Coupled valence carrier and core-exciton dynamics in WS$_2$ probed by few-femtosecond extreme ultraviolet transient absorption spectroscopy


I. INTRODUCTION

Studying the dynamics of elementary excitations in semiconductors such as photoexcited carriers, phonons, and excitons has been crucial to the success of electronic devices [1,2]. While most measurements on the photophysical and photochemical properties of semiconductors are performed within the optical domain, few-femtosecond to attosecond core-level transient absorption (TA) and transient reflectivity spectroscopy have recently been utilized to investigate carrier dynamics in semiconductors and two-dimensional materials [3–10]. Core-level TA spectroscopy in semiconductors typically consists of an optical pulse to excite the carriers in the sample and an extreme ultraviolet (XUV) or x-ray pulse to record the changes in the core-level absorption spectra. In many semiconductors, the core-level absorption spectra can be mapped onto the conduction band (CB) density of states (DOS) due to significant dielectric screening [14], and the core-level TA spectra directly reflect the carrier distributions as a function of energy, thereby providing real-time tracking of carrier dynamics [4–7].

In contrast, the core-level absorption spectra of many insulators, in particular, ionic solids with poor dielectric screening, exhibit sharp peaks below the onset of core-to-CB edges [15]. These discrete transitions are termed “core excitons,” which are formed by the Coulomb attraction between the excited electron and the core hole [16,17]. The electron-hole binding results in longer lifetimes of the core excitons compared with the typical <1 fs decay time of core-to-CB transitions [18,19]. The observation of the decay of core excitons is enabled through attosecond transient absorption spectroscopy in the extreme ultraviolet, a core-level TA spectroscopy utilizing subfemtosecond XUV pulses in combination with <5-fs-long optical pulses. As opposed to the typical XUV TA measurement in semiconductors where the XUV pulse probes the valence electronic state after optical excitation [3,4,7,8,20], those experiments probing core-exciton states use the subfemtosecond XUV pulse to excite the core excitons, and...
the core-exciton transition dipoles are subsequently perturbed with the optical pulse [11–13], analogous to the studies on the decay of atomic Rydberg states and autoionization states [21]. Due to the large band gap in insulators, which exceeds the photon energy of available visible and ultraviolet light pulses, the observation of the effect of carrier dynamics on a core-excitonic system via core-level TA spectroscopy has not been achieved, and the effect of valence electron-hole pairs on core-exciton transitions and their dynamics remains elusive.

In this paper, we report the observation of core-exciton transitions within the W N 6 edge (32–37 eV) of WS2 and a smooth core-level absorption edge at the nearby W O 3 transition (37–45 eV), which consists of core-to-CB transitions that can be understood within the single-particle mean-field picture. The proximity of the two different types of core-level absorption edges presents an excellent opportunity in simultaneously observing the dynamics of carriers in the valence shell and their influence on the dynamics of core excitons. A single experiment thus probes the carriers at the W O 3 edge and the discrete core-exciton transitions at the W N 6,7 edge. Tungsten disulfide is a group VI transition metal dichalcogenide and a semiconducting two-dimensional (2D) layered material. In its mono- and bilayer form, the electronic structure and photophysics of WS2 have been extensively studied for potential applications in optoelectronics and 2D valleytronics [22–34]. Recently, interlayer charge-transfer excitations and novel elementary excitations such as moiré excitons were observed in heterostructures containing WS2 layers [35–38].

Here, by conducting core-level transient absorption spectroscopy in the XUV on WS2 thin films, picosecond hole relaxation and carrier recombination times are obtained from the core-level TA spectra at the W O 3 edge. An ~10 fs coherence lifetime of core excitons at the W N 6,7 edge is also measured. In contrast to the attosecond XUV TA studies on insulators, where the observed dynamics are dominated by coupling of core-exciton states with the optical field [11–13], here the core-exciton lineshape is primarily influenced by the change in electronic screening and band filling due to carriers excited by the optical pulse. This optical-XUV transient absorption study at the W O 3 and N 6,7 edges provides a prototypical example for measuring the carrier-induced modification of core excitons and paves the way for exploring carrier dynamics in 2D heterostructures and superlattices involving transition metal dichalcogenides, where the element specificity of core-level TA spectroscopy can be employed to enable layer-selective probing of photophysical and photochemical phenomena.

II. EXPERIMENTAL SCHEME

The details of sample preparation and the scheme of the XUV TA spectroscopy experiment are provided in Appendices A and B, respectively. In brief, 40-nm-thick WS2 films were synthesized on 30-nm-thick silicon nitride windows by atomic layer deposition (ALD) of WO3 thin films and subsequent sulfurization with H2S [39]. In the XUV TA experiments, the samples were irradiated with a broadband optical pulse (500–1000 nm) with nominal duration of 4 fs and a time-delayed broadband XUV pulse (30–50 eV) produced by high-harmonic generation using a near-single-cycle optical pulse in a Kr gas jet.

