Dissecting Interlayer Hole and Electron Transfer in Transition Metal Dichalcogenide Heterostructures via Two-Dimensional Electronic Spectroscopy

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ABSTRACT: Monolayer transition metal dichalcogenides (ML-TMDs) are two-dimensional semiconductors that stack to form heterostructures (HSs) with tailored electronic and optical properties. TMD/TMD-HSs like WS₂/MoS₂ have type II band alignment and form long-lived (nanosecond) interlayer excitons following sub-100 fs interlayer charge transfer (ICT) from the photoexcited intralayer exciton. While many studies have demonstrated the ultrafast nature of ICT processes, we still lack a clear physical understanding of ICT due to the trade-off between temporal and frequency resolution in conventional transient absorption spectroscopy. Here, we perform two-dimensional electronic spectroscopy (2DES), a method with both high frequency and temporal resolution, on a large-area WS₂/MoS₂ HS where we unambiguously time resolve both interlayer hole and electron transfer with $34 \pm 14$ and $69 \pm 9$ fs time constants, respectively. We simultaneously resolve additional optoelectronic processes including band gap renormalization and intralayer exciton coupling. This study demonstrates the advantages of 2DES in comprehensively resolving ultrafast processes in TMD-HS, including ICT.

KEYWORDS: transition metal dichalcogenides, van der Waals heterostructure, interlayer charge transfer, ultrafast spectroscopy

S emiconducting monolayer transition metal dichalcogenides (ML-TMDs) are a class of two-dimensional crystalline materials with enticing electronic and optical properties, including a direct band gap in the visible range, strong light–matter coupling, enhanced excitonic correlations, and spin/valley locking. A growing area of TMD research involves vertical stacks of ML-TMDs, referred to as TMD heterostructures (HSs). Weak interlayer van der Waals forces between ML-TMDs bypass the lattice parameter matching constraints of conventional semiconductor HSs and preserve the electronic structure of each constituent, enabling the creation of new materials with tailored electronic and optical properties which differ from those of the isolated MLs.

Several TMD-HSs present a type II (staggered) band alignment, in which the valence band (VB) maximum and the conduction band (CB) minimum at the K/K′ points reside in different layers. This alignment favors charge separation and the formation of interlayer (IL) excitons via interlayer charge transfer (ICT). IL excitons are of particular interest as they feature recombination time scales (i.e., $1–100$ ns) orders of magnitude longer than those of intralayer excitons in isolated ML-TMDs (i.e., $\sim 100$ ps). The strong enhancement of the IL exciton lifetime, due to a reduced spatial overlap of the electron and hole wave functions, is responsible for the observation of several phenomena including the diffusion of IL excitons over micrometer length scales and extremely long valley polarization retention, among others. Enhanced IL exciton lifetimes make TMD-HSs with type II band alignment excellent candidates for optoelectronic and light-harvesting applications.

A variety of spectroscopic techniques, including photoluminescence and ultrafast transient absorption (TA) spectroscopy, have revealed ICT in TMD-HSs to be extremely rapid and very efficient. ICT has been previously reported to occur on a sub-100 fs time scale, though there remains uncertainty in the rates of ICT and a lack of information about differences in electron versus hole transfer times within the same system. Several studies on IL electron transfer have used above-resonance excitation, which introduces additional many-body effects into the system, and it is

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unclear how to disentangle these effects from ICT processes. Experimental studies show that the time scales and efficiencies of ICT are independent of the IL twist angle. Several theories as to how ICT proceeds without hindrance by interlayer momentum mismatch include mediation via local structural inhomogeneity, excess electronic energy, phonon-mediated intermediate scattering through hybridized valleys, and long-lived quantum coherences at the interface. Obtaining a comprehensive picture of ICT processes in TMD-HSs is of critical importance both for understanding the fundamental physics at play and for developing optoelectronic applications. One key experimental limitation in ultrafast TA studies of ICT thus far has been the trade-off between temporal and spectral resolution. Ultrafast TA spectroscopy is limited in its ability to distinguish between transitions in spectrally congested systems, where electronic transitions are close in energy. In fact, spectral selection of one transition requires narrowband pump pulses which, due to the Fourier transform (FT) limitation, reduce the temporal resolution of the experiment. Two-dimensional electronic spectroscopy (2DES) is a multidimensional spectroscopy technique that measures the third-order material polarization using a sequence of three pulses: two excitation pulses separated by the so-called coherence time, \( t_1 \), and a detection pulse that is delayed by the waiting time, \( t_2 \), with respect to the second excitation pulse (Figure 1a). FT over the coherence time \( t_1 \) allows for resolution over the excitation frequency axis, while resolution over the detection frequency axis is typically obtained by dispersing the detection pulse in a spectrometer. 2DES can obtain high excitation frequency resolution while using broadband pulses and thus simultaneously maximize spectral and temporal resolution. 2DES is very well-suited for studying ultrafast dynamics in systems with a high degree of spectral congestion and has been exploited to measure inhomogeneous/homogeneous line widths, excitonic coupling, exciton valley coherence, and biexcitons in ML-TMDs. Here, we use 2DES to simultaneously measure both interlayer electron and hole transfer dynamics in a large-area WS\(_2\)/MoS\(_2\) HS prepared using a novel mechanical exfoliation technique. This is achieved by using extremely short (sub-20 fs) and broadband pulses with spectra covering the A and B exciton of MoS\(_2\) and the A exciton of WS\(_2\). We observe signatures of intra- and interlayer coupling between the A/B excitons of MoS\(_2\) and the A exciton of WS\(_2\) immediately after photoexcitation. We unambiguously resolve interlayer hole transfer (IHT) to WS\(_2\) following selective excitation of the A exciton of MoS\(_2\) with a 34 ± 14 fs time constant and furthermore resolve interlayer electron transfer (IET) to MoS\(_2\) following excitation on resonance with the A exciton of WS\(_2\) with a 69 ± 9 fs time constant. We corroborate our findings by comparing these results with 2DES measurements of the individual layered components of the HS.

