To understand the path to high EQE, we can deconvolute the EQE into the product of two terms: the absorbance and internal quantum efficiency (IQE). High EQE devices exhibit both high absorption and IQE, i.e., carrier generation and collection efficiency per absorbed photon. To date, reports of van der Waals-based photovoltaic devices have not considered both of these concepts and separately evaluated them as criteria for high-efficiency photovoltaics.

Coupling electromagnetic simulations with absorption and EQE measurements enable quantitative characterization of few-atomic-layer thickness optoelectronic devices in van der Waals heterostructures. In this paper, we demonstrate EQEs >50% (Figure 1a), indicating that van der Waals heterostructures have

Received: May 6, 2017
Accepted: June 2, 2017
considerable potential for efficient photovoltaics. We show that high EQE results from both high optical absorption and efficient electronic charge carrier collection. We analyze the optical response using electromagnetic simulations to explain how near-unity absorption can be achieved in heterostructures (Figure 1b). We find that experimental absorption results for van der Waals heterostructures match well with these electromagnetic simulations. Thus, we can separate optical absorption and electronic transport to quantitatively compare their effects on charge collection efficiency for both pn heterojunctions and Schottky junctions (Figure 1c). In addition, we analyze the role of few-layer graphene (FLG) as a transparent top contact (Figure 1d). Finally, we outline important considerations for designing high-efficiency photovoltaic devices. By simultaneously maximizing both external radiative efficiency and EQE in a single device, van der Waals material-based photovoltaic devices could in principle achieve efficiencies close to the Shockley–Queisser limit for their bandgaps.

RESULTS AND DISCUSSION

Optoelectronic Device Characteristics. We analyzed the optoelectronic device characteristics of a high-performance device consisting of a vertical van der Waals heterostructure device of 0.6 nm-thick FLG/9 nm WSe$_2$/3 nm MoS$_2$/Au (see Supporting Information S1 for optical and photocurrent images). Its optoelectronic and device characteristics are shown in Figure 2. First, we find that this device exhibits an EQE > 50% (Figure 2a) with absorbance >90% from approximately 500 to 600 nm. Spectral features such as the exciton resonances of MoS$_2$ and WSe$_2$ are well reproduced in the EQE spectrum. In addition, we observe a maximum single-wavelength power conversion efficiency (PCE) of 3.4% under 740 W/cm$^2$ of 633 nm laser illumination (Figure 2b). Since the high-performance device is electrically in parallel with other devices, typical macroscopically large spot size (~cm) AM 1.5G illumination measurements would yield device characteristics substantially different from the high-performing one. Thus, we estimated the AM 1.5G performance using extracted device parameters of a diode fit under laser illumination (see Supporting Information S2 for details). We estimate the AM 1.5G PCE of this device to be ~0.4%. This value is presently too low to be useful for photovoltaics, but the high EQE values reported here indicate promise for high-efficiency devices, when device engineering efforts are able to also achieve correspondingly high open-circuit voltages in van der Waals-based photovoltaics.

