We report a van der Waals heterostructure formed by monolayers of MoS$_2$ and ReS$_2$ with a type-I band alignment. First-principle calculations show that in this heterostructure, both the conduction band minimum and the valence band maximum are located in the ReS$_2$ layer. This configuration is different from previously accomplished type-II van der Waals heterostructures where electrons and holes reside in different layers. The type-I nature of this heterostructure is evident by photocarrier dynamics observed by transient absorption measurements. We found that carriers injected in MoS$_2$ transfer to ReS$_2$ in about 1 ps, while no charge transfer was observed when carriers are injected in ReS$_2$. The carrier lifetime in the heterostructure is similar to that in monolayer ReS$_2$, further confirming the lack of charge separation. We attribute the slower transfer time to the incoherent nature of the charge transfer due to the different crystal structures of the two materials forming the heterostructure. The demonstrated type-I semiconducting van der Waals heterostructure provides new ways to utilize two-dimensional materials for light emission applications, and a new platform to study light–matter interaction in atomically thin materials with strong confinement of electrons and holes.

1 Introduction

Forming heterostructures by combining two materials is one of the key approaches to control electrons in semiconductors, which has led to discoveries such as integer, fractional, and quantum spin Hall effects, as well as exciton-polariton condensations. Functional materials based on heterostructures are the cornerstone of many applications including semiconductor lasers and light emitting diodes. Depending on the work functions of the involved materials, two types of band alignments can be formed in heterostructures. A type-I alignment refers to the case where both the conduction band minimum (CBM) and the valence band maximum (VBM) are located in the material with the narrower bandgap, as shown in Fig. 1(a) and (b). Electrons and holes excited in the wide-gap material transfer to the narrow-gap material, as indicated by the arrows. The quantum confinement of electrons and holes in the same region facilitates their radiative recombination, which is desirable in light-emitting applications. Carriers excited in the narrow-gap material are prohibited from interlayer transfer due to their lower energies. When the CBM and VBM are located in different materials, the alignment is called type-II. Fig. 1(c) and (d) show the situation where the CBM and VBM are located in the narrow-gap and wide-gap materials, respectively. In this case, excitation of the...
wide-gap material is followed by the transfer of electrons, but not the holes, while opposite charge transfer occurs when the narrow-gap material is excited. The separation of the electrons and holes to different layers can increase their lifetime, and is desired for applications in photovoltaics and photodetection.7–10

The newly developed two-dimensional materials brought the semiconductor heterostructures to a new regime exhibiting an ultimate thinness, atomically sharp and near-defect-free interfaces, exceptional mechanical flexibility, and chemical stability.11,12 More importantly, van der Waals interlayer coupling imposes almost no restrictions on the selection of component materials, opening up a huge number of combinations to explore and utilize. So far, most semiconducting van der Waals heterostructures are composed of monolayers (MLs) of different materials in the family of transition metal dichalcogenides (TMDs). Using the four most extensively studied TMDs, namely MoS2, MoSe2, WS2, and WSe2, 6 combinations exist, and have all been studied: MoS2–MoSe2,13 MoS2–WS2,14–18 MoS2–WSe2,19–25 MoSe2–WSe2,26 MoSe2–WSe2,27 and WS2–WSe2.28 These heterostructures were all found to be type-II, with qualitatively similar electronic and optical properties. Hence, demonstration of a semiconducting van der Waals heterostructure with type-I band alignment is of fundamental importance for control of electrons in atomically thin materials. Such a structure can facilitate applications of van der Waals heterostructures for light emitting devices and enable designs of sophisticated bandgap landscaping in functional materials. It will also provide a platform to study light–matter interaction in atomically thin systems with adjustable confinement energy, and to study the transport of electron–hole pairs and excitons across van der Waals interfaces. Here we demonstrate the first type-I semiconducting van der Waals heterostructure, MoS2–ReS2.