III. RESULTS AND DISCUSSION

The core-level absorption spectrum of the 40-nm-thick WS2 film is displayed in Fig. 1(a) (red line). The static spectrum below 37 eV (marked with the dashed cyan line) exhibits four distinct peaks labeled as A–D. Peaks A and B occur on top of the absorption edges between 33 and 34 eV. Peak C exhibits a Fano-type asymmetric lineshape at approximately 35.5 eV with fine-structure peak D occurring at approximately 36.6 eV [40]. A smooth absorption feature extends from 38.5 eV to beyond 45 eV. By comparing the measured spectrum with the calculated imaginary part of the dielectric function using the all-electron full-potential linearized augmented plane wave (FP-LAPW) method (Appendix C) with random phase approximation [41,42], the smooth absorption feature above 38.5 eV is assigned to the transition between the W 5p3/2 core bands and the CB (W O 3 edge). Peaks A and B are assigned to the W 4f7/2 transitions (W N 7 edge), and peaks C and D are assigned to the transitions from W 4f5/2 core levels to the CB (W N 6 edge). The comparison between the measured W 5p3/2 absorption edge and the calculated dielectric function [Fig. 1(a), dashed black line] indicates significant lifetime broadening of the W 5p3/2→CB transitions. In addition, the peaks measured at the absorption edges between 32 and 37 eV are clearly different from the smooth onset of the 4f→CB transitions [Fig. 1(a), dash-dotted black line] calculated with mean-field approximation, suggesting that many-body interactions between the electron and the core hole contribute to the measured discrete lineshape. Note that the broadband XUV pulse (30–50 eV) covers the core-level transitions from both the W 4f and 5p orbitals, enabling simultaneous observation of dynamics at the two different edges.

Typical core-level TA spectra between ~40 fs and 2.6 ps time delay are displayed in Fig. 1(b). In this paper, the time delay is defined as the arrival time of the XUV pulse subtracted from the time of the optical pulse, and positive time delay indicates that the samples are probed by the XUV pulse “after” optical excitation. The change in absorbance ∆A(t) at a specific time delay t is defined as ∆A(t) = A(t) − A(t = −40 fs) with A denoting absorbance. The pump-excited carrier density is estimated to be 1×1021 cm−3 or 6×1013 cm−2 per layer (Appendix D). Between 37 and 40 eV below the W O 3 edge, two weak, broad positive features are observed [Fig. 1(b)]. Although the difference in static absorbance below and above the W N 6,7 edge, or “edge jump,” is much smaller than the W O 3 edge [Fig. 1(a)], the XUV TA signal occurring near transitions A, B, and C is narrow and much stronger than the TA signal above 37 eV [Fig. 1(b)]. Clearly, the nature of the W N 6,7 edge transitions (peaks A–D) and the W O 3 edge transitions are different, and separate treatment is needed to understand their corresponding XUV TA spectra.

When a core electron is excited into the CB, the excited electron can interact with the core hole via Coulomb attraction. In many semiconductors, the electronic screening reduces the Coulomb interaction such that the core-level transitions can still be understood in a single-particle picture [14].
As the core bands are dispersionless, the core-level transitions map the CB density of states and, with optically excited carriers in the valence band (VB) and CB, core-level transitions probe the electronic occupation in the valence shell and the energy shifts of the VB and CB due to carrier and phonon excitations [Fig. 1(c)] [4–7]. This scheme corresponds to the smooth W\ O\ _3 edge transitions above 37 eV but cannot describe the transitions at the W\ N_{6,7} edge.

Discrete peaks form in core-level absorption spectra when the Coulomb attraction between the electron and the core hole is non-negligible. The interaction between the core hole and the electron excited by the XUV or the x-ray renormalizes the core-level absorption, and discrete “core-exciton” peaks can form near the critical points of the core-to-CB transitions [Fig. 1(d)] [16]. Note that excitonic interactions between the excited electron and the core hole are present for all core-to-CB absorptions. Therefore the renormalization of the core-level absorption spectra is not limited to the near-edge transitions, but may extend several eV above the edge [43,44]. The behavior of transitions at the W\ N_{6,7} edge below 37 eV is consistent with the description of core excitons. To verify this, we compare the XUV TA spectra of the W\ N_{6,7} and W\ O_3 edges near zero time delay.