Further measurements were performed at different laser powers under 633 nm laser illumination (Figure 3), yielding various power-dependent characteristics. Examination of the short-circuit current $I_{sc}$ yielded nearly linear dependence on laser power, as expected in ideal photovoltaic devices. The dashed blue line represents the fit to the expression $I_{sc} = AP^\tau$, where $A$ is a constant of proportionality, $P$ is the incident power, and $\tau$ represents the degree of nonlinearity in this device ($\tau = 1$ is the linear case). We find that $\tau = 0.98$ in our device, indicating nearly linear behavior under short-circuit conditions. In addition, in an ideal photovoltaic device, the open-circuit voltage is expected to grow logarithmically with the input power, since $V_{oc} = \frac{k_bT}{q} \ln \left( \frac{1}{J_{dark}} + 1 \right) \approx \frac{k_bT}{q} \ln \left( \frac{1}{J_{dark}} \right)$ for large illumination current densities $J_{sc}$. Here, $J_{dark}$ is the dark current density, $n$ is the ideality factor, $k_b$ is the Boltzmann constant, $T$ is the temperature of the device, and $q$ is the fundamental unit of charge, so that $\frac{k_bT}{q} \approx 0.0258V$ at room temperature. In
Figure 3b we see that the experimental data match well with the diode fit (dashed black line, see Supporting Information S2 for fitting details), suggesting an ideality factor of $n = 1.75$ and a dark current density $J_{\text{dark}} = 0.65 \text{mA}/(\text{cm}^2)$ assuming a 30 $\mu$m $\times$ 30 $\mu$m device area. Also, since the PCE is given as $PCE = J_{sc}V_{oc}FF/P_{in}$ where $J_{sc}$ is the short-circuit current density, $V_{oc}$ is the open-circuit voltage, FF is the fill fraction, and $P_{in}$ in the incident power density, we would expect the PCE to scale roughly logarithmically as well. This is true for laser powers up to $\sim 740 \text{ W/cm}^2$ (Figure 3c). However, for larger input power, the PCE decreases with increasing power. Such a drop in PCE can be attributed to series resistances in the device, either at the contacts or at the junction. This is corroborated by the match between the experimental data (dots) and the fitted expression (dashed line), yielding the diode fitting parameters in the lower right-hand corner of the plot in Figure 3c. The fit for the $V_{oc}$ was simultaneously done with the PCE, therefore yielding the same set of parameters and a good match between experiment and extracted device parameters. Finally, we observed a decrease in the EQE at 633 nm with increasing power (Figure 3d). Using the above fitted parameters, series resistance can only be used to partially explain a decrease in the EQE at higher powers. Thus, the additional decrease in EQE at higher powers may be due to the onset of carrier density-dependent nonradiative processes such as Auger or biexcitonic recombination which are not accounted for in the diode fit used above, where the dark current is fixed for all powers.

**Absorption in van der Waals Heterostructures.** We first investigate the absorption and optical properties of van der Waals heterostructures. We formed a heterostructure composed of hexagonal boron nitride (hBN)/FLG/WSe$_2$/MoS$_2$/Au. The composite heterostructure has various regions (inset of Figure 4a), corresponding to different vertical heterostructures. Given the sensitivity of the performance of van der Waals materials to different environmental conditions and device fabrication procedures, the samples fabricated here allow us to...
study optical and electronic features of different heterostructures in a systematic manner by probing specific heterostructures fabricated on the same monolithic substrate. This is enabled by the small spot size of our laser, which additionally allows us to properly normalize the spectral response without artificially including geometric factors (see Methods for details).

As an example, consider the optical response at the location of the blue dot in the inset of Figure 4a. The vertical heterostructure there is composed of 1.5 nm FLG/4 nm WSe$_2$/5 nm MoS$_2$/Au. This location can be probed spectrally for its absorption characteristics (Figure 4a), revealing near-unity absorption in van der Waals heterostructures. The peaks at $\sim$610, $\sim$670, and $\sim$770 nm correspond to the resonant excitation of the MoS$_2$ B exciton, MoS$_2$ A exciton, and WSe$_2$ A exciton, respectively.$^{22}$ On the other hand, the broad mode at $\sim$550 nm corresponds to the photonic mode that leads to near-unity absorption.$^{23}$ Measurements of the absorption can be corroborated with electromagnetic simulations, unveiling both the accuracy between simulation and experimental results as well as the fraction of photon flux absorbed into individual layers of the heterostructure stack (Figure 4b). Despite the near-unity absorption observed in the heterostructure stack, there is parasitic absorption in both the underlying gold substrate and in the FLG that accounts for $\sim$20% of the total absorbance. Such parasitic absorption can be reduced by using a silver back reflector, as shown in Figure 4c,d. We find that the simulated and measured absorbance is also in good agreement for the case of a silver back reflector. Thus, the optical response of a van der Waals heterostructure can be modeled accurately using full wave electromagnetic simulations, and our method of measurement yields accurate and reliable results.