2 Results and discussion

Our first-principle computation results of the electronic structures of ReS2 and MoS2 MLs, as well as the MoS2–ReS2 heterostructure, are shown in Fig. 2(a) and (b). Details of the first-principle computation are given in the ESI† Fig. 2(a) shows the computed band alignment for ML MoS2 and ReS2. Clearly, the VBM of ReS2 is higher than that of MoS2, while the CBM of ReS2 is lower than that of MoS2. Hence, the band-alignment information alone already suggests that the MoS2–ReS2 heterostructure is likely to exhibit type I band alignment. Computed band structures of MoS2–ReS2 heterostructure is depicted in Fig. 2(b). It can be seen that the CBM is located at the Γ point while the VBM is located at the point between K and Γ. Nevertheless, both CBM and VBM are contributed in ReS2, consistent with the band alignment computation. Note also that the computed bandgap of ReS2 (1.32 eV) is slightly less than the measured optical gap (1.52 eV, see below) because computation based on generalized gradient approximation (see ESI†) tends to underestimate the bandgap.

Fig. 2(c) shows the fabricated heterostructure sample of MoS2/ReS2. The individual MLs were mechanically exfoliated from bulk crystals onto polydimethylsiloxane substrates and sequentially transferred to a Si/SiO2 (90 nm) substrate. The sample was then annealed at 200 °C in an Ar/H2 (100 sccm/5 sccm) environment at a base pressure of about 3 Torr. The thickness of the flakes was confirmed to be ML by optical contrast, photoluminescence, and Raman measurements. Fig. 2(d) shows optical contrast along the yellow line in (c) from the green channel of the image. The contrast of about 11 to 13% for each layer is consistent with previous reports for TMD MLs,29,30 and in particular, ReS2 MLs.31 Photoluminescence of the MoS2 flake is also consistent with its ML thickness (see ESI†). Furthermore, Raman spectra of both the ReS2 (Green) and MoS2 (Red) flakes are shown in Fig. 2(e), providing further evidence of ML thickness.32–36 Specifically, the separation of the E2g and A1g peaks is used to confirm the ML thickness of the MoS2 flake,32 while the separation of the I and III peaks is consistent with previous results of ReS2 MLs33 (see ESI†). The ML flakes were large enough to provide a sufficiently large overlapped region as well as isolated ML regions for a thorough control experiment. The sample was under ambient conditions in all the measurements.

The nature of the band alignment of MoS2–ReS2 was revealed by studying its photocarrier dynamics with transient absorption measurements. First, a pump pulse of 1.67 eV and 300 fs was focused to the ReS2 ML region, with a spot size of about 2.2 µm and a fluence of 7 µJ cm−2. Since the measured optical bandgap of ML ReS2 is about 1.52 eV,37 the pump pulse injects photocarriers in ReS2 by interband absorption. With an absorption coefficient of 8 × 106 m−1 (bulk value38), the peak injected...
density is about $1.4 \times 10^{11}$ cm$^{-2}$. The dynamics of these photocarriers is monitored by measuring the differential reflection of a 1.53 eV probe pulse, tuned slightly above the band gap of ReS$_2$. The differential reflection is defined as $D/R_0 = (R/ R_0)/R_0$, where $R$ and $R_0$ are the reflection of the probe with and without the presence of photocarriers, respectively. This quantity is proportional to the carrier density in all the configurations used in this study (see ESI†). The measured differential reflection signal is shown as the open red squares in Fig. 3. The decay of the signal can be fit by a single-exponential function with a background (black curve). The decay constant of 65 ps can be attributed to the recombination lifetime of the photocarriers.

