

Picosecond energy transfer in a transition metal dichalcogenide–graphene heterostructure revealed by transient Raman spectroscopy

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Intense light-matter interactions and unique structural and electrical properties make van der Waals heterostructures composed by graphene (Gr) and monolayer transition metal dichalcogenides (TMD) promising building blocks for tunneling transistors and flexible electronics, as well as optoelectronic devices, including photodetectors, photovoltaics, and quantum light emitting devices (QLEDs), bright and narrow-line emitters using minimal amounts of active absorber material. The performance of such devices is critically ruled by interlayer interactions which are still poorly understood in many respects. Specifically, two classes of coupling mechanisms have been proposed, charge transfer (CT) and energy transfer (ET), but their relative efficiency and the underlying physics are open questions. Here, building on a time-resolved Raman scattering experiment, we determine the electronic temperature profile of Gr in response to TMD photoexcitation, tracking the picosecond dynamics of the G and 2D Raman bands. Compelling evidence for a dominant role of the ET process accomplished within a characteristic time of ~ 4 ps is provided. Our results suggest the existence of an intermediate process between the observed picosecond ET and the generation of a net charge underlying the slower electric signals detected in optoelectronic applications.

van der Waals heterostructures | Raman scattering | energy transfer | ultrafast spectroscopy | graphene

The van der Waals (vdW) heterostructures have recently emerged as a versatile platform to combine, in a single unit, key properties of layered materials. For instance, TMD–graphene (Gr) heterostructures have been thoroughly studied (1–3) as they represent a new class of truly two-dimensional metal-semiconductor junctions useful for engineering broadband, efficient, and ultrafast photodectors (4, 5) as well as narrow-line light-emitters (6). These systems jointly exploit the rich TMD photophysics (7–10) and their high photodetection efficiency (11) together with the unique electronic properties and subpicosecond photoresponse of Gr (12, 13).

Remarkably, vdW heterostructures feature atomically sharp heterointerfaces whereat efficient charge tunneling and/or energy funneling from the TMD to Gr layers can be achieved (4, 14–17). While a solid ground has been established to describe such near-field couplings in nanoscale systems (18, 19), the ultimate thinness of two-dimensional materials and their particularly strong Coulomb and light–matter interactions lead to new regimes where the shares of charge and energy transfer and the associated timescales are challenging to determine (14, 20).

In order to conceive high-performance optoelectronic devices based on vdW heterostructures, a key challenge consists of spatially and temporally tracking charge carriers and excitons in tightly coupled layers. Along these lines, early measurements on TMD-Gr heterostructures have employed transient absorption (TA) spectroscopy (16) and optoelectronic studies based on photoconductivity and photocurrent measurements (4). TA spectroscopy benefits from a short time resolution, ultimately limited by the laser pulse duration. Although accelerated picosecond transient dynamics of TMD excitons in TMD-Gr heterostructures has been observed (14-17, 21-24), TA does not make it possible to disentangle contributions from net charge tunneling and energy transfer, both when the latter arises from long-range dipole–dipole (Förster energy transfer) (25) or short-range exchange-mediated (Dexter energy transfer) (26) interactions. In electrically contacted devices, photoinduced charge transfer from the TMD to Gr leads to photogating (27). Such photogating processes (14, 28-30) are environment-dependent and slow, involving charge transfer to Gr and subsequent back-transfer to the TMD monolayer on timescales ranging from nanoseconds to seconds (31). Alternatively, direct photocurrent generation is observed on faster (subnanosecond) timescales in vertically biased Gr/few-layer

Significance

Hot carrier-based energy harvesting is critically implied in the performances of optoelectronic devices based on van der Waals heterostructures composed by graphene (Gr) and monolayer transition metal dichalcogenides (TMD). The way electron-hole couples initially photogenerated in the TMD are converted into an electric current in Gr is a controversial issue. In this work we identify the interlayer interaction occurring during the first picoseconds following photoexcitation as an energy transfer process that is much faster than (other) photogating phenomena implied in optoelectronic applications.

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TMD/Gr junctions. This mechanism, however, becomes quite inefficient when the active TMD region is thinned down to a monolayer (4), likely due to picosecond nonradiative transfer to Gr.