To note, the subwavelength dimension of the total heterostructure thickness is critical for achieving near-unity absorption. Indeed, the entire stack can be treated as a single effective medium, where small phase shifts are present between layers, and therefore the material discontinuities are effectively imperceptible to the incident light (see Supporting Information S3 for details). Ultimately, the van der Waals heterostructure-on-metal behaves as a single absorbing material with effective medium optical properties. Therefore, as previously demonstrated, near-unity absorption at different wavelengths can be achieved for an absorber layer with the appropriate thickness$^{23,24}$ ($\sim$10–15 nm total thickness for TMD heterostructures).

**Carrier Collection in van der Waals Semiconductor Junctions.** As discussed above, another criterion for high EQE is efficient carrier collection. Given the large exciton binding energies in TMDs ($\sim$50–100 meV in the bulk),$^{25,26}$ the large internal electric field at the semiconductor heterojunction may...
play a role in exciton dissociation and subsequent carrier collection. Charge carrier separation in TMDs can be accomplished using either a pn junction or a Schottky junction, and we find that a pn heterojunction dramatically enhances the EQE when compared with a Schottky junction.

The heterostructure described in Figure 4a,b can be probed as an optoelectronic device with the formation of a top electrode (see inset of Figure 5a). Since the back reflector (gold) can simultaneously serve as a back contact to the entire vertical heterostructure, we can use this scheme to compare the electronic performance of various vertical heterostructures. Given the work function between WSe2 (p-type) and Au, it is expected that a Schottky junction will form between the two materials (see Figure 1c), whereas WSe2 (p-type) on top of MoS2 (n-type) is expected to form a pn heterojunction. High spatial resolution scanning photocurrent microscopy allows us to examine the two heterostructure devices in detail (Figure 5a). We observe large photocurrent for the pn heterojunction geometry (yellow region) compared to the Schottky junction geometry (light blue region). The decrease of the photocurrent in the left-side of the yellow region in Figure 5a is due to shadowing from the electrical probes. A line cut of the spatial photocurrent map shown in Figure 5b provides a clearer distinction between the two junctions, demonstrating ∼6× more photocurrent for the pn junction relative to the Schottky junction.

The photocurrent density is directly related to the EQE and therefore the product of the absorbance and IQE. In order to quantitatively compare the electronic differences between the two junctions, we need to normalize out the different optical absorption in the two devices, i.e., compute the IQE of each device:

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Figure 5. Charge transport and collection in vertical pn and Schottky junction geometries: (a) Spatial photocurrent map of the fabricated van der Waals heterostructure device using a 633 nm laser excitation. The inset is an optical image of the device (scale bar = 20 μm). (b) The line profile of the dotted red line arrow in (a), illustrating the different photocurrent intensities depending on the device geometry (Schottky and pn junction). (c, i) Experimentally measured spectral characteristics of the absorbance (blue) and EQE (red) in the 1.5 nm FLG/4 nm WSe2/Au (Schottky geometry) device along with the (ii) experimentally derived IQE (purple) and the calculated active layer IQE (green). The inset is a cross-sectional schematic of the measured device, at the orange dot in (a). (d) Same as in (c) except with a 1.5 nm FLG/4 nm WSe2/5 nm MoS2/Au (pn geometry) device. The inset is a cross-sectional schematic of the measured device, at the purple dot in (a).
\[ \text{IQE}_{\text{exp}}(\lambda) = \frac{\text{EQE}(\lambda)}{\text{Abs}(\lambda)} \]  

where \( \text{EQE}(\lambda) \) and \( \text{Abs}(\lambda) \) are the experimentally measured EQE and absorbance of their respective devices (Figure 5c,i and d,i). A plot of the experimentally derived IQE (i.e., \( \text{IQE}_{\text{exp}} \)) is shown in purple in Figure 5c,ii and d,ii. This plot also confirms that a pn junction geometry (with \( \text{IQE}_{\text{exp}} \sim 40\% \)) formed of van der Waals materials is more efficient for carrier collection than a Schottky junction geometry (with \( \text{IQE}_{\text{exp}} \sim 10\% \)).