Similar results were obtained when the measurement was performed on the MoS$_2$–ReS$_2$ heterostructure, shown as the solid red squares in Fig. 3. With the energy of the lowest excitonic state at 1.85 eV, the MoS$_2$ layer of the heterostructure was neither pumped nor probed. However, the MoS$_2$ layer could serve as a channel for charge transfer if the band alignment were type-II. Specifically, if the CBM (VBM) were lower (higher) in MoS$_2$ compared to ReS$_2$, electrons (holes) would be expected to transfer to MoS$_2$ [see Fig. 1(c) and (d)]. If either transfer occurred, one would expect a smaller signal magnitude (by about a factor of 2) due to the loss of one type of carrier in ReS$_2$ and a slower decay of the signal due to a longer carrier lifetime, resulting from the spatial separation of the electrons and holes. Hence, this result indicates the lack of charge transfer and indirect exciton formation in MoS$_2$–ReS$_2$, both of which have been generally observed in type-II TMD heterostructures.$^{13-28}$
MoS\textsubscript{2} with a peak density of 4.8 \times 10^{11} \text{ cm}^{-2} (according to an absorption coefficient of 3 \times 10^{4} \text{ m}^{-1}).\textsuperscript{39} For type-I alignment, these excitons should transfer to ReS\textsubscript{2}, resulting in an increase of the signal from the 1.67 eV pump. The signal (solid green squares) is indeed increased, by about a factor of 3, and decays with a similar time constant (62 ps, blue curve) as the ReS\textsubscript{2} ML. Both features suggest that excitons injected in MoS\textsubscript{2} transfer to ReS\textsubscript{2}, confirming the high quality of the interface. To confirm that the increased signal is not from additional carriers injected in ReS\textsubscript{2}, due to the larger pump photon energy, we repeated the measurement on the ReS\textsubscript{2} ML, shown as the open green squares. The signal is similar to that obtained with the 1.67 eV pump (open red squares), confirming that the signal from ReS\textsubscript{2} alone is rather insensitive to the pump photon energy in this range. This is also consistent with previous observations that the absorption coefficient of ReS\textsubscript{2} does not change significantly in the range of 1.67–1.85 eV.\textsuperscript{10}

Further evidence of the type-I band alignment was obtained by probing the MoS\textsubscript{2} layer, as summarized in Fig. 4. At first, a 3.06 eV pump pulse was used to inject photocarriers in the MoS\textsubscript{2} ML region. From the fluence of 12 \mu J cm\textsuperscript{-2}, a peak injection density of 3.4 \times 10^{12} \text{ cm}^{-2} is estimated, using an absorption coefficient of 1.5 \times 10^{5} \text{ m}^{-1}.\textsuperscript{39} Shown as the solid squares in Fig. 4, a differential reflection signal on the order of 10\textsuperscript{-3} was observed with a 1.85 eV probe, tuned to the A-exciton of ML MoS\textsubscript{2}. Next, we used a 1.53 eV pump with a fluence of 33 \mu J cm\textsuperscript{-2} to inject photocarriers in the ReS\textsubscript{2} layer of the heterostructure with a peak carrier density of 7.6 \times 10^{11} \text{ cm}^{-2}. If the band alignment were type-II, either electrons or holes would transfer to MoS\textsubscript{2} [Fig. 1(c) and (d)]. With such a density, the transferred carriers are expected to induce a differential reflection signal of the 1.85 eV probe on the order of 10\textsuperscript{-4}. However, as shown by the open squares in Fig. 4, no signal above 10\textsuperscript{-4} was observed. This further confirmed the lack of charge transfer from ReS\textsubscript{2} to MoS\textsubscript{2} in the heterostructure.

The demonstrated type-I band alignment of MoS\textsubscript{2}–ReS\textsubscript{2} allows us to study the interlayer transfer process of photocarriers in this configuration, as summarized in Fig. 5. In this figure, all the differential reflection signals are normalized in order to facilitate a visual comparison. First, a 1.85 eV pulse with a peak fluence of 7 \mu J cm\textsuperscript{-2} was used to pump the heterostructure. According to the absorption coefficients of the two MLs, the peak densities of photocarriers injected in the MoS\textsubscript{2} and ReS\textsubscript{2} layers are 4.8 \times 10^{11} \text{ cm}^{-2} and 1.3 \times 10^{11} \text{ cm}^{-2}, respectively. By probing the ReS\textsubscript{2} layer with a 1.53 eV pulse, we monitored the evolution of photocarrier population in that layer, and hence time resolved the transfer of photocarriers from the MoS\textsubscript{2} layer to the ReS\textsubscript{2} layer. We found a rather slow rise of the signal, as shown by the solid green squares. The contribution of the photocarriers directly injected in ReS\textsubscript{2} can be isolated by the control experiment on ML ReS\textsubscript{2}, under the same conditions, as shown by the open green squares. The actual magnitude of this signal is about one third of the signal from the heterostructure, which is consistent with the ratio of the carriers injected in the two layers. Clearly, the rising of the signal is much faster in ML ReS\textsubscript{2}, confirming that the slower rising observed in the heterostructure originates from the transfer of photocarriers across the interface. We found that the rising part of the signal can be fit by \[ N_1[1 - \exp(-t/\tau_2)] + N_2, \text{ where } N_1 \text{ and } N_2 \text{ represent the photocarrier densities injected in MoS}_2 \text{ and ReS}_2 \text{ layers, respectively, and } \tau_2 \text{ the transfer time. As shown by the solid green curve in Fig. 5, this expression can satisfactorily describe the data, with a transfer time of } 1.3 \pm 0.1 \text{ ps.} \]