A key technique for characterizing vdW heterostructures is Raman spectroscopy. The Raman spectrum of Gr is dominated by two features, known as the G and 2D modes, near 1,580 and 2,600 cm⁻¹, respectively. The G- and 2D-mode processes arise from nonresonant inelastic scattering with one zone-center longitudinal and transverse optical phonon and from a resonant process involving a pair of near-zone edge transverse optical phonons, respectively (32-35). Both modes are sensitive to external perturbations and in particular to doping (36-38) and temperature rises (39-41). For instance, the integrated intensity of the 2D-mode feature (I_{2D}) is routinely used to determine the Fermi level in Gr (34, 38, 42) as well as of the defect density (43) and the electronic temperature (41, 44) as summarized in Eq. 1. Building upon this sensitivity, quantitative Raman-based methods have been developed to characterize Gr-based systems. Recently, a combination of photoluminescence (PL) spectroscopy and steady state spontaneous Raman spectroscopy have revealed the interplay between slow, extrinsic photoinduced net charge transfer and picosecond energy transfer, pointing toward the dominant role of energy transfer in TMD-Gr heterostructures (14), without, however, determining the associated temporal dynamics.

Extending such sensing capabilities to the out-of-equilibrium regime would offer a unique way to address the ultrafast response of Gr following photoexcitation of the TMD above its optical bandgap. Indeed, both charge and energy transfer processes may in principle generate hot carriers in Gr, leading to an increase in its electronic temperature $T_{\rm e}$, which can be quantified by monitoring hot PL and out-of-equilibrium Raman scattering. The former directly yields T_e (45), while the latter is sensitive to both transient Fermi level shifts and temperature rises (46, 47). Assuming that photoexcitation of the TMD-Gr heterostructure predominantly yields a net charge flow (i.e., charge transfer) across the coupled layers, a long-lived doped state is expected to develop in Gr after electron-phonon relaxation, affecting the line shape and the intensity of the G-mode and 2D-mode features. In contrast, in an energy transfer-dominated regime, Gr would only exhibit a transient increase of T_e , the dynamics of which is governed by the interplay between energy transfer and phonon relaxation of the electron-hole (e-h) pairs injected in Gr occurred in the subpicosecond timescale (12, 45, 48, 49). This scenario would result in a subpicosecond decrease of I_{2D} combined with a broadening of the G-mode feature. Charge and energy transfer can thus be distinguished based on the transient dynamics of the Raman features of Gr.

Here we provide direct mechanistic and temporal insights into interlayer coupling in monolayer tungsten disulfide (WS₂)–Gr heterostructures by exploiting recent developments in ultrafast Raman spectroscopy (44, 46, 47, 50). Specifically, extracting the temporal profile of T_e in Gr following optical excitation of the WS₂ monolayer slightly above its optical bandgap, we demonstrate highly efficient energy transfer from TMD to Gr in a 4-ps timescale, with no evidence for a net charge transfer.

Results

Our model system to track ultrafast interlayer transfers is a vdW heterostructure made from a Gr monolayer stacked onto WS_2 monolayer supported by a silica substrate. In our experiments, the sample is probed using 1-ps pulses at 1.58 eV, i.e., around 400 meV below the WS_2 optical bandgap. The role of the probe pulse

is twofold: 1) it induces a combination of linear and nonlinear PL, which we use as an imaging tool for preliminary mapping the heterostructure, and 2) it serves as a time-delayed Ramaninducing beam following photoexcitation by 1-ps pulses at 2.07 eV, slightly above the WS₂ optical bandgap. The time duration of the probe pulse is imposed by the necessary trade-off between a pulse duration (δt) as short as the exciton lifetime in the WS₂ and a narrow excitation bandwidth ($\delta \nu$) required to yield a sufficient spectral resolution (these two quantities are fundamentally constrained by the Fourier relation: $\delta \nu \delta t \geq 14.7 \text{ cm}^{-1}\text{ps}$). The experiment concept is shown in Fig. 1 *A* and *D* and further detailed in *Materials and Methods*.

PL. PL spectroscopy is one of the main ways to characterize the TMD–Gr coupling. Bare TMD monolayers exhibit indeed strong PL: the radiative recombination upon optical excitation is favored by the presence of a direct bandgap in absence of competitive nonradiative decay channels (51, 52). Conversely, Gr does not exhibit significant PL (53) as efficient nonradiative decay channels (electron–phonon coupling) are active at any excitation wavelength owing to its gapless band structure (54). Accordingly, massive quenching of the TMD PL indicates efficient near-field coupling to Gr (4, 14–16, 55) (see also *SI Appendix*, Fig. S4).