Embedded in the above analysis is yet another convolution of the optical and electronic properties. As per Figure 4b, we found that absorption in FLG and Au accounted for \( \sim 20\% \) of the absorbance of the total heterostructure. Assuming very few photons absorbed in those layers ultimately are extracted as free carriers (i.e., \( \text{IQE}_{\text{FLG}} \approx \text{IQE}_{\text{Au}} \approx 0 \)), the IQE defined above convolutes the parasitic optical loss with the electronic loss in the device. Thus, another useful metric we shall define is \( \text{IQE}_{\text{active}} \), the active layer IQE:

\[ \text{IQE}_{\text{active}}(\lambda) = \frac{\text{EQE}(\lambda)}{\text{Abs}(\lambda) - \text{Abs}_{p}(\lambda)} \]  

where the additional term \( \text{Abs}_{p}(\lambda) \) corresponds to the parasitic absorption in the other layers of the device that do not contribute to current (i.e., Au and FLG in this device). Thus, \( \text{IQE}_{\text{active}}(\lambda) \) is a measure of the carrier generation and collection efficiency only in the active layer (i.e., WS\textsubscript{2} and MoS\textsubscript{2}) of the device and is purely an electronic efficiency as defined above. We shall use this quantity to accurately compare electronic geometries. Given the good agreement between simulations and experiment shown in Figure 4, a simple method of estimating the parasitic absorption described above is therefore through electromagnetic simulations. \( \text{IQE}_{\text{active}} \) of the Schottky
and pn heterojunction geometries calculated with eq 2 is shown in Figure 5c,i and d,ii with dotted green curves.

Analysis of these plots reveals several important points. First, IQE-active for the pn junction geometry is ∼3× higher than in the Schottky junction geometry when spectrally averaged. Though yet to be fully clarified, we attribute higher IQE in pn heterojunctions to the larger electric fields in a pn heterojunction that may lead to a higher exciton dissociation efficiency and consequently IQE. Second, compared to the IQE which included the parasitic absorption (purple dots in Figure 5e,ii and d,ii, the active layer IQE curves (green dots) are spectrally flat within measurement error and calculations (ΔIQE/IQE ≈ 0.07). Thus, the few broad peaks around the exciton energies of WSe₂ (∼770 nm) and MoS₂ (∼610 nm and ∼670 nm) in IQE-exp are not attributed to, e.g., resonant excitonic transport phenomena, but rather as a simple convolution of the optical and electronic effects when calculating the electronic IQE. In other words, consideration of parasitic absorption is critical when analyzing the electronic characteristics of thin optoelectronic devices. However, IQE-exp is still a useful metric, as it effectively sets a lower bound on the true IQE. Generally, we expect IQE-exp ≤ IQEtrue ≤ IQE-silver as electromagnetic simulations tend to slightly overestimate the absorption when compared with experimental results. Thus, in this paper, we shall plot both expressions when comparing different electronic device geometries. Finally, it is important to mention that an active layer IQE of ∼70% is achieved in van der Waals heterostructures without complete optimization of the electronic configuration of the device, such as the band profiles and the specific choice of contacts. With careful electronic design, we suggest it may be possible to achieve active layer IQEs >90%.

Optically Transparent Contacts for Carrier Extraction. As another aspect of analysis, we studied the role of vertical carrier collection compared to lateral carrier collection in van der Waals heterostructures. Graphene and its few-layer counterpart can form a transparent conducting contact allowing for vertical carrier collection, in contrast to in-plane collection (see Figure 1d). The strong, in-plane covalent bonds of van der Waals materials suggest that in-plane conduction may be favorable when contrasted with the weak out-of-plane van der Waals interaction. However, the length scale for carrier transport in-plane (∼μm) is orders of magnitude larger than in the vertical direction (∼nm). Therefore, transport in a regime in between these two limiting cases is not surprising.