Figure 1 reports a sketch of the band alignment at the K point for the WS\(_2\)/MoS\(_2\) HS as calculated in ref 46. For clarity, we identify the A and B excitons of each layer of the HS with a subscript corresponding to the layer’s transition metal (AX and BX where X = Mo or W). Single crystal millimeter-scale ML-TMDs are prepared via gold-tape exfoliation of bulk crystals. The large-area WS\(_2\)/MoS\(_2\) HS is obtained by vertically stacking the MLs and transferring them onto a transparent 200-μm-thick SiO\(_2\) substrate. an interlayer twist angle of 29.5 ± 0.9° was determined by polarization-resolved second harmonic generation measurements (Figure S3). The absorption spectrum of the HS at 80 K is shown in Figure 1c. The 2DES experiments are performed at 80 K using a partially collinear pump–probe geometry and measuring the real absorptive third-order nonlinear signal. The home-built 2DES instrument generates the pump–probe pair with birefringent wedges, which scan the coherence time, \( t_1 \), with high phase stability.

2DES maps plot the correlation between excitation energy (\( \hbar \omega_1 \)) and detection energy (\( \hbar \omega_2 \)) for a given waiting time, \( t_2 \). We begin by examining the 2DES maps of the real absorptive signal of the ML-MoS\(_2\) (Figure 2a). The 2DES map at \( t_2 = 0 \)
displays two positive diagonal (ℏω1 = ℏω3) peaks corresponding to photobleaching (PB) of AMo and BMo excitons. At early times, the diagonal peaks are inhomogeneously broadened along the diagonal due to the spatial heterogeneity of the sample, while the homogeneous line width lies along the antidiagonal direction. Spectral signatures of the off-diagonal cross-peaks (ℏω1 ≠ ℏω3) between AMo and BMo at t2 = 0 fs indicate strong intralayer excitonic coupling, whereas the t2-dependent behavior reports on intralayer exciton dynamics and coupling strengths. Strong coupling between the A and B excitons in ML-MoS243,44,48 and other ML-TMDs38 has been previously observed and ascribed to several mechanisms: light-induced band gap renormalization (BGR),49 Dexter-like intervalley exciton coupling, and intervalley exchange interaction,43 among others.49,51,52 At later times, all four peaks exhibit horizontal elongation along the excitation axis, consistent with previous observations, using ultrafast TA, of a delayed excitonic PB following above-resonance excitation.53

The early temporal dynamics of the ML-MoS2 excitonic signatures are plotted as a function of waiting time, t2, in Figure 2b. With the exception of the above-diagonal cross-peak at (ℏω1,ℏω3) = (AMo, BMo) (pink circle), the signatures display an instantaneous rise and a double exponential decay within the 1 ps temporal window of the measurement. The ultrafast decay is consistent with previous observations and has been attributed to carrier thermalization and cooling.34 The above-diagonal cross-peak is characterized by a slightly delayed PB formation followed by a single exponential decay. Similar dynamics of the (AMo,BMo) peak were recently reported in a 2DES study of chemical vapor deposition grown ML-MoS2 using a circularly polarized configuration and attributed to an excitonic exchange interaction.52 Negative peaks in the 2D maps (blue, Figure 2a) are due to photoinduced absorption and are located at detection energies below the positive PB peaks. These features are attributed to many-body effects which are referred to cumulatively as BGR.44,48

The 2DES maps of the WS2/MoS2 HS (Figure 3a) show many distinct features compared to the ML-MoS2. At t2 = 0 fs, we see prominent diagonal peaks of the AMo and BMo PB, as in the ML-MoS2 sample, with an additional diagonal peak from the AW exciton. The AW peak is red-shifted in comparison to the AMo of the ML-WS2 (Figure S4) and is well separated spectrally from the AMo and BMo excitons. We attribute the spectral shift of AW to differences in the dielectric environment of the WS2 layer in the HS.55,56 At t2 = 0 fs, the WS2/MoS2 HS 2DES maps show significant broadening along the diagonal, due to overlap of the three inhomogeneously broadened excitons in close spectral proximity to one another. Similar to ML-MoS2, Figure 3a shows signatures of strong intralayer coupling between the AMo and BMo excitons, resulting in cross-peaks present at t2 = 0 fs. We are unable to resolve intralayer cross-peaks of the WS2 layer as the BW exciton falls outside our spectral detection window.