A different scenario arises when photoexciting Gr with ultrashort laser pulses. Under large photon fluxes, the nonradiative recombination channel is significantly depleted, and hot PL (albeit weaker than the TMD PL) is observed from Gr (45, 56). Here we take advantage of such transient regime to introduce an imaging protocol, based on the PL detection upon pulsed laser excitation, for simultaneously mapping Gr and WS₂. Specifically, photoexciting the WS₂–Gr heterostructure below the WS₂ optical bandgap using our probe pulse, the WS₂ PL stems from a lowerefficiency two-photon absorption process (*SI Appendix*, Fig. S6), whose intensity is comparable (as shown in Fig. 1*C*) with the Gr hot PL and is emitted in a different spectral region.

By recording a map of the integrated PL intensity in distinct spectral regions selected to maximize the contrast (Fig. 1*B*), we are able to isolate the regions with Gr and WS₂ only, as well as the heterostructure with strongly and weakly coupled areas. Such simultaneous mapping, not achievable with continuous wave (cw) excitation, is a powerful method for imaging Gr-based heterostructures. The green contour of Fig. 1*B* highlights the region of coupled WS₂–Gr, characterized by PL quenching relative to bare WS₂ and by the presence of hot PL from the coupled Gr monolayer (Fig. 1*C*). The Raman measurements presented hereafter are performed in this coupled region of the heterostructure.

Time-Resolved Raman Scattering. In our experiments the Raman signal induced by probe pulses is used to monitor Gr response following optical excitation by the higher-energy pump photons absorbed by the WS₂ monolayer. Probe pulses also lead to an out-of-equilibrium increase of the electronic temperature $T_e^{\rm pr} \sim 1,550$ K in Gr (*Materials and Methods*), due to the e–h pairs injected upon absorption ($A_{\rm Gr} \sim 2.3\%$). In contrast, the pump pulses are mainly absorbed by the WS₂ layer [optical absorptance $A_{\rm X} \sim 5.4\%$ at 2.07 eV in the same preparation condition (57, 58)], generating excitons in WS₂. However, a fraction of the pump pulse is absorbed also by the Gr layer [$A = (1 - A_{\rm X})A_{\rm Gr} \sim 2.2\%$] and induces a photoinduced electronic heating of Gr (45) (for 70 μ W, $T_e < 1,400$ K [*SI Appendix*, Fig. S8], implying an e–h pair density $n_{\rm eh} < 4.5 \times 10^{11}$ cm⁻²; *SI Appendix*, Fig. S9). Importantly, probe pulses have a negligible interaction with WS₂, as testified by the weak two-photon fluorescence (*SI Appendix*, Fig. S6).



Fig. 1. Optical characterization of a WS 2-Gr heterostructure. (*A*) Time-delayed pump and probe beams focused onto a diffraction-limited WS2-Gr spot. (*B*) PL map generated by probe only (1.58 eV) after integrating the emission spectra in the ranges 1.88 to 2.05 eV (WS2, two-photon PL) and 1.65 to 1.85 eV (Gr, hot luminescence) and the corresponding optical image (*Right*). Contour lines indicate bare Gr (blue), few-layer Gr (yellow), WS2 (red), coupled (green), and uncoupled (black) WS2-Gr regions. (*C*) PL spectra for the spots indicated in the map with corresponding colors. (*D*) Time-resolved, probe-generated Raman spectra of Gr modes in the WS2-Gr heterostructure for two time delays, corrected from hot PL background (*Materials and Methods*).

Fig. 2 shows time-resolved Raman spectra of the WS₂-Gr heterostructure compared with reference measurements performed on the nearby bare Gr region. Our measurements without pump pulse reveal a slightly lower value of I2D in WS2-Gr, with respect to Gr, due to static charge transfer to Gr (dashed line in Fig. 2G), in agreement with cw experiments (SI Appendix, Fig. S5D and refs. 6, 14). As shown in Figs. 1D and 2G, the integrated area of the 2D-mode feature depends on the time delay. This effect appears prominently in Fig. 2 C and D, where the differential 2D mode spectra (upon subtraction of the Raman spectra recorded at a $\Delta t = -30$ ps) exhibit a clear intensity depletion. Moreover, Fig. 2G indicates the presence of a small pump-induced heat transfer from the electronic to the phononic degree of freedom, rising I_{2D} above the pump off value (dashed lines in Fig. 2G) at large (due to transient D phonon heating) and at negative (increased lattice temperature) Δt . In contrast, the differential spectra of the G-mode feature (Fig. 2C and SI Appendix, Fig. S12) do not exhibit any sizeable transient intensity change. Remarkably, while the transient decrease of I_{2D} sets in and decays almost entirely within our temporal resolution in the Gr reference, the same effect decays on a longer timescale of $\sim 4 \text{ ps}$ in WS₂–Gr.