Silver exhibits lower absorption in the visible than gold, suggesting it could be an optimal back reflector for photovoltaic devices, as seen in Figure 4. Thus, we contrast the case of in-plane and out-of-plane conduction concurrently with the presence of two different back reflectors that simultaneously function as an electronic back contact (gold vs. silver) to a pn heterojunction, as in Figure 6a,b. Optical and photocurrent images of the devices are shown in Supporting Information S1.

Our results in Figure 6c,d show the distinctions between the various contacting schemes. In the case of both silver and gold, a transparent top contact such as FLG seems to enhance the carrier collection efficiency. This is particularly true in the case of silver, where IQE-active enhancements of ∼5× is apparent. In the case of gold, the IQE is enhanced by about ∼1.5× when parasitic absorption is taken into account. By analyzing the work functions of gold (∼4.83 eV) and silver (∼4.26 eV), the Schottky–Mott rule suggests in both cases that a Schottky barrier should form equal to ϕ_g = ϕ_M − χ, where ϕ_M is the Schottky barrier height, ϕ_M is the work function of the metal, and χ is the electron affinity of the semiconductor. However, several reports have indicated that gold appears to form an electrically ohmic contact to MoS₂, which we observe here. Conversely, the above data suggest that silver and MoS₂ follow the traditional Schottky–Mott rule, leading to the formation of a small Schottky barrier of ∼0.26 eV. Given that the energy barrier is about 10kT, very few electrons can be extracted out of the pn heterojunction when silver is used as a back contact, leading to very low IQEs. By taking into account just the active layer (dashed lines), we see that gold is ∼2× better as an electronic contact than silver.

Finally, we examine the role of vertical carrier collection on the I–V characteristics of the two devices (Figure 6e,f). In the case of gold, we can see purely an enhancement of the short-circuit current with vertical carrier collection. On the other hand, vertical carrier collection for silver drastically increases both the short-circuit current density and the open-circuit voltage. This phenomenon is consistent with the previously described nature of gold (ohmic) and silver (Schottky) contacts. Namely, on silver in the absence of a transparent top contact, due to both the Schottky barrier and the large in-plane propagation distance, carriers are collected with poor efficiency leading to a small I_sc. Consequently, a high recombination rate of the generated carriers which are inefficiently extracted leads to small V_oc values. On the other hand, even in the absence of a transparent electrode, gold enables efficient extraction of electrons from the pn heterojunction as an ohmic contact. Thus, the short-circuit current and open-circuit voltage in gold are higher compared to the silver back contact, even in the absence of a transparent electrode. When introducing FLG as a transparent top contact, the propagation distance is significantly reduced in the silver device, and carriers can be extracted with much higher efficiency, leading to a large enhancement of both the current and voltage. Whereas for gold, the FLG enhances the already high carrier collection (yielding larger I_sc), but only has a negligibly small enhancement effect on the open-circuit voltage. Overall, these results demonstrate that vertical carrier collection plays a crucial role in high photovoltaic device performance in van der Waals heterostructures.

Thickness Dependence on Charge Collection Efficiency. As a final point of analysis, we briefly examined the effect of thickness on IQE-active under vertical carrier collection. We compared the optoelectronic characteristics of a thicker pn heterojunction (11 nm hBN/1.5 nm FLG/4 nm WSe₂/9 nm MoS₂/Au) with a thinner pn heterojunction (1.5 nm FLG/4 nm WSe₂/5 nm MoS₂/Au). The experimentally measured absorbance and EQE are plotted in Figure S4 for reference. By normalizing out the differences in absorption between the pn junctions, we see a somewhat surprising result when we analyze the active layer IQE (dashed lines, Figure 7). In particular, despite the roughly 50% more length in active layer thicknesses (13 nm vs. 9 nm) and qualitatively different absorption and EQE spectra, the thick pn junction exhibits nearly the same active layer IQE compared to the thin pn junction. In fact, it appears to be slightly more efficient, but this is within the error bar of the measurement and simulations (ΔIQE/IQE ≈ 0.07, see Methods for details of errors). This observation is corroborated with the experimentally derived IQE (dotted curve, Figure 7), which has nearly the same spectrum between the two thicknesses, but differ in magnitude due to differences in parasitic absorption. This result suggests that the ultrathin
limit (~10 nm) of van der Waals heterostructures with vertical carrier collection, the IQE has a weak dependence on active layer thickness. This weak dependence may be due to a combination of increased scattering competing with charge transfer, tunneling, and exciton quenching effects as the vdW heterostructure becomes thicker. The exact role of each of these effects, as well as possibly other effects, will require a new theoretical framework and experimental measurements to analyze their relative contributions to charge collection efficiency.