Figure 2(a) displays the XUV TA spectra between −25 and +25 fs time delay; lineouts of the XUV TA spectra at five different time delays between −20 and +7 fs are plotted in Fig. 2(b). At positive delays, the XUV light probes the changes due to photoexcitations in the valence shell, while at negative time delays, the optical pulse perturbs the core-level transition dipole before its decay by both carrier photoexcitation and the direct coupling of the core-excitonic transitions with the optical field [11–13]. The distinction between the transitions at the W\ N_{6,7} edge and the W\ O_3 edge can be visualized in the XUV TA signal at negative delays. While the TA signal at the W\ O_3 edge (37–40 eV) diminishes to zero at less than −4 fs time delays, TA signals near peaks A, B, and C are still visible at less than or equal to −10 fs time delays. The experimental results thus indicate that transitions A, B, and C are more long-lived than the transitions below the W\ O_3 edge.

The comparison of core-level transition lifetimes corroborates the assignment that the peaks within the W\ N_{6,7} edge [Fig. 1(a)] are core excitons because the electron–core-hole attraction of the core exciton stabilizes the core-excited state and enables a longer lifetime [18]. The broad positive feature above 37 eV at positive time delays can then be interpreted as photoexcited holes in the VB. In the following, we first focus on the measured dynamics induced by photoexcited carriers at the W\ O_3 edge. Next, we discuss the measured TA spectra of core-exciton transitions at the W\ N_{6,7} edge at both positive and negative time delays.
FIG. 2. (a) XUV TA signal between −25 and +25 fs time delay. The static spectrum is plotted as a dashed black line as a reference. (b) shows the XUV TA lineouts at five different time delays between −20 and 10 fs.

A. Carrier dynamics at the W O₃ edge

To understand the XUV TA signal at the W O₃ edge, XUV TA spectra at five different time delays between +7 fs and +2 ps are presented in Fig. 3(a). At energies above the edge (41–45 eV), the XUV TA spectra exhibit a weak decrease in absorbance (negative ΔA) throughout the entire range of delays, whereas two positive features with different dynamical behavior are observed between 37 and 40 eV below the edge. The feature spanning 37–38.4 eV decays with time, while the feature between 38.4 and 39.5 eV, which is barely observable near time zero, increases in magnitude with the time delay.

The positive and negative ΔA below and above the edge might initially suggest that the positive feature is due to holes in the VB, which open up new excitation pathways from the core, and the negative feature is due to electrons in the CB, which blocks the core-level excitations into the CB. The different dynamical behaviors between 37–38.4 eV and 38.4–39.5 eV could then be assigned to relaxation of the photoexcited hot hole to the VB edge. However, such an assignment implies that the transition from the core to the VB edge is at approximately 39.5 eV and the initially photoexcited holes are located approximately 2 eV below the VB maximum. Given the approximately 2 eV direct band gap of bulk WS₂ [22], this suggests an optical transition energy of approximately 4 eV, far exceeding the maximum photon energy at 2.5 eV of the optical pulse. Therefore the positive feature between 38.4 and 39.5 eV cannot be assigned to holes in the VB. Instead, it can be interpreted as a redshift of the W O₃ edge due to band gap renormalization. Both the change in electronic screening and phonon heating due to photoexcited electrons and holes can lead to band gap renormalization, resulting in the lowering of the CB edge and therefore a redshift of the core-to-CB transition energies [4]. As energy dissipates from the electronic domain, the increase in phonon temperature and the heat-induced lattice expansion can further enhance band gap renormalization and lowering of the CB [45], which causes the positive features to grow with time delay. In addition, with the assignment of the positive feature spanning 38.4–39.5 eV as due to the phonon-induced edge shift, the feature between 37 and 38.4 eV, which diminishes with increasing time delay, can then be assigned to holes.

To track the hole relaxation process, the median energy of the hole signal $E_{h,med} = \frac{\int E \Delta A dE}{\int \Delta A dE}$ is plotted in Fig. 3(b). The median energy $E_{h,med}(t)$ shifts from 37.7 to 37.9 eV with respect to time and can be fitted by a single exponential with a time constant of $1.2 \pm 0.3$ ps. In addition, as holes relax to the VB edge at the long-time limit, the core-to-VB edge transition energy can be determined from $E_{h,med}(t \to \infty)$. The extracted transition energy from the core to the VB edge from the exponential fitting is 37.9 eV. The proximity between the hole feature and the positive feature due to band gap renormalization leads us to assign the core-to-VB edge transition with the median energy of the hole feature at the long-time limit rather than use the maximum energy cutoff of the hole feature as in the core-level TA studies of germanium and 2H-MoTe₂ [4,7].

Consulting the band structure diagram of WS₂ [Fig. 4(b)] and the initial photoexcited carrier distribution that is
of states (PDOS) of WS$_2$ calculated with the FP-LAPW method as a function of crystal momentum [Fig. 4(a)], the initial proportional to the number of photons available for excitation

\[ \text{int} \mathcal{N}_{\text{ph,exc}} = \sum_{\text{states}} |\langle \psi_{\text{exc}} | \phi_n \rangle|^2 \]

where $\langle \psi_{\text{exc}} | \phi_n \rangle$ is the sum of the modulus square of the wave-function projections on atom $n$.

