Ultrafast formation of interlayer hot excitons in atomically thin MoS$_2$/WS$_2$ heterostructures

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Van der Waals heterostructures composed of two-dimensional transition-metal dichalcogenides layers have recently emerged as a new family of materials, with great potential for atomically thin opto-electronic and photovoltaic applications. It is puzzling, however, that the photocurrent is yielded so efficiently in these structures, despite the apparent momentum mismatch between the intralayer/interlayer excitons during the charge transfer, as well as the tightly bound nature of the excitons in 2D geometry. Using the energy-state-resolved ultrafast visible/infrared microspectroscopy, we herein obtain unambiguous experimental evidence of the charge transfer intermediate state with excess energy, during the transition from an intralayer exciton to an interlayer exciton at the interface of a WS$_2$/MoS$_2$ heterostructure, and free carriers moving across the interface much faster than recombining into the intralayer excitons. The observations therefore explain how the remarkable charge transfer rate and photocurrent generation are achieved even with the aforementioned momentum mismatch and excitonic localization in 2D heterostructures and devices.

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Two-dimensional (2D) materials, including graphene, hexagonal boron nitride, layered transition-metal dichalcogenides (TMDCs), as well as black phosphorus, have demonstrated a wide variety of unique optical, electrical, thermal and mechanical properties. Atomically thin opto-electronic devices produced by stacking layers of 2D materials to form Van der Waals (VDW) heterostructures have demonstrated fascinating effects, including high mobility, high carrier inhomogeneity and high on/off ratios in vertical tunnelling transistors.

Heterostructures composed of TMDC monolayers are of particular interest for fabricating light-harvesting and photodetectors. TMDCs are known as the direct bandgap semiconductors with remarkably strong light–matter interactions. The efficient separation of photo-excited electrons/holes and generation of across-layer flow of carriers and photocurrent were observed in TMDC heterostructures. The observations further corroborate the promising perspective of the heterostructures as atomically thin opto-electronic devices.

Here, to unravel this fundamentally important puzzle, we carry out ultrafast visible/infrared (IR) microspectroscopy studies on the cross-layer charge transport dynamics in WS2/MoS2 heterostructure. Different from the visible detection light used in previous works, the energy of mid-IR detection pulse is well below the thermal energy of the intermediate state (denoted as hot exciton) available allows the sampling of a broader range of momentum space than that determined by the conduction band minimum (CBM) and valence band maximum (VBM) of the stacked heterojunction. Momentum conservation can therefore be overcome. At the same time, the excess energy in the hot intermediates leads to lower-binding energy and a larger electron–hole distance than those of the interlayer excitons. As the result, the intermediates may easily dissociate and contribute to the photocurrent in the presence of the internal/external potentials before relaxing to the interlayer excitons. The results therefore reveal the mechanism how the momentum mismatch and excitonic localization in 2D heterostructures are overcome to reach fast and efficient charge transfers, and photocurrent generation.

Results

Charges can efficiently transfer between MoS2 and WS2. One optical image of the atomically thin MoS2/WS2 heterostructures is displayed in Fig. 1a. The bright triangles are the monolayer WS2 single crystals underneath a monolayer of MoS2 that covers the entire image. The samples were epitaxially grown via a two-step chemical vapour deposition method. The well-defined WS2 triangular domains with ~100 µm lateral size were firstly grown on SiO2/Si followed by a second time growth of overlaid MoS2 layers. The top MoS2 layers only have 0° and 60° relative orientations to the bottom WS2 layer corresponding to a AA and AB stacking, respectively. By controlling growth conditions, the top MoS2 domains coalesced into a continuous monolayer film as illustrated in Fig. 1a. Then, the as-grown WS2 single crystals and MoS2/WS2 VDW heterostructures were transferred onto CaF2 substrates for spectroscopic investigations. Raman spectroscopy and atomic force microscopy measurements (Supplementary Figs 1–3) confirm that each component of the hybrid is a monolayer.