CONCLUSIONS

Our results suggest important challenges that must be addressed to enable high photovoltaic efficiency. For example, despite the usefulness of gold as an electrical back contact, we found from electromagnetic simulations that it accounts for nearly 20% of the parasitic loss in the heterostructures reported here. Schemes using optically transparent carrier selective contacts could be used to avoid this parasitic optical loss. Another open question is the role and importance of exciton dissociation and transport. Indeed, the large exciton binding energies in TMDs (~50–100 meV in the bulk) suggests that a significant exciton population is generated immediately after illumination. However, it is not yet clear whether such an exciton population fundamentally limits the IQE of the device, posing an upper limit on the maximum achievable IQE in van der Waals material-based photovoltaic devices. Finally, the problem of open-circuit voltage must also be addressed. For example, the type-II band alignment between ultrathin MoS2 and WSe2 suggests a renormalized bandgap of ~400–500 meV given by the minimum conduction band energy and maximum valence band energy of the two materials. In accordance with the Shockley–Queisser limit, this would severely reduce the maximum PCE attainable by a factor of ~3. Therefore, to achieve higher open-circuit voltages, a monolithic device structure may be required to avoid low-energy interlayer recombination states.

However, our results described here also suggest a different approach in addressing the optical and electronic considerations for ultrathin van der Waals heterostructures when compared with conventional photovoltaic structures. For example, our observation that ultrathin van der Waals heterostructures can be optically treated as a single effective medium is a regime of optics that is uncommon for the visible to near-infrared wavelengths analyzed in photovoltaic devices. Likewise, our observation of weak thickness dependence of the charge collection efficiency represents a realm of electronic transport that is also quite unconventional and unexplored when compared to traditional photovoltaic structures. Thus, the combination of the above observations may enable entirely different photovoltaic device physics and architectures moving forward.

To summarize, we have shown that EQEs >50% and active layer IQEs >70% are possible in vertical van der Waals heterostructures. We experimentally demonstrated absorbance >90% in van der Waals heterostructures with good agreement to electromagnetic simulations. We further used the active layer IQE to quantitatively compare the electronic charge collection efficiencies of different device geometries made with van der Waals materials. By further reducing parasitic optical losses and performing a careful study on exciton dissociation and charge transport while simultaneously engineering the band profiles and contacts, van der Waals photovoltaic devices may be able to achieve EQEs >90%. Our results presented here show a promising and exciting route to designing and achieving efficient ultrathin photovoltaics composed of van der Waals heterostructures.

METHODS

Metal Substrates Preparation. Atomically smooth metal substrates were prepared using the template stripping technique. We prepared the substrates using polished silicon wafers (University Wafer) with native oxide and then cleaned the silicon substrates via sonication in acetone (10 min), followed by sonication in isopropyl alcohol (10 min). Samples were then blow dried with nitrogen gas before cleaning with oxygen plasma (5 min, 100 W, 300 mTorr under O2 flow).