The assignment of the VB maximum at 37.9 eV suggests that a negative TA feature due to excited electrons is expected at 39.9 eV (37.9 + 2 eV band gap). While a weak negative TA signal barely above the noise level is indeed observed spanning 41–45 eV, assigning this to the CB electrons plus the VB maximum at 37.9 eV would suggest an electron-hole energy separation of at least 3 eV, which exceeds the maximum photon energy of 2.5 eV in the optical pulse. In addition, no significant dynamics are observed for the negative feature between 41 and 45 eV, whereas the electrons are expected to recombine with the holes within the observed 3.1 ps recombination time. This indicates that the weak signal at $> 41$ eV may be caused by changes in core-hole lifetime, which affects the spectral broadening of the edge, or a decrease in oscillator strength of the W O$_3$ edge due to carrier and phonon excitations [7,46], rather than directly due to the photoexcited electrons in the CB.

The absence of the XUV TA signal due to occupation of photoexcited electrons in the CB can be explained by the significant spectral broadening above the W O$_3$ edge, where the experimentally measured static spectrum is much wider and smoother than the calculated dielectric function according to the projected CB DOS [Fig. 1(a)]. Here, the fine structure in the projected CB DOS due to critical points in the CB is completely lost in the measured absorption edge. This is in contrast to the recently studied L$_{2,3}$ edge in Si and Te N$_{4,5}$ edge in 2H-MoTe$_2$ where the critical points in the CB can be directly mapped onto the core-level absorption spectrum [7,47]. The broadening in the W O$_3$ edge increases the overlap between the expected negative XUV TA feature due to electron occupation in the CB and the positive feature due to CB redshift. The overlapping negative and positive features thus lead to the cancellation between the two and make extracting the electron distribution in the CB from core-level TA spectra here unreliable.

**B. Dynamics and lifetimes of core excitons at the W N$_{6,7}$ edge**

While the core-level TA spectra at the W O$_3$ edge can be explained by the redshift of CB and electronic occupation in the VB and CB, XUV TA signals of core-exciton transitions within the W N$_{6,7}$ edge cannot be interpreted with the same approach. Note that unlike the insulators where core-exciton dynamics have only been observed when XUV light arrives first [11–13], here the XUV TA signal at the core-exciton transitions extends throughout the entire range of positive time delays [Figs. 1(b) and 2(a)]. This difference arises because in insulators, the band gap exceeds the photon energy range of the optical pulse, so electron-hole pairs in the VB and CB are not excited. Thus the core excitons can only be perturbed by optical-field-induced coupling of core-exciton states. Here, photoexcited carriers with picosecond lifetimes can instantaneously “dress” the XUV-excited core excitons by the carrier-induced change in band filling and electronic screening [48,49]. Therefore carrier-induced modification of the core-exciton lineshape occurs at both negative and positive time delays.

The presence of the XUV TA signal at short (<25 fs) positive delays is solely due to photoexcited carriers since dynamics caused by electron-phonon scattering occur on a
timescale of $10^2$ fs and can be ignored [50–52]. While photoexcited carriers are the sole contributor to the modification of core excitons at short positive time delays, at negative delays direct field-induced changes can also modify the core-exciton lineshape. It has been shown that similar to atomic autoionizing states [21,53–56], the optical pulse can cause energy shifts of the core-exciton transitions through the AC Stark effect and resonant coupling between the core-exciton states or with the ionization continuum [11–13]. Formal treatment of the XUV TA spectra at negative time delays requires computation of the free induction decay of the core-exciton transition dipoles by solving a time-dependent Schrödinger equation, including couplings of the core-level transitions with both the photoexcited carriers and the optical field [11–13,54]. However, due to the complexity in including the many-body interactions between the core-level transitions and the photoexcited carriers, which is detailed in the next section, we propose an alternative method to separate the contribution from photoexcited carriers and the optical field by their different time behaviors.

Here, we analyze the different contributions by applying global fitting to the XUV TA spectra $\Delta A$ through singular value decomposition (SVD) (Appendix E): $\Delta A(t, E) = \sum_n s_n(t) w_n(E)$. Functions $u_n(t)$ and $v_n(E)$ are singular vectors, or components, ranked by singular values $s_n$ in descending order. Transition D is excluded from the analysis due to poor signal in that spectral region. The XUV TA signal from the largest component at transitions A and B is shown in Fig. 5(a), showing good agreement with experimental data [Fig. 2(a)] and indicating that the dynamics at transitions A and B can be described by a single component. The corresponding singular vector [Fig. 5(b)] directly reflects the TA signal measured at $+10$ fs time delay [Fig. 2(b)], and the dynamics of the component [Fig. 5(c)] exhibits an exponential decay at negative delay and becomes constant when $t > 0$. This indicates that transitions A and B have similar decay dynamics and lifetimes, and although the optical pulse can potentially affect the core excitons directly through coupling the core-exciton transition dipoles with the optical field, the carriers excited in the valence shell remain the dominant influence on core excitons A and B.