The band alignment of the MoS2/WS2 heterostructure is illustrated in Fig. 1b. The MoS2/WS2 heterostructure forms a type II heterojunction with the highest VBM and the lowest CBM residing in WS2 and MoS2 layers, respectively. The offset between two CBMs (VBM) is 0.27 eV (0.35 eV). The optical excitation

Figure 1 | Properties of the MoS2/WS2 heterostructure. (a) Optical microscope image of an as-grown WS2/MoS2 heterostructure sample. Bright triangle areas correspond to WS2 single crystals covered with the MoS2 monolayer in the entire image. (b) Illustration of the band alignment of a MoS2/WS2 heterostructure. C, CBM; V, VBM. Optical excitation of the MoS2/WS2 heterostructure leads to tightly bound interlayer excitons, with electrons (e−) in MoS2 layer and holes (h+) in WS2 layer. (c) Photoluminescence (PL) spectra of as-grown WS2 monolayer, MoS2 monolayer and MoS2/WS2 heterostructure. The strong PL at 1.96 eV of the WS2 monolayer and at 1.86 eV of the MoS2 monolayer arise from their A-exciton resonances. Photoluminescences in the MoS2/WS2 heterostructure are quenched, compared with those in the standalone samples, indicating interlayer charge transfers in the heterostructure.
of the MoS\textsubscript{2}/WS\textsubscript{2} heterostructure can lead to a tightly bound interlayer exciton with the electron (e\textsuperscript{−}) in MoS\textsubscript{2} and the hole (h\textsuperscript{+}) in WS\textsubscript{2} (refs 19,25). The efficient charge transfer in the MoS\textsubscript{2}/WS\textsubscript{2} heterostructure is confirmed by the photoluminescence (PL) measurements. Figure 1c displays the PL spectra of individual monolayer WS\textsubscript{2}, MoS\textsubscript{2} and as-grown MoS\textsubscript{2}/WS\textsubscript{2} heterostructure. The sharp peak at 1.96 eV of WS\textsubscript{2} (blue) and the peak at 1.86 eV of MoS\textsubscript{2} (red) arise from their A-exciton resonances. In the MoS\textsubscript{2}/WS\textsubscript{2} heterojunction, the peak of MoS\textsubscript{2} slightly red shifts, and that of WS\textsubscript{2} slightly blue shifts, compared with the corresponding peaks of individual monolayers. The energy shifts are caused by the interlayer coupling. The coupling strength is calculated to be \(0.05\) eV. Calculation details are shown in Supplementary Note 1.

The PL intensity of WS\textsubscript{2} in the heterostructure is significantly reduced by \(\sim 95\%\) compared with that of the monolayer, indicating a very effective charge transfer in the heterojunction since a pair of spatially separated electron and hole in two TMDC layers cannot emit efficiently\(^\text{19,27}\).

Hot excitons form rapidly after the charge transfer. The real time charge transfer dynamics across the heterojunction were directly monitored with the ultrafast visible/IR microspectroscopy (Fig. 2a). In these experiments, a femtosecond energy-tunable visible pulse excites electrons in the sample from the valence band to the conduction band. The evolution of the photo-excited charge carriers is then monitored by recording the absorption change in the mid-IR region with a femtosecond ultra-broadband mid-IR pulse. In the studies, three sets of experiments were carried out, with the energy values of the visible excitation pulses centred at the A-exciton energy of MoS\textsubscript{2} (1.85 eV), at the A-exciton energy of WS\textsubscript{2} (1.95 eV) and at the free carrier energy (3.1 eV), respectively. Different carriers and charge transport dynamical processes were therefore generated in the heterostructure and monitored by the IR detection pulses, providing a comprehensive picture for the across-layer charge transfer dynamics.

Figure 2b displays an image of transient absorption change for a MoS\textsubscript{2}/WS\textsubscript{2} heterostructure sample. The top layer represents the transient absorption change 100 fs after the excitation of MoS\textsubscript{2} with a pulse centred at 1.85 eV. The intensity of signal from the MoS\textsubscript{2}/WS\textsubscript{2} heterojunction region is about twice of that from the MoS\textsubscript{2}-only region. The bottom layer of the figure is the optical image of the measured area, where the heterojunction is located at the centre.
Figure 2c (raw data) and Fig. 2d (normalized data) display the evolutions of the photo-excitation-induced absorption changes in MoS2 monolayer (red), WS2 monolayer (blue) and the MoS2/WS2 heterostructure (green) by exciting the samples with a pulse centred at 670 nm (1.85 eV) and detecting at \( \lambda \approx 2,000 \text{ cm}^{-1} \) (\( \sim 0.25 \text{ eV} \)). Signals in all samples quickly reach maximum within fs time scales, and then decay in ps time scales. In the WS2 monolayer, the rising time is 120 fs, while in MoS2 and the heterostructure it is 50 fs. A marked enhancement of the signal maximum is observed (nearly 100% larger than the sum of the two standalone monolayers) in the heterostructure (Fig. 2c,e).