Metal was then deposited via electron beam evaporation on the polished and cleaned surface of the silicon wafer. For gold (Plasmatic Materials, 99.99% purity), base pressures of ~3 × 10−7 Torr was achieved before depositing at 0.3 A/s. This continued until a thickness of ~20 nm was achieved. Then, the rate was slowly ramped to 1 A/s and then held there until a total thickness of 120 nm was reached. For silver (Plasmatic Materials, 99.99% purity), following McPeak et al., we deposited at a base pressure of ~3 × 10−7 Torr at 40 A/s for a final thickness of 150 nm. After deposition of the metal, an adhesive handle was formed using a thermal epoxy (Epo-Tek 375, Epoxy Technology). One g of part A Epo-Tek 375 and 0.1 g of part B Epo-Tek 375 were mixed in a glass vial and were allowed to settle for ~30 min. Afterward, individual droplets of the mixture were added directly onto the metallic surface before placing cleaned silicon chips (~1 cm2) on top. The droplet of epoxy was allowed to settle under the weight of the silicon chip before placing on a hot plate (~80 °C) for 2 h. Individual chips were then cleaved with a razor blade, forming the final substrate consisting of atomically smooth metal/thermal epoxy/silicon. Typical RMS surface roughness of the metal was <0.3 nm using this technique (examined via AFM).

van der Waals Heterostructure Fabrication. The bottom-most layer of the van der Waals heterostructure (e.g., MoS2) was directly exfoliated onto the metallic substrates prepared using the above technique. Exfoliation was performed using bulk crystals purchased from HQ Graphene using Scotch tape. Subsequent layers were formed using a viscoelastic dry transfer technique using a home-built setup at
room temperature. Dry transfer was performed using PF-20-X4 Gel Film from Gel-Pak as the transparent polymer. van der Waals materials were directly exfoliated onto the polymer using Nitto tape and then mechanically transferred onto the MoS\textsubscript{2}/metal substrate. Samples were examined in the optical microscope during each layer of the process, and an AFM scan was performed afterward to extract the thicknesses of individual layers. Thicknesses were then corroborated with optical measurements and calculations.

A top electrode was patterned using standard photolithography techniques. NR-9 1000 PY was used as a negative resist. The resist was spun at 5000 rpm for 55 s before baking at 150 °C for 1 min. A mask aligner with a prepatterned mask was used to define the features and aligned on top of the van der Waals heterostructure. After exposure for ~18 s under 10 mW of UV light (λ = 365 nm), the resist was postbaked at 105 °C for 1 min and cooled to room temperature. Finally, the resist was developed using RD-6 developer for 10–15 s before rinsing in deionized water for 35 s. The sample was then blow-dried with nitrogen and examined under an optical microscope.

Electron beam deposition was then used to form the top ring electrodes (10 nm Ti/90 nm Au). Base pressures of ~3 × 10\textsuperscript{−6} Torr was achieved before the beginning of the deposition. For titanium, a deposition rate of 0.3 A/s was used for the entirety of 10 nm. Immediately afterward, gold was deposited at a rate of 0.3 A/s for 15 nm. The rate was slowly ramped to 0.6 A/s for 10 nm and then to 0.9 A/s for another 10 nm. At 35 nm of total gold thickness, the rate was finally ramped to 1.0 A/s until the total gold thickness was 90 nm. The resist was then lifted-off using heated acetone (40–45 °C) for 30 min. If needed, the samples were sonicated in 5 s intervals in acetone to remove the resist. The sample was then rinsed in isopropanol alcohol and blow dried with nitrogen.

**Spatial Photocurrent Map and IV Measurements.** Samples were contacted on the top electrode and bottom metallic substrate using piezoelectric controlled probes (MiBots, Imina Technologies) were contacted on the top electrode and bottom metallic substrate and cross-referenced with the EQE spectrum of the measured device.