Within their decay time, core excitons A and B are modulated by the valence electron-hole pairs. By fitting the decay dynamics with a single exponential [Fig. 5(c)], a core-exciton coherence lifetime ($T_2$) of $10.9 \pm 0.4$ fs is extracted. The decoherence in core-exciton transitions can be caused by population decay through Auger processes or by exciton-phonon coupling [11–13]. Previous studies on the decay of core excitons in insulators show that when exciton-phonon coupling prevails over other decoherence channels, the free induction decay of the core-exciton transition dipole moment exhibits a Gaussian decay [11–13] that leads to a Gaussian spectral profile [57,58]. Here, the decay of XUV TA signal at negative time delays is exponential rather than Gaussian [Figs. 5(c) and 5(f)]; a comparison between Gaussian and exponential fitting is detailed in Appendix E, suggesting that phonon-induced dephasing is insignificant and Auger processes are the dominant contributor to core-exciton decay. Thus a population decay time ($T_1 \approx T_2/2$) at $5.5 \pm 0.2$ fs can be inferred [59].

To quantitatively reproduce the XUV TA spectra at transition C [Fig. 2(a)], the two largest components in the SVD are required [Fig. 5(d)]. The dynamics of the two components [Fig. 5(f)] show that at $> 10$ fs delays, the largest (first) component is constant whereas the second component is zero. In addition, the second component only becomes nonzero either at negative time delays or during pulse overlap. This indicates that the largest component represents the influence of carriers on the core exciton and the second component originates from the direct coupling of the core exciton to the optical field, because in contrast to the direct coupling to the optical field that can only occur when the field overlaps with the transition dipole before its decay, the carriers are much longer lived than the transient optical pulse and can cause spectral changes at $>10$ fs time delays. The field-induced TA component [Fig. 5(e), blue line] exhibits a negative amplitude below the edge and a positive amplitude above [cf. Fig. 6(b)], the asymmetry of the XUV TA amplitude centered around the edge suggests that it may relate to the change in the Fano $q$ factor of the transition, which can be induced by the optical pulse through a ponderomotive phase shift [54], or direct coupling to neighboring core-excited states [55]. Fitting the

FIG. 5. (a) XUV TA signal from the largest component in SVD for transitions A and B; the corresponding singular vector with respect to energy and time is shown in (b) and (c), respectively. (d) exhibits the XUV TA signal from the two highest ranked components in SVD for transition C; the corresponding singular vectors for the first and second largest singular values as a function of energy and time are shown in (e) and (f), respectively. The XUV TA signal from only the largest component in SVD for transition C is shown in Fig. 10 (Appendix E). Note that the magnitude of the singular value of each component is included in the amplitude shown in (c) and (f).
dynamics of the largest component with a single exponential yields a core-exciton coherence lifetime of 9.6 ± 0.1 fs.

C. Carrier-induced modification of core-exciton transitions within the W N₆.₇ edge

In this section, we focus on the modification of the core-exciton absorption lineshape by photoexcited carriers. First, we compare the effect of photoexcited carriers on core-exciton and core-to-band absorption at +10 fs time delay [Fig. 2(b)]. In contrast, the positive TA signal observed below the W O₃ edge, no positive signal is observed between 30 and 33 eV [Fig. 1(b)], and the majority of the TA signal occurs near core-exciton transitions A–C. The lack of XUV TA signal directly from the electronic state blocking of carriers as at the W O₃ edge can be attributed to the renormalization of core-level absorption spectra due to core-exciton formation. The formation of core excitons concentrates the oscillator strength of core-to-CB transitions to the bound core-excitonic states, and the core-level absorption spectra no longer map to the CB DOS. In addition, the oscillator strength at the W N₆.₇ edge is lower than the W O₃ edge. Compared with the edge change (jump) of 1 optical density (OD) below and above the W O₃ edge, the W N₆.₇ edge has an edge jump of ≤0.5 OD [Fig. 1(a)]. This indicates that the signal of electronic state blocking, if present at the W N₆.₇ edge, would be much lower than the signal below the W O₃ edge (37.4–38.4 eV), which is already close to the noise level.