In the experiments, the optical excitation energy is centred at the A-exciton energy (1.85 eV) of MoS2 with a width \( \sim 0.1 \text{ eV} \), lower than that of WS2 (1.95 eV) (refs 19,25). In MoS2, therefore, electrons are excited to both excitonic band and conduction band because the sharp excitonic band absorption overlaps with the broad conduction band absorption33, forming both tightly bound excitons and other less tightly bound electron/hole pairs34. More detailed discussions about the carrier excitations are provided in Supplementary Fig. 6. The detection energy of \( \sim 0.25 \text{ eV} \) is well below the excitonic-binding energy (\( \sim 0.17 \text{ eV} \)).

Because of this, the excitons similar to a charge-neutrally insulating gas34,35 can hardly absorb the detection IR photons. Consequently, the detection signal is predominantly contributed by the absorption of the weakly bound electron/hole pairs. The recombinations of these electron/hole pairs and their reorganizations to form excitons result in the signal decay. In WS2, on the other hand, electrons are excited to the tail of the conduction band (absorption spectrum is in literature33 and Supplementary Fig. 6), predominantly forming weakly bound electron/hole pairs, and very few A-excitons are generated. In the heterostructure, both MoS2 and WS2 are excited, and the signal is mainly contributed from the MoS2 excitation (Fig. 2c).

At the interface, charges of both intralayer excitons and weakly bound electron/hole pairs can transfer across the boundary26,27. Electrons move to MoS2 and holes to WS2. The crossing-layer charge transfer is supported by the fast decay dynamics (\( \sim 0.8 \text{ ps} \)) of the heterostructure that is essentially the same as that of MoS2, but apparently faster than that (\( \sim 2.0 \text{ ps} \)) of WS2 (Fig. 2d, and also Fig. 3b,e), independent of the relative intensities of the two monolayers. (If charges did not transfer between two layers, the heterostructure decay dynamics would be the weighted average of the two monolayers). Absorption spectrum of MoS2 monolayers33 reveals that roughly half of the absorbed photons are converted to the energy of the intralayer excitons directly, with the rest going to the weakly bound electron/hole pairs. Detailed discussions of the carrier ratio estimation are in Supplementary Fig. 6. Before charge transfers, the intralayer excitons do not contribute to the detection signals because of their high-binding energy. After the transfer, the intralayer excitons become interlayer charge complexes. The binding energy of the charge complexes is smaller than the detection IR energy, and they absorb the detection IR photons and enhance the signals, as observed in the experiments. Since the additional IR absorbers generated by the charge transfer of excitons are as many as those generated by the charge transfer of other carriers, the signal is almost doubled. This transfer mechanism is further supported by experiments displayed in Fig. 3d,e, where only free carriers are generated by photo excitation (3.10 eV) and the interlayer charge transfers only change the signal decay dynamics, but not the signal intensity. Detailed discussions will be provided later. Note that this signal enhancement observed might also be caused by the absorption change at the interface, as well as the interlayer coupling. However, the absorption of a heterostructure prepared by vertical heteropitaxial growth is smaller than the sum of the two standalone monolayers due to interlayer interactions36,37. Therefore, the absorption change can be excluded as the reason. The increase of signal maximum cannot be caused by the interlayer coupling either, since no signal increase is observed at 3.10 eV excitation (Fig. 3d,f), which should be also affected by interlayer coupling. Multiple reflections between the two monolayers cannot cause the large intensity change either, because of the very small refractive index difference between MoS2 and WS2 that are shown in Supplementary Fig. 4.

The interlayer charge transfer rate, derived from the signal rising edge of the heterostructures (Fig. 2c,d) is faster than 50 fs. The derivation is shown in Supplementary Note 2. Following the charge transfers, the signal decays very fast. The majority of the signal decays within 800 \( \pm \) 100 fs, and a small portion decays slower than \( 13 \pm 2 \text{ ps} \). The decays are significantly faster than those of the interlayer excitons detected with visible lights26,27 and near IR (tens to hundreds of ps; Fig. 3h). The fast decay dynamics, therefore, indicate that the charge transfers do not generate the tightly bound interlayer excitons directly. Instead, the interlayer charge separation immediately produces intermediate hot excitons with binding energy lower than the mid-IR detection energy \( \sim 0.25 \text{ eV} \). The electrons of WS2 transfer to the conduction bands above CBM of MoS2 and the holes of MoS2 move to the valence bands below VBM of WS2. The majority of the hot excitons then dissipates the excess energy to the environment and forms the tightly bound interlayer excitons within 800 fs, and a small portion of the hot excitons recombines with 13 ps because of defects or phonon scattering.