Discussion

The PL quenching of WS₂ may stem either from charge transfer or from energy transfer. Here we numerically simulate the Raman experimental results considering only energy transfer from WS₂ to Gr with near-unity efficiency.

We first focus on the time-resolved Raman response of bare Gr. In this case, the fast drop and recovery of I_{2D} can be ascribed to the e-h pair density instantaneously photogenerated upon pump absorption and relaxing on an ultrafast (resolution-limited) timescale, followed by a slower D phonon heating. In order to isolate the latter, the Gr experimental profiles have been preliminarily fitted as the sum of a Gaussian dip and an exponential saturation term (Fig. 2), which takes into account the electronic and subsequent lattice heating, respectively. The latter (ΔI_{2D}^{ph}), expected to be weakly sensitive to the WS₂ coupling, is reported in Fig. 2*G*. The genuine modification of the 2D area induced by the electronic heating (I_{2D}^{e}) is hence obtained for both Gr and WS₂– Gr by subtracting the phononic term, as $I_{2D}^{e} = I_{2D} - \Delta I_{2D}^{ph}$.

Gr by subtracting the phononic term, as $I_{2D}^e = I_{2D} - \Delta I_{2D}^{ph}$. In order to quantitatively evaluate the e-h pair density injected by the probe pulse and the corresponding T_e increase, the relation between the Gr electronic properties and the I_{2D} can be exploited (42):

$$\begin{split} I_{2\mathrm{D}}^{\mathrm{e}} &\propto \left(\frac{\gamma_{\mathrm{K}}(E)}{\gamma(E, T_{\mathrm{e}})}\right)^{2} = \\ &= \left(\frac{\gamma_{\mathrm{K}}(E)}{\gamma_{\mathrm{K}}(E) + \gamma_{\Gamma}(E) + \gamma_{\mathrm{def}}(E) + \gamma_{\mathrm{ee}}(E, T_{\mathrm{e}})}\right)^{2}, \end{split} \tag{1}$$

where *E* denotes the energy of the electronic state and 2γ is the scattering rate of electrons and holes, equal to the sum of $\gamma_{\rm K}$, γ_{Γ} , $\gamma_{\rm def}$, and $\gamma_{\rm ee}$, which are the scattering rate of electrons with 2D-mode and G-mode phonons near the K and Γ points of the Brillouin zone, defects, and other electrons, respectively. Notably, $\gamma_{\rm ce}$ is the only term that depends on $T_{\rm e}$ (36, 37, 39, 42, 43, 59). Specifically (60),

$$\gamma_{\rm ee}(E, T_{\rm e}) = \alpha T_{\rm e}, \qquad [2]$$

with $\alpha = 0.51 \text{ cm}^{-1} \text{ K}^{-1}$ for E > 100 meV (46).

From the full width at half maximum (FWHM) of 2D mode ($\Gamma_{\rm 2D}$) measured without pump pulse, we can estimate $\gamma \approx 195 \ {\rm meV}$ for WS₂–Gr at $T_{\rm e} = T_{\rm e}^{\rm pr}$ (35). In the presence of pump pulse, an increase of $T_{\rm e}$ is expected. However, for negative or large pump–probe delays (> 4 ps) the electronic heating of Gr



Fig. 2. Time-resolved Raman spectra. (A and B) Pump-off Raman spectra of the G and 2D modes for WS2-Gr and bare Gr. Transient differential Raman spectra $\Delta I(\Delta t, E) = I(\Delta t, E) - I(-30ps, E)$: (C and D) color maps and (E and F) vertically offset slices for selected time delays in ps. The 2D mode intensity I_{2D} , as opposed to I_{G} , decreases around zero delay. This effect is observed to a lesser extent in bare Gr. (G) While I_{2D} drop—due to electronic heating recovers in bare Gr within a timescale comparable with the pump-probe temporal overlap (black symbols and guideline), it takes longer in WS2-Gr (blue symbols and guideline). The dashed blue (black) lines show I_{2D} without the pump beam in WS₂-Gr (bare Gr), indicating the presence of a small photoinduced phonon heating. The bare Gr profile has been modeled (red thick line) with a picosecond drop (fast electronic term broadened by instrumental resolution) and its convolution (
) with an exponential term (transient phononic contribution; red thin line): $f(\Delta t) = C + A \exp \left| -\Delta t^2 / (2\sigma^2) \right| + \Delta l_{2D}^{\text{ph}}(\Delta t)$, where $\Delta l_{2D}^{\text{ph}}(\Delta t) = B \exp\left(-\Delta t^2/2\sigma^2\right) \circledast \left[\theta(\Delta t) \left(1 - e^{-(\Delta t)/\tau}\right)\right]$. *A*, *B*, and *C* are fitting parameters. $\sigma = 0.66$ ps corresponds to the autocorrelation of the 1-ps FWHM pump and probe pulses, and $\tau = 5$ ps from ref. 39. $\Delta l_{2D}^{\rm ph}$ is also reported with a vertical offset to emphasize its role in the WS2-Gr case (dotted red line). The accuracy in the I_{2D} measurement is ~ 1%.