In addition to time resolving the photocarrier build-up process in ReS\textsubscript{2}, an alternative way to study the photocarrier transfer is to monitor the decay of population in MoS\textsubscript{2}. For that purpose, we used a 3.06 eV pump pulse to inject photocarriers in the heterostructure and a probe pulse of 1.85 eV to monitor the population of carriers in MoS\textsubscript{2}. Carriers injected in ReS\textsubscript{2} are

![Fig. 4](image-url) Differential reflection of a 1.85 eV probe from MoS\textsubscript{2} monolayer (ML) with a 3.06 eV pump (solid squares) and MoS\textsubscript{2}–ReS\textsubscript{2} heterostructure with a pump of 1.53 eV (open squares).

![Fig. 5](image-url) Differential reflection signal of the 1.53 eV probe with 1.85 eV pump measured from MoS\textsubscript{2}–ReS\textsubscript{2} (solid green squares) and monolayer (ML) ReS\textsubscript{2} (open green squares). The solid blue squares are measured from MoS\textsubscript{2}–ReS\textsubscript{2} heterostructure with a pump of 3.06 eV and probe of 1.85 eV. The solid curves are fits to the data.
not sensed by this probe. The results are shown as the blue squares in Fig. 5. The signal decays single-exponentially with a time constant of 1.1 ± 0.2 ps, as indicated by the blue curve. Such a decay is significantly faster than that obtained from the MoS$_2$ ML (black squares in Fig. 4), which reflects the exciton recombination time. This fast decay is due to the transfer of photocarriers to ReS$_2$, and the time constant matches $\tau_T$, obtained by monitoring the build-up of population in ReS$_2$, very well.

Recent studies have revealed that charge transfer in type-II heterostructures formed by TMDs occurs on an ultrafast time scale, shorter than 100 fs, which can be facilitated by quantum coherence. The transfer process observed here is different from the previous studies in two aspects. First, in type-II heterostructures, electrons and holes transfer to opposite layers, and the Coulomb field is along the opposite direction of charge transfer. Here, the transfer is ambipolar, with both types of carriers moving together. Second, the previously studied heterostructures were formed by TMDs with the same 2H crystal structure. Here, the heterostructure is formed by 2H-MoS$_2$ and ReS$_2$ with a distorted 1T structure. It is possible that the different lattice structures suppress the coherent transfer process, and result in an incoherent, and thus slower, transfer process.

3 Conclusions

We have shown, both theoretically and experimentally, that MoS$_2$ and ReS$_2$ MLs form a type-I van der Waals heterostructure with both the CBM and VBM located in ReS$_2$. By selectively exciting photocarriers into the different layers of the heterostructure and probing the ReS$_2$ layer, we obtained evidence for carrier transfer from the MoS$_2$ layer to the ReS$_2$ layer. When probing the exciton resonance of MoS$_2$, a fast decay time in the heterostructure relative to ML MoS$_2$ is a further indication that both electrons and holes are transferring to the ReS$_2$. We observed that the transfer time of these carriers occurs on the order of 1 ps, which is significantly slower than the interlayer transport observed in previously studied TMD heterostructures with type-II band alignment. We attribute this to the incoherent nature of the transfer process between the two materials with different lattice structures. Overall, this observation of a type-I TMD heterostructure further proves the versatility of this family of materials to reach a broad range of applications, where band alignment plays a crucial role in charge transfer phenomena. The results also shed light on the fundamental understanding of charge transfer at van der Waals interfaces.

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