The signal enhancement within 50 fs of the heterostructure (Figs 2e and 3c) and the decay dynamics of the heterostructure almost as fast as that of MoS2 (Figs 2d and 3b,e) indicate that the formation of interlayer hot excitons (faster than 50 fs) is significantly faster than the formation of intralayer excitons from free carriers or weakly bound charge complexes (600 fs–2.0 ps, Figs 2d and 3b,e).

The charges of excitons in WS2 transferring to MoS2 lead to a similar phenomena. In the second set of the experiments, the optical excitation energy was centred at 630 nm (1.95 eV). In WS2, both the A-excitons (1.95 eV) and weakly bound electron/hole pairs are generated. In MoS2, mainly the weakly bound electron/hole pairs are excited, and very few excitons (1.85 eV) are directly generated upon the optical excitation due to their relatively low energy. Similar to the results of experiments with 1.85 eV excitation, the maximum signal intensity of the heterostructure is apparently larger than the signal sum of the two standalone monolayers as displayed in Fig. 3a,c. As discussed above, the signal enhancement is also caused by the formation of interlayer hot excitons intermediates after the electron transfer of the A-excitons of WS2 to MoS2. The signal dynamics of the heterostructure are identical to those excited by the 1.85 eV photons, despite the relative signal intensity in each monolayer is different under the two excitations.

**Formation of intralayer excitons is relatively slow.** When free carriers are created and move across the heterojunction afterwards, no signal enhancement is observed. In the third set of experiments, the excitation pulse is tuned to 400 nm (3.10 eV). The photon energy is much higher than the bandgaps of MoS2 (2.39 eV) and WS2 (2.31 eV)19,25. Therefore, free carriers instead of excitons are generated88. In the standalone monolayer samples, the free carriers relax to form excitons, contributing to the fast signal decays16 (600 \( \pm \) 100 fs in MoS2 monolayer and 1.3 \( \pm \) 0.2 ps in WS2 monolayer) observed in Fig. 3d,e. The slow portion of the signal decays (\( \sim 10 \text{ ps} \)) is attributed to the recombination of the free carriers through other ways, for example, the carrier-phonon...
scattering or defect assisted electron–hole recombination\textsuperscript{34}. In the heterostructure, both MoS\textsubscript{2} and WS\textsubscript{2} are excited by the 3.10 eV light. More carriers are generated in the WS\textsubscript{2} layer than in MoS\textsubscript{2} (Fig. 3d). Similar to the observations with excitations at lower energy, the free carriers rapidly transfer across the interlayer boundary, resulting in a much faster signal decay of the heterostructure compared with that of WS\textsubscript{2}. The results clearly demonstrate that the interlayer charge transfers occur before the formation of intralayer excitons. After crossing the interlayer, the free carriers still remain free and able to absorb the mid-IR photons. The population of photo-induced free carriers is unchanged, and the absorption coefficients of carriers are almost unchanged because of the similar environment in MoS\textsubscript{2} and WS\textsubscript{2} monolayers, leading to that the interlayer charge transfers following the 3.10 eV excitation do not enhance the signal intensity (Fig. 3f). This is different from the observations with the excitation energy on resonance with the A-excitons (Figs 2e and 3c), where interlayer charge transfers of excitons produce extra mid-IR absorbers, as discussed above.

The observed fast signal decay in the MoS\textsubscript{2}/WS\textsubscript{2} heterostructure is repeatable in different areas of the sample with different incident excitation fluences (from 5 to 20 \(\mu J/cm^2\)). As the fluence increases, the excitation-induced absorption change increases with the same proportion and has not reached the nonlinear regime until the incident fluence reaches 20 \(\mu J/cm^2\) (Supplementary Fig. 5). Under the excitation conditions, it is possible that biexcitons or other multi-particle species can be generated\textsuperscript{39}. These multi-particle species can evolve into exciton pairs with excess energy in a fission-type process. However, the binding energy (~50 meV)\textsuperscript{39} of these species is lower than the mid-IR-probe photon energy, and they contribute to the experimental signals in a way similar to free carriers that is not
affected by interlayer charge transfer. Therefore, the generations of these multi-particle species do not affect the conclusion that the charge transfer of intralayer excitons results in interlayer hot excitons. More discussions about possible high-phothoeitation-density effects on the dynamics are provided in Supplementary Fig. 5. Figure 3g also suggests that the photo-induced dynamics in the heterostructure remain constant in the detection energy range from 1,390 (0.17 eV) to 2,910 cm\(^{-1}\) (0.36 eV), indicating that the binding energy of the interlayer hot exciton intermediate is <0.17 eV and that of the interlayer excitons is >0.36 eV.