induced by the direct interaction with pump pulse is not observed within the probe pulse duration due to the ps recombination of the e-h pairs. The electronic heating effect can only be observed around the time overlap of pump and probe pulse, generating a drop in I_{2D}^e (Fig. 2*G*). Taking into account this experimental evidence and Eq. **1**, the temporal profile of T_e in WS₂–Gr and in bare Gr can be extracted (Fig. 3*A* and *Materials and Methods*).

As shown in Fig. 3*A*, the additional energy flux injected through energy transfer from the WS₂ layer to Gr leads to a different T_e profile as compared to reference measurements in bare Gr, taking into account that the energy associated with a WS₂ exciton transfer is ~ 2 eV, and using the Fermi distribution with the density of states in Gr, we can derive the relation between the additional e–h pair density induced by the pump pulse and the relative increase of T_e above T_e^{pr} (right axis of Fig. 3*A* and *Materials and Methods*).

At this point, we model the dynamics of e–h pairs in Gr taking into account the transfer pathways reported in Fig. 3B. The pump pulse creates a population of e–h pairs in the WS₂ and Gr monolayers proportional to their absorptances. The WS₂ layer with Gr on top exhibits a fast decay channel (τ_T), which inhibits its much longer decay $\tau_0 \sim 100 \text{ ps} - 1 \text{ ns}$ (14–16, 61) and exciton–

exciton annihilation (6). Due to the suppression of this slow linear recombination in bare WS₂, the temporal evolution of the exciton density in WS₂, $N_X(t)$, can be described by the following differential equation:

$$\frac{dN_{\rm X}}{dt} = -\frac{N_{\rm X}}{\tau_{\rm T}} + A_{\rm X}I_{\rm PUP}(t),$$
[3]

where $I_{PUP}(t)$ is the temporal profile of the pump pulse, assumed Gaussian with a FWHM of 1 ps.

On the other hand, Gr exhibits an intrinsic decay time for the thermalization of the electronic system with the phononic bath and substrate (τ_G). Interlayer energy transfer induces e–h pair injection in Gr ($N_G(t)$), leading to a rise in T_e . Its temporal evolution can be expressed as

$$\frac{dN_{\rm G}}{dt} = -\frac{N_{\rm G}}{\tau_{\rm G}} + \frac{N_{\rm X}}{\tau_{\rm T}} + AI_{\rm PUP}(t), \qquad [4]$$

where the coupling term $\frac{N_{\rm X}}{\tau_{\rm T}}$ obviously vanishes in the bare Gr case. Performing a best fit of the experimental data reported in Fig. 3A with $N_{\rm G}(t)$ obtained from Eqs. 3 and 4, convoluted with the 1-ps probe pulse temporal profile, we obtain the black and blue solid lines for bare Gr and WS₂–Gr, respectively, with characteristic times $\tau_{\rm G} = 0.6$ ps and $\tau_{\rm T} = 4.3$ ps.

The decay time $\tau_{\rm G}$ in bare Gr is close to the temporal resolution of our setup and is consistent with previous reports on Gr (62– 64). Most importantly, we are able to determine the relevant energy transfer timescale from WS₂ to Gr at room temperature. Remarkably, $\tau_{\rm T} \approx 4$ ps is longer than $\tau_{\rm G}$, indicating that energy transfer–mediated e–h pair injection in Gr (the term $\frac{N_{\rm X}}{\tau_{\rm T}}$ in Eq. 4) is responsible for the longer decay of $N_{\rm G}(t)$ observed in WS₂–Gr.

Since we are investigating a system separated by a subnanometer vdW gap, Förster and Dexter energy transfer are both likely to occur. The $\tau_{\rm T}$ value that we directly determine here, however, provides a possible rationale to previous observations of PL quenching (14) and TA spectroscopy (15–17) and agrees with theoretical calculations based on a dominant Förster-type energy transfer mechanism (25).