To understand the effect of carriers on core-exciton transitions, the core-level absorption spectra at +10 fs time delay with three different optical pump fluences are displayed in Fig. 6. While transition D is broadened with increasing pump fluence, the carrier-induced changes at A, B, and C are much more complex. At transitions A, B, and C, the absorption edge shifts to higher energy with increasing pump fluence. However, the changes in absorbance below and above the edge are nonmonotonic with increasing optical excitation. At fluences between 0 and 21 mJ/cm², the absorbance below the edges shifts to higher energy with increasing pump fluence. At a fluence of 29 mJ/cm², the absorption above the edge increases rather than decreases compared with the absorbance at 21 mJ/cm² pump fluence. In addition, shoulders and ripples start appearing around transitions A, B, and C at a fluence of 29 mJ/cm². New small features appear below the edge of A and C, and a dip appears below absorption edge B. Clearly, the carrier-induced changes to the core-exciton transitions cannot be simply described by an energy shift or a broadening of the lineshape.

The behavior of the core-exciton spectral change at +10 fs delay with respect to increasing pump fluence is reminiscent of the changes in optical absorption of valence excitons in highly excited semiconductors [49]. For example, it has been observed that in highly excited GaAs, the absorption peak of bound excitons decreases in magnitude and new features below the onset of excitons appear in the optical absorption spectra [48]. Using a generalized Elliott formula [60–62], Lee et al. showed that the contributions to the optical absorption spectra of highly excited semiconductors can be divided into three categories [48]. First, the ionization of bound excitons due to the screening of the photoexcited electron-hole plasma causes suppression of the bound exciton transitions and an overall blueshift of the absorption edge. Second, the increasing band filling causes ripples to appear around the exciton transitions, especially below the absorption onset. Third, the carrier-induced band gap renormalization introduces a redshift of the CB edge that partially compensates the blueshift of absorption onset due to suppression of bound exciton transitions.

The phenomena observed for optical excitons can analogously explain the spectral changes in core-exciton transitions here. The increase in electronic screening due to electron-hole excitations in the VB and CB suppresses bound core-exciton transitions and contributes to the overall blueshift of the absorption onset. The changes in band filling in the valence shell modulate the core-excited states’ energies, oscillator
The similar behavior between the carrier-dressed core-exciton lineshape and optical absorption in highly excited semiconductors, which can be simulated analytically, leads us to explore the possibility of extending the formalism [61] to quantitatively extract parameters such as core-exciton radii and binding energies. The generalized Elliott formula [60–62] is based on a parabolic two-band model that only incorporates a single CB minimum. As core-exciton features B and D are clearly embedded in the core-to-CB continuum transitions, the parabolic two-band approximation is no longer applicable. In addition, due to the dispersionless core bands, core excitons can form at multiple CB minima, e.g., at the K, Δ, and Σ valleys (Fig. 4), and the wave functions at those CB minima can further hybridize. Therefore quantitative treatment of the core-exciton transitions here, and by extension, their modification due to carriers, will require Bethe-Salpeter equation calculations including the full band structure of WS$_2$ [63–65], which is beyond the scope of this work.

**D. Picosecond XUV transient absorption signal at the W N$_6$,$\bar{7}$ edge**

As photoexcited carriers are the major contributor to the modulations of the core-exciton spectra at negative and short positive time delays, we consider here the possibility of using the TA spectra of core excitons to extract carrier dynamics. Although picosecond carrier relaxation and recombination would suggest a decay of TA signal at the core-exciton transitions, a growth of TA signal (Fig. 7, black arrows) is observed below transitions A and C with increasing time delay, and no significant TA change is measured above the edge at transitions A, B, and C throughout 0–2.6 ps. This indicates that in addition to photoexcited carriers, the excitation of phonons through electron-phonon interactions also contribute to the spectral changes in core excitons at long time delays, as phonon-induced band gap renormalization can induce a redshift of CB that is consistent with the positive TA signal observed below the transitions. Therefore the core-exciton transitions at the W N$_{6,7}$ edge here are poorly configured for extraction of carrier dynamics, because the spectral changes due to carriers and phonons at hundreds-of-femtoseconds to picosecond timescales cannot be easily separated.

**E. Comparison between the core-exciton and core-to-conduction-band transitions**

The contrasting behavior of core-exciton transitions at the W N$_{6,7}$ edge and the core-to-band transitions at the W O$_3$ edge in the same energy range is highly unique and suggests that factors other than the macroscopic screening, which are experienced by transitions from both W 4$f$ and 5$p$ through XUV photons, are contributing to the core-exciton formation. As the CB minima are dominated by W 5$d$ orbitals that are accessible from both W 4$f$ and 5$p$ through XUV photons, it is thus suggested that the contributor to the difference between the W N$_{6,7}$ edge and the W O$_3$ edge absorption lies in the properties of the core orbitals. The W 4$f$ orbitals involved at the W N$_{6,7}$ edge are far more localized than the W 5$p$ orbitals for the O$_3$ edge transitions. The localized core hole may then act as a point positive charge and modulate the electronic wave functions in the CB to form a core exciton [17].