In the above experiments, signals following the interlayer charge transfers decay significantly faster than that of the interlayer excitons observed with visible\(^{19}\) and near-IR detections (1.55 eV; Fig. 3h). As discussed above, experiments in this work detect free carriers and intermediate hot excitons with binding energy lower than the detection mid-IR photon energy. Charge complexes, for example, A-excitons, with higher binding energy do not contribute to the experimental signals. The binding energy of interlayer exciton in the MoS\(_2\)/WS\(_2\) heterostructure has not been previously experimentally determined. Our experiments show that the 1.55 eV near-IR photon can excite the interlayer excitons (Fig. 3h), but the 0.36 eV mid-IR photon cannot (Fig. 3g), indicating that the binding energy of the interlayer excitons resides between 0.36 and 1.55 eV. Our calculations based on a model\(^{25}\) also suggest that the interlayer binding energy is 0.4–0.6 eV (Supplementary Fig. 7; Supplementary Note 3), slightly lower than that (0.5–0.7 eV) of a single layer\(^{24–26}\).

Figure 4 | Charge separation processes in the heterostructure. After the excitation of WS\(_2\) (MoS\(_2\)) monolayer, the electron (hole) transfers from the CBM (VBM) of WS\(_2\) (MoS\(_2\)) monolayer to the conduction (valence) bands just above (below) CBM (VBM) of the MoS\(_2\) (WS\(_2\)) monolayer within 50 fs, generating a hot interlayer exciton with a relatively long distance. The hot exciton then reorganizes and dissipates the excess energy to form the tightly bound interlayer exciton within 800 fs.

The charge transfers include two major steps. Our measurements herein suggest that the interlayer charge transfer process in the MoS\(_2\)/WS\(_2\) heterostructure includes two major steps: the immediate formation of interlayer hot excitons following the ~50 fs interlayer charge separation, and then the relaxation of hot excitons to generate tightly bound excitons within ~800 fs. The process is schematically illustrated in Figs 4 and 5. The experimental observations support one of the two theories (the hot exciton theory) recently proposed\(^{28}\) to explain why the charge transfer dynamics in 2D heterostructures are little dependent on the momentum mismatch of intra- and interlayer excitons: the momentum conservation during the charge transfer can be achieved by the formation of a hot interlayer exciton that employs the excess energy through the transfer. Specifically, after photo excitation, charge transfers don’t form interlayer excitons directly. Instead, the electron (hole) transfers from WS\(_2\) (MoS\(_2\)) to the conduction (valence) bands above (below) CBM (VBM) of the MoS\(_2\) (WS\(_2\)) generating a hot interlayer exciton. The hot exciton has a relatively long distance between the electron and hole, and therefore is only weakly bound. The hot exciton then reorganizes and dissipates the excess energy to form the tightly bound interlayer exciton.

Discussion
The experiments herein demonstrate that the ultrafast charge transfer between the WS\(_2\) and MoS\(_2\) monolayers results in the formation of interlayer hot excitons. The existence of hot excitons explain why the interlayer charge transfer is independent on the apparent momentum mismatch between the intralayer/interlayer excitons, and also answers why an electron–hole pair can overcome the large Coulomb potential to generate photocurrent in ultrathin photovoltaic cells with MoS\(_2\)/WSe\(_2\) or MoS\(_2\)/WS\(_2\) as the active p–n junction and conventional metal or graphene as
interlayer excitons. That carriers other than excitons created before they turn into the tightly bound exciton can help the electron/hole pair easily dissociate in the heterostructure move across the interface with a much higher rate than recombining into the intralayer excitons also favours the generation of photocurrent. The semi-free character of the hot interlayer excitons can greatly facilitate the charge separation and photocurrent generation processes, which is critical for many applications of 2D heterostructures in optoelectronics and photovoltaics.