Notably, the WS₂ PL quenching and the I_{2D} drop (42) may be compatible with a charge transfer mechanism involving a net transient flux of electrons (or holes) to Gr. Critically, in this scenario, the doping generated in Gr upon charge transfer would be expected to relax on a timescale much longer than the picosecond dynamics reported in Fig. 2*G* (14, 31). Moreover, the measured temporal profile of the G band FWHM is also not compatible with a charge transfer–dominated scenario, as detailed in *SI Appendix*.

The energy transfer nature of the process established here suggests that the photogating effects (14, 28) and associated photodetection capabilities demonstrated in TMD–Gr-based optoelectronic devices (27, 31) result from an additional conversion mechanism into a much slower, less efficient, net charge transfer. Moreover, this evidence confirms the dominant contribution of energy transfer, measured with Raman spectroscopy (14), and at the same time the ps simultaneous transfer of e–h of the exciton provides an explanation for the low internal quantum efficiency of photocurrent generation in photodetectors made from TMD monolayers (4).

Conclusion

We have measured the time-resolved Raman spectrum of Gr in a WS_2 -Gr heterostructure upon resonant photoexcitation of band-edge WS_2 excitons. By comparing the Raman response



Fig. 3. Modeling energy transfer in a WS₂-Gr heterostructure. (A) T_e and e-h pair density at different time delays extracted from the dynamics of I_{2D} for WS₂-Gr (open symbols) and bare Gr (full symbols) are compared with the simulated profiles (solid lines). The uncertainty on the T_e is 20 K. (B) Sketch of the kinetic model used in the simulation. The pump pulse can 1) generate an exciton population in WS₂ (thick orange arrow) and 2) populate the electronic states of Gr with e-h pairs (thin orange arrow). The e-h pairs (represented as blue and white circles, respectively) in Gr decay with a timescale τ_G . In contrast, the excitons in bare WS₂ have a long lifetime τ_0 . Exciton decay is strongly accelerated in WS₂-Gr due to energy transfer to Gr with a characteristic time τ_T .

of bare and WS₂-coupled Gr, we have unveiled a slower relaxation of the 2D mode integrated intensity occurring on a 4-ps timescale in WS2-Gr. Our experimental data are rationalized by a kinetic model based solely on energy transfer. Although photocurrent generation occurs with relatively high efficiency in vertically biased TMD-Gr heterojunctions (4, 65) made from few-layer TMD films, our results strongly suggest that net photoinduced charge transfer has negligible efficiency in ultimately thin monolayer TMD-Gr heterostructures, explaining the low efficiency of associated photodevices. Instead, we demonstrate that a TMD monolayer can boost carrier injection in Gr through picosecond energy transfer, with a room-temperature transfer yield approaching unity considering the longer excitonic lifetimes in bare TMD monolayers. Besides showing the possibility to exploit photothermionic effects for efficient hot carrier injection from Gr to TMDs under subbandgap excitation (5, 17), our results provide an essential step for the microscopic understanding of ultrafast interlayer coupling in TMD-Gr heterostructures, whose relevance for optoelectronics is thus consolidated. They also indicate as a key technological challenge the efficient funneling of hot carriers generated in Gr before they release their energy into heat, either through Gr phonon emission or through coupling with the substrate.

Materials and Methods

Sample Preparation. Our WS₂–Gr heterostructure was fabricated as described previously (14) using a dry, viscoelastic transfer technique (66). In brief, a WS₂ monolayer and a Gr monolayer were mechanically exfoliated from bulk crystals and stacked onto a fused silica substrate using a home-built transfer station. The sample was characterized using cw PL and Raman spectroscopies in ambient conditions (see *SI Appendix*, Figs. S4 and S5, respectively) before performing time-resolved studies.

PL Measurements. The experimental setup is the same as the time-resolved Raman experiment, but in this case, only the probe pulse is sent on the sample. The PL spectra reported in Fig. 1 *B* and *C* are collected in 2 s per pixel, and the sample scan is performed with two mechanical stages. This experimental scheme allows us to perform a simultaneous imaging of the two monolayers. Moreover, using the nonlinear two-photon PL for WS₂, the spatial resolution is higher as compared to measurements in the cw regime performed using the same optical elements.