**IV. CONCLUSION**

In summary, photoinduced dynamics at W N$_{6,7}$ and O$_3$ edges in WS$_2$ are simultaneously measured by XUV core-level transient absorption spectroscopy. Picosecond hole relaxation and recombination dynamics in the valence band are extracted from the transient absorption spectra of the core-to-conduction-band transitions at the W O$_3$ edge. Lifetimes of core-excited states at the W O$_3$ edge and the W N$_{6,7}$ edge are obtained from XUV transient absorption spectra at negative time delays. While the lifetimes of W O$_3$ edge transitions are well below the duration of the optical pulse (~4 fs), core-exciton coherence lifetimes up to 11 fs are observed at the W N$_{6,7}$ edge. Global fitting of the XUV transient absorption spectra at short time delays reveals that in contrast to the direct field-induced core-exciton dynamics observed in insulators [11–13], carrier-induced modulation of core-exciton states dominates the dynamics at the few- to-femtoseconds timescale.

The drastically different behaviors between the absorption from the W 5$p$ and 4$f$ core orbitals in the same energy region suggest that in addition to macroscopic screening, the degree of localization of the core orbitals can contribute significantly to the core-level absorption lineshape and the formation of core excitons. The observation of carrier-modulated core-exciton transitions can serve as an initial step in further understanding and manipulating the dynamics of core excitons in condensed matter, and the extraction of hole dynamics at the W O$_3$ edge further advances the use of core-level TA spectroscopy in measuring carrier dynamics in transition metal dichalcogenides and their heterostructures.

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APPENDIX A: SAMPLE PREPARATION

The WS\(_2\) sample was synthesized by atomic layer deposition of WO\(_3\) thin films on 30-nm-thick silicon nitride membranes (Norcada Inc.). The tungsten oxide film was subsequently converted into sulfide in a tube furnace with H\(_2\)S. Before atomic layer deposition, 16-nm-thick silicon nitride films were deposited onto the Si frame of the silicon nitride windows using plasma-enhanced chemical vapor deposition (PECVD) to prevent silicon sulfide formation during the reaction with H\(_2\)S [66]. The passivated windows were then coated with WO\(_3\) using atomic layer deposition in an oxygen plasma [39]. The thickness of WO\(_3\) was calculated from the required thickness of WS\(_2\) using the ratio of the density between the two assuming no W loss in the reaction with H\(_2\)S. The thickness of the oxide film was characterized by in\(_{\text{situ}}\) spectroscopic ellipsometry. After the oxide deposition, the windows were put in a quartz boat and transferred into a tube furnace which was heated up to 600 \(^\circ\)C. H\(_2\)S [5 cm\(^3\)/min at STP (sccm)] and Ar (100 sccm) as a buffer gas was flowed into the tube to react with WO\(_3\). After 1 h of reaction, the H\(_2\)S flow was turned off while maintaining the Ar flow to prevent contamination from outside air, and the furnace was left to cool down. After the temperature decreased below 200 \(^\circ\)C, the Ar flow was switched to N\(_2\), and the samples were taken out after the instrument reached room temperature. To verify that the absorption peaks below 37 eV are not due to defect-induced color centers, the XUV absorption spectrum of the synthesized film was compared with the total electron yield (TEY) spectrum of single-crystal WS\(_2\) (2Dsemiconduc-

FIG. 8. (a) A typical XUV spectrum produced by high-harmonic generation in Kr and (b) the XUV spectra as a function of the carrier envelope phase of the driving pulse.