Electromagnetic Simulations.** Calculations were performed using the transfer matrix method\textsuperscript{20} with optical constants taken from literature for each of the TMDs.\textsuperscript{21} We assumed that for the TMD thicknesses analyzed in this paper, their optical response can be represented by the bulk optical permittivities. Permittivities of Ag and Au were taken from McPeak\textsuperscript{45} and Olman,\textsuperscript{47} respectively. The optical response of FLG was assumed to be like graphite, with its dielectric constant stemming from the assumption of normal incidence for both absorption and EQE measurements while using a NA = 0.5S objective, fluctuations in the laser power during the measurement, and sample contact stability. In addition, we have observed in our laser that there is relatively little power for λ < 450 nm. Additionally, there is relatively high absorbance in the 50× objective for λ > 700 nm. Combined with the fact that the simulated parasitic absorption accounts for a larger fraction of the total absorption for λ < 450 nm and λ > 700 nm, the significantly noisier spectra in the active layer IQE at these wavelengths can be attributed to the factors described above.

**Spectral Absorbance and EQE Measurements.** Quantitative absorbance and EQE measurements were performed using a home-built optical setup. A supercontinuum laser (Fianium) was coupled to a monochromator to provide monochromatic incident light. A series of apertures and mirrors were used to collimate the beam before being focused on the sample with a long working distance (NA = 0.55) 50X objective to provide a small spot size (~1 μm lateral resolution). Power-dependent IV measurements were performed at particular locations of the device using the spatial photocurrent maps. Illumination power was modified using neutral density filters in the microscope, and the incident power was measured by a photodetector and cross-referenced with the EQE spectrum of the measured device.

**Absorption and EQE Measurements.** For absorption measurements, the sample itself was used as a photodetector. The top ring electrode and bottom metallic substrate was probed using MiBots. Laser light was then focused on a particular spot, and the current was collected by the probes and sent through a lock-in amplifier for homodyne detection, as in the reflection spectrum case. After measurement of the current signals from the sample, another spectral scan was performed with the optical system in the same configuration using a NIST calibrated photodetector (818-ST2-UV/DB, Newport). Power fluctuations between scans were again accounted for by using a small amount of beam-split light and sending it to a photodetector. The measured currents were normalized to this photodetector’s current before being normalized to the calibrated photodetector to yield the absolute EQE.

Despite the various steps of calibration used for normalization, we still estimate measurement errors of δAbs/Abs ≈ 0.02 and δEQE/EQE ≈ 0.05 stemming from the assumption of normal incidence for both absorption and EQE measurements while using a NA = 0.5S objective, fluctuations in the laser power during the measurement, and sample contact stability. In addition, we have observed in our laser that there is relatively little power for λ < 450 nm. Additionally, there is relatively high absorbance in the 50× objective for λ > 700 nm. Combined with the fact that the simulated parasitic absorption accounts for a larger fraction of the total absorption for λ < 450 nm and λ > 700 nm, the significantly noisier spectra in the active layer IQE at these wavelengths can be attributed to the factors described above.

Given that there is sample-to-sample variation of the dielectric constant, it is likely that the literature values of the dielectric constant differ from the samples measured here. This difference we estimate leads to absorption simulation errors of ~5%. Assuming this is true, the estimated error for the active layer IQE can be approximated as

\[
\delta \text{IQE} \approx \sqrt{\left(\frac{\delta \text{Abs}}{\text{Abs}}\right)^2 + \left(\frac{\delta \text{Abs}}{\text{Abs}}\right)^2 + \left(\frac{\delta \text{EQE}}{\text{EQE}}\right)^2},
\]

which is about 7%.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.7b03148.

Additional experimental data, calculations, and analysis accompany this paper (PDF).

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Acknowledgments

This work is part of the "Light-Material Interactions in Energy Conversion" Energy Frontier Research Center funded by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under award no. DE-SC0001293. D.J., A.R.D., and M.C.S. acknowledge additional support from the Space Energy Sciences under award no. DE-SC0001293. D.J., A.R.D., and M.C.S. acknowledge additional support from the Kavli Nanoscience Institute Graduate and Postdoctoral Fellowships. A.R.D. also acknowledges financial support. J.W. acknowledges support in part from the Kavli Nanoscience Institute Graduate Research Fellowship under grant no. 1144469. K.T. would like to thank the Caltech SURF program and the Northrop Grumman Corporation for financial support.

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