Gr, having no band gap (67), is not expected to have any radiative decay channel for charge carriers. However, under high excitation densities it emits light (45) over a wide spectral range owing to the inhibition of a nonradiative recombination channel. Indeed, in an out-of-equilibrium configuration, the large density of charge carriers in the conduction band cannot fully relax down to the

Fermi level via electron-phonon decay pathways. Thus, the electronic subsystem also relaxes radiatively through hot PL, which is well described by Planck's law. Hence, in order to extract the electronic temperature (T_e), we fit the hot PL spectra of Gr with Planck's law (black lines in *SI Appendix*, Fig. S7):

$$I(\hbar\omega, T_e) = \eta_{em} \tau_{em} E(\hbar\omega) \frac{\hbar\omega^3}{2\pi^2 c^2} \frac{1}{e^{\frac{\hbar\omega}{kT_e}} - 1},$$
[5]

where $\eta_{\rm em}$ is the emissivity of the blackbody, $\tau_{\rm em}$ is the emission time of the blackbody, and $E(\hbar\omega)$ is the responsivity of the detection chain taking into account the efficiency of the charge-coupled device, which depends on the photon energy. $\hbar\omega$ is the energy of the emitted photon, T_e is the electronic temperature, and k is Boltzmann's constant.

In *SI Appendix*, Fig. S7, we show the hot PL spectra obtained performing a probe pulse only experiment on Gr (b) and on WS₂–Gr (a) at different powers of the probe pulse. The peak located near 1.8 eV is present in both Gr and in WS₂–Gr and is assigned to the anti-Stokes Raman G mode, while the broad peak at around 2 eV is assigned to PL from the A-exciton of WS₂ excited by a two-photon absorption process.

In *SI Appendix*, Fig. S8, we show the temperature extracted from fit as a function of PRP powers for both bare Gr (diamond) and WS₂–Gr (square). From the same figure, we estimate that the probe pulses (with power of 210 μ W) used in our time-resolved Raman scattering experiment lead to an electronic temperature $T_e^{pr} \sim 1,550$ K. The temporally overlapped pump pulses induce an additional temperature increase, which originated from both direct electronic heating of Gr and energy transfer from WS₂.

Time-Resolved Raman Experiments. We use a two-module Toptica FemtoFiber Pro source as in ref. 50. This setup is able to generate, at a repetition rate of 40 MHz, 1-ps probe pulses at 1.58 eV (PRP) and a supercontinuum (SC) output between 0.89 and 1.38 eV. The SC spectral intensity can be tuned with a motorized Si-prism pair compressor. A periodically poled lithium niobate crystal with a fan-out grating (a poling period changing along the transverse direction) is exploited to produce broadly tunable (from 1.97 to 2.30 eV) narrowband 1-ps pump pulses, with a power < 10 mW (68). A dichroic mirror is used to combine the two beams, whose relative temporal delay is tuned with a motorized optical delay line. A long-working distance 20× objective (numerical aperture NA = 0.4) focuses the pulses onto the same ${\sim}2~\mu m$ spot on the sample. Before objective, the probe and pump power are 210 and 70 μ W, respectively. The Raman emission on the Stokes side is collected by a condenser (numerical aperture 0.75), and the pump-probe pulses are filtered out using interference filters. The Raman spectrum is measured with a monochromator (Acton Spectra Pro-2500i) coupled to a cooled charge-coupled device array (CCD Princeton Instruments Pixis 100).

In Fig. 2 the Stokes Raman signals of Gr generated by PRP are reported at several time delays, first on Gr and then on WS₂–Gr. For each delay we acquire Raman spectrum for 10 min and repeat the acquisition six times. In *SI Appendix*, Fig. S10, we show a typical spectrum in WS₂–Gr with and without pump. Raman spectra are affected by an intense background due to the substrate. In order to remove this background, we collect, with same acquisition time, the signal spectrum from the substrate, and we subtract it (with a scaling of the intensity) from the Raman signal. In *SI Appendix*, Fig. S11*A*, we show two selected spectra (pump off and 0.1 ps) in which we have removed the substrate background. In order to better clean data we remove a baseline (red) and get the spectra as in *SI Appendix*, Fig. S11*B*. We do the same procedure on the spectra obtained on Gr alone. The two peaks at \sim 1,585 and \sim 2,620 cm⁻¹ are the G and 2D modes, respectively.

*T*_e and e-h Pair Density Calculation. To extract the *T*_e from Eq. **1**, the value of $\gamma(T_e^{\rm pr})$ is required, which can be extracted from the FWHM of the 2D-mode feature, Γ_{2D} (35), which writes

$$\Gamma_{\rm 2D} = 4\sqrt{2^{2/3} - 1} \frac{1}{2} \frac{\partial \omega_{\rm 2D}}{\partial (h\nu_{\rm laser})} \gamma(T_{\rm e}), \qquad [6]$$