Methods

MoS2/WS2 heterostructures growth. Our heterostructure growth follows a two-step process: the WS2 domains were firstly grown on SiO2 substrates followed by second growth of MoS2 on the top of WS2. SiO2 substrate was pre-cleaned with acetone and isopropanol, followed by treated in piranha solution (H2SO4:H2O2—3:1) for 2 h. The triangular WS2 domains were firstly grown on SiO2 (300 nm)/p + + + Si substrates. Growth process was done in three-zone chemical vapour deposition system with 1 inch quartz tube using WO3 (Alfa Aesar 99.999%), MoO3 (Alfa Aesar 99.999%) and S (Alfa Aesar 99.9%) powders as the precursor. For MoS2 growth, typical temperatures for three zones are 115°C, 560°C and 800°C, respectively. All of three temperature zones were heated to preset values at a rate of 25 °C min−1. Sulfur and MoO3 powders were separately loaded in two 10 mm-radius mini- quartz tubes, which enable a stable evaporation of S and MoO3 sources by preventing any cross-contaminations during the growth. All temperature zones were kept stable for 20 min before growth. The precursor powders were pre-placed outside of the furnace and rapidly loaded from outside into each zone to start the growth. During the growth process, argon was used as a carrying gas at a flow rate of 130 sccm, and the vacuum pressure was kept at 0.7 torr.

Heterostructures transfer. The as-grown MoS2/WS2 VDW heterostructures are transferred onto CaF2 substrate for transmission mode laser experiments. The process is as follows: at first, a thin layer of poly(methyl methacrylate) (PMMA; PMMA/CH176/C176B4) is spin-coated onto the sample on SiO2 substrate twice at the speed of 4000 rpm, then the sample is carefully placed in the buffer HF (1:5) and let it float. After 20 h, the PMMA sheets is separated from the SiO2 substrate and dried with glass plate, move to deionized-water for rinsing a few times, then pre-cleaned CaF2 windows are used to fish the PMMA sheets. The CaF2 windows with PMMA and samples stay in room temperature and vacuum for 24 h, finally acetone is used to remove the PMMA, the MoS2/WS2 VDW heterostructures stayed on the CaF2 substrate and confirm by optical microscope.

Raman and PL measurements. Raman and PL spectroscopy were carried out using a Horiba Jobin Yvon LabRAM HR-Evolution Raman microscope. The excitation light is a 532 nm laser, with an estimated laser spot size of 1 μm. Raman and PL spectroscopy were carried out using a Horiba Jobin Yvon LabRAM HR-Evolution Raman microscope. The excitation light is a 532 nm laser, with an estimated laser spot size of 1 μm.

Ultrafast visible/IR microspectroscopy. The experimental setup of the ultrafast visible/IR microspectroscopy is illustrated in Fig. 2a and Supplementary Fig. 4. In brief, the output of a femtosecond amplifier laser system (at a repetition rate of 1 kHz, 1.6 mJ energy per pulse, 800 nm central wavelength and a pulse duration of ~50 fs, Uptek Solutions Inc.) was split into two parts. One was used to pump a home-built nonlinear optical parametric amplifier to generate visible laser pulses with tunable wavelengths, and the other was directed to generate an ultra-broadband super-continuum pulse that covers almost the whole mid-IR region. In ultrafast experiments, the visible pulse is the pump light with the central wavelength and excitation power adjusted based on need. The interaction spot on samples varies from 120 to 250 μm. The mid-IR super-continuum pulse acts as the probe light, which was focused on the sample by a reflective objective lens (× 150/0.28 numerical aperture, Edmund Optics Inc.) to reduce the spot size to the level of sample area (~40 μm). A 300-megapixel digital camera was used to align the pump/probe beam to proper sample area. The probe light was detected by a liquid-nitrogen-cooled mercury–cadmium–telluride array detector after frequency resolved by a spectograph with a resolution of 1–3 cm−1 that is dependent on the central frequency. The time delay between the pump light and probe light was controlled by a motorized delay stage.

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

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**Author contributions**

J.Z. and H.C. designed the experiments. J.Z. supervised the project. H.C. and X.W. performed the ultrafast experiments. X.W., J.Z., Y.G., X.Z., J.Y., C.Y., J.L., P.A. and G.Z. prepared materials. T.W. and W.Z. performed the binding energy calculations. J.Z., H.C., W.Z. and G.Z., prepared and revised the manuscript.

**Additional information**

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