We can identify several spectral signatures unique to the HS which are the result of interlayer coupling and ICT processes. Several below-diagonal signatures of interlayer coupling are visible at t2 = 0 fs at (ℏω1,ℏω3) = (BMo,AW) and (AW,AMo). These signatures show an instantaneous rise similar to the diagonal peaks (Figure S7). At later times, additional interlayer cross-peaks grow in above the diagonal. The 2D map at t2 = 200 fs shows a grid of positive PB signals between every possible combination of the three excitonic states along with weak negative BGR signals red-shifted along ℏω3 from each excitonic resonance. The above-diagonal cross-peaks (green and cyan circles, Figure 3a) display a delayed PB signal. We assign these interlayer cross-peaks to ICT processes: IHT and IET for the peaks located at (ℏω1,ℏω3) = (AMo,AW) and (AW,AMo), respectively.

The delayed PB at (ℏω1,ℏω3) = (AMo,AW) (Figure 3b) is consistent with a dissociation of AMo excitons, leading to ultrafast hole scattering from the upper VB of the MoS2 to the higher-energy upper VB in the WS2 at the K′(K) point (Figure 3c). To confirm this assignment, we compare to the 2DES data of ML-MoS2 and ML-WS2. The t2 dynamics of the ML-MoS2 sample taken at the same excitation/detection energy as the IHT signature shows a completely different behavior: an instantaneous buildup and a negative signal due to the BGR upon AMo resonant photoexcitation (Figure S8). The isolated ML-WS2 sample (Figure S4) gives rise to a negligible signal because the excitation energy is below the optical gap of the

**Figure 3.** (a) 2DES maps of the real absorptive signal of WS2/MoS2 HS at 80 K for t2 = 0 fs (left) and 200 fs (right). Dashed lines indicate the energies of AMo (ℏω = 1.92 eV), AW (ℏω = 1.99 eV), and BMo (ℏω = 2.07 eV) excitons. The pump incident fluence is 13 μJ/cm2. The photoinduced carrier density is estimated to be ∼5.5 × 1012 cm−2 by considering the absorption coefficients of the individual layers. (b) Values of t2 dynamics of ICT signatures. Signatures of IHT following excitation of the AMo exciton (left) show a delayed rising component at the detection energy of AW. Signatures of IET following excitation of the AW exciton (right) show a delayed rising component at the detection energy of BMo. The t2 traces are shown with fits color coded for dots in (a). The (AW,WMo) t2 trace is multiplied by a factor of 2 for better comparison. (c) Cartoon illustrating the process of interlayer hole and electron transfer upon optical excitation of the MoS2 and WS2 layers, respectively.
We have focused on the two above-diagonal signatures of ICT for our analysis thus far, though two additional signatures of ICT are expected below the diagonal in Figure 3a. As described above, the below-diagonal ($A_{Mo}B_{Mo}$) and ($B_{Mo}A_{Mo}$) peaks are characterized by pulse-width-limited buildup dynamics with overlapping signals which are determined by both intralayer and interlayer scattering processes. The absence of delayed formation in the PB peaks of the below diagonal cross-peaks is a signature that the former of these processes prevails and determines the formation dynamics. This result is supported by recent experiments showing that electron–hole pairs photoexcited above the optical gap of TMDs relax on the bottom (top) of the CB (VB) at the K point on time scale faster than 20 fs.55

In conclusion, we have performed broadband 2DES on a large-area WS$_2$/MoS$_2$ HS, covering simultaneously the A and B excitons of MoS$_2$ and the A exciton of WS$_2$. We unambiguously observe IHT to WS$_2$ with 34 ± 14 fs time constant following excitation of the $A_{Mo}$ exciton and IET to MoS$_2$ with 69 ± 9 fs time constant following excitation of the $A_{Mo}$ exciton. Our results demonstrate the ability of 2DES, due to its unique combination of high temporal and spectral resolution, to distinguish between various excitonic dynamics in a spectrally congested TMD-HSs, overcoming the limitations of conventional TA spectroscopy. Continuing to implement 2DES on different large-area TMD-HS samples will allow us to characterize the extremely rapid exciton and charge carrier dynamics and their dependence on interlayer tilt angle, among many other variables.

**ASSOCIATED CONTENT**

*Supporting Information*

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.1c01098.

Two-dimensional electronic spectroscopy (2DES); 2DES setup; pulse characterization; large-area sample preparation and characterization of interlayer twist angle; 2DES map of isolated WS$_2$; fitting procedure and time constants; supplementary data (PDF)

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Author Contributions
X.Y.Z., S.D.C., and G.C. conceived the research. F.L. prepared the TMD-HS samples. V.R.P. and M.R. performed the 2DES experiments. V.R.P., M.R., C.T., M.M., and S.D.C. analyzed and interpreted the experimental data. Y.B. performed the SHG experiment. The paper was written by V.R.P. and S.D.C. with input from all authors. X.Y.Z. and G.C. supervised and coordinated the project.

Notes
The authors declare no competing financial interest.

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