where ω_{2D} is the 2D-mode frequency, $[\partial \omega_{2D}/\partial (h\nu_{laser})]/2 = \frac{1}{ch}v_{ph}/v_F \sim 100$ cm⁻¹ eV⁻¹ (32, 69), i.e., the ratio between the phonon and Fermi velocity, defined as the slope of phononic (electronic) dispersion at the phonon (electron) momentum corresponding to a given excitation laser energy $h\nu_{laser}$ (32). *c* and *h* indicate the speed of light and the Planck constant, respectively. However, Eq. **6** does not consider the bimodal spectral feature of the 2D mode, due to the contributions of so-called inner and outer processes (70, 71). For this reason, we isolate each contribution, following the procedure reported in ref. 69. The fitted bandwidth of the 2D mode contributions, obtained in cw measurements at room temperature, is 19 cm⁻¹, which we use to estimate $\gamma(300 \text{ K}) = 940 \text{ cm}^{-1}$. Considering Eq. **2**, $\gamma(T_c) = 1,600 \text{ cm}^{-1}$ at $T_c^{\text{pr}} = 1,550 \text{ K}$, as estimated above. To evaluate T_c from T_{2D}^{p} , Eq. **2** is replaced in Eq. **1**, obtaining

$$I_{\rm 2D}^{\rm e}(T_{\rm e}) \propto \left(\frac{\gamma_{\rm K}(E)}{\gamma(T_{\rm e}^{\rm pr}) + \alpha(T_{\rm e} - T_{\rm e}^{\rm pr})}\right)^2.$$
 [7]

The ratio between I_{2D}^{e} at different T_{e} is

$$\frac{I_{2D}^{e}(I_{e})}{I_{2D}^{e}(I_{e}^{pr})} = \left(\frac{\gamma(I_{e}^{pr})}{\gamma(I_{e}^{pr}) + \alpha(I_{e} - I_{e}^{pr})}\right)^{2}.$$
[8]

 $T_{\rm e}$ extracted from the previous equation is

$$I_{\rm e} - I_{\rm e}^{\rm pr} = \frac{\gamma(I_{\rm e}^{\rm pr})}{\alpha} \left(\sqrt{\frac{I_{\rm 2D}^{\rm e}(I_{\rm e}^{\rm pr})}{I_{\rm 2D}^{\rm e}(I_{\rm e})}} - 1 \right).$$
 [9]

Summing up, Eq. **9** is obtained from the dependence of I_{2D}^c on T_c (Eq. **7**). The ratio between I_{2D}^e at different temperatures allows us to remove the dependence on $\gamma_K(E)$ (Eq. **8**), and T_c can be conveniently isolated (Eq. **9**), considering that $\gamma(T_e^{pr}) = 1,600 \text{ cm}^{-1}$. However, the model in Fig. 3*A* is designed to reproduce the e-h pair dynamics. Consequently, a conversion of T_c in e-h pairs (n_{ch}) is required. In our experimental configuration the pump photon energy (E_{PUP}) matches the TMD bandgap ($\sim 2 \text{ eV}$), and therefore, the energy of each e-h pair

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was either generated from the pump photons or transferred from WS_2 . Hence, the carrier density is related to the total energy density (per unit surface) associated to a given T_c as

$$n_{\rm eh} = \frac{\varepsilon_{\rm TOT}(I_{\rm e})}{E_{\rm PUP}} = \frac{1}{E_{\rm PUP}} \int_{-\infty}^{\infty} g(E)f(E, E_{\rm F}, T_{\rm e})dE =$$

$$= \frac{1}{E_{\rm PUP}} \int_{-\infty}^{\infty} \frac{2|E|}{\pi \hbar^2 v_{\rm F}} \frac{1}{e^{(E-E_{\rm F})/(kT)} + 1} dE,$$
[10]

where g(E) is the density of states in Gr, $f(E, E_F, T_e)$ is the Fermi-Dirac distribution, E_F and v_F are the Fermi energy and Fermi velocity, and k is the Boltzmann constant. Spontaneous Raman measurements (*SI Appendix*, Fig. S5) indicate a relatively low doping below 6×10^{11} cm⁻², i.e., a Fermi level $|E_F| < 100$ meV at room temperature. Moreover, under our experimental condition, the probe pulse alone induces $T_e = T_e^{\text{pr}}$ in Gr, further reducing the upper bound for the Fermi level down to 30 meV (72). In *SI Appendix*, Fig. S9, we report n_{eh} as a function of T_e for $E_F = 0$ meV and $E_F = 30$ meV. Such a doping effect mainly results in a minor vertical shift, not relevant for the incremental carrier generation N_G (additional to the probe-induced n_{eh}) associated with a temperature rise from T_e^{pr} to the pumpinduced time-dependent T_e (Fig. 3*B*).

Data Availability. All study data are included in the article and/or SI Appendix.

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