APPENDIX B: EXPERIMENTAL SETUP

The optical and XUV pulses in the experiment were produced by a Ti:sapphire carrier envelope phase (CEP) stabilized laser operating at 1 kHz (Femtoperpower Compact Pro seeded by Femtolaser Rainbow CEP3). The output of the Ti:sapphire laser was 1.8 mJ in pulse energy and approximately 30 fs in pulse duration. The laser beam was focused into a 1-m-long Ne-filled hollow core fiber to generate a supercontinuum spanning 500–1000 nm wavelength with self-phase modulation. A mechanical chopper was installed after the hollow core fiber to chop down the repetition rate to 100 Hz to prevent sample damage through excessive heating. The dispersion accumulated during pulse propagation was compensated by a set of broadband double-angle chirped mirrors (PC70, UltraFast Innovations) and a 2-mm-thick ammonium diphosphate crystal [67]. The beam was then separated into the probe and pump arm by a 9:1 broadband beamsplitter. Each arm was equipped with a pair of UV-graded fused silica wedges for dispersion fine-tuning. The probe beam was subsequently focused into a Kr gas jet to produce broadband XUV pulses (30–50 eV) via high-harmonic generation (Fig. 8). The XUV beam then traveled through a 100-nm-thick Al filter blocking the high-harmonic driving field and was focused onto the sample with a Au-coated toroidal mirror. The pump beam was time delayed with respect to the probe by a piezo-driven optical delay stage and was subsequently recombinated with the probe arm by an annular mirror. A 200-nm-thick Al filter is placed after the sample to prevent the pump beam reaching the XUV spectrometer. The XUV beam passing through the sample and the Al filter was dispersed by a flat-field grating onto an XUV CCD camera. The spectral energies were calibrated with the autoionization lines of Ar 3s3p\(^3\)np and Ne 2s2p\(^5\)np states [68,69]. The duration of the pump pulse was characterized by dispersion scan [70] to be \(\tau_{\text{pump}} = 4.2 \pm 0.1\) fs, and the spectrum and temporal profile of the pump pulse are shown.
in Figs. 9(a) and 9(b), respectively. The pulse energy of the pump beam was controlled by an iris, and the beam profile of the pump pulses was imaged directly at the sample position with a complementary metal-oxide semiconductor (CMOS) camera to calculate the pump fluence. During the XUV transient absorption experiment, the sample was raster scanned to prevent heat damage.

To correct for the drift of time delay during the experiments, an optical-XUV transient absorption measurement on Ar was conducted after each WS 2 transient absorption scan [4,7]. The Ar gas cell was mounted alongside the WS 2 sample. The suppression of Ar 3s3p6np autoionization lines by the optical pulse at 26–37 eV photon energies was measured [Fig. 9(c)] [53,69], and the time reference of each scan was determined by fitting the integrated absolute value of the transient absorption signal of the Ar 3s3p64p state along the energy axis and fitting it with a Gaussian error function [4]. The time axis of each WS 2 transient absorption scan was shifted according to its time-zero reference and the transient absorption signal \( \Delta A \) interpolated onto a uniform time delay grid. In addition to time-zero referencing, the cross-correlation time between the pump and probe pulses was estimated by the width of the error function rise [4,53], and the estimated pump-probe cross-correlation time is \( \tau_{\text{cr}} = 4.1 \pm 0.5 \) fs. The maximal cross-correlation time of the 4.2 fs pump pulse and a single-cycle driving pulse for high-harmonic generation centered at 730 nm [Fig. 9(a)] is \( \sqrt{4.2^2 + 2.4^2} \approx 4.8 \) fs. The experimentally measured cross-correlation time of 4.1 ± 0.5 fs is within the cross correlation of the pump pulse and a single-cycle driving pulse, and the maximum width of the XUV pulse train envelope is estimated to be \( \sqrt{(\max \tau_{\text{cr}})^2 - (\min \tau_{\text{pump}})^2} \approx 2.1 \) fs, less than two half cycles of the optical driving field. This indicates that the attosecond XUV pulse train consists of \( \leq 2 \) XUV bursts.

To provide a reference for future studies on carrier effects on core excitons in solids that cannot be prepared as thin films, we performed XUV transient reflectivity experiments on 40-nm-thick WS 2 thin films deposited on silicon wafers, which were synthesized alongside the samples for XUV transient absorption experiments (Appendix A). The measurements were taken on a beamline almost identical to the one for XUV transient absorption, except for the interaction geometry at the sample [71]. The optical pump and XUV probe pulses (\( p \) and s polarized, respectively) impinged on the sample surface with a 66° angle from the sample normal. The reflected XUV beam was directed into a spectrometer identical to the one used in absorption. A gold mirror was used as a reference to extract the absolute reflectivity of the WS 2 sample [71], and because of the relatively weak change in reflectivity, the data were processed using edge-pixel referencing [72]. The results of the XUV reflectivity measurements are detailed in Appendix G.

**APPENDIX C: ELECTRONIC STRUCTURE CALCULATIONS**

The electronic structure of bulk WS 2 is computed with the all-electron full-potential linearized augmented plane wave (FP-LAPW) method using the ELK code [41,73].
density functional theory (DFT) computation is conducted within the local spin density approximation (LSDA) [74]. Spin-orbit coupling effects are included, and the calculations are converged with a $k$ grid of $10 \times 10 \times 3$ $k$ points. A 4 eV blueshift is added to the calculated dielectric function in Fig. 1(a) to compensate the underestimated gap between the core levels and the CB in DFT calculations.

The number of photons available in the optical pulse to excite valence electron-hole pairs as a function of $k$ points (Fig. 4) is calculated with the formula

$$N_{ph,exc}(k) = \sum_{n_r,n_h} \int dE \, N_{ph}(E) \delta[E_{n_r}(k) - E_{n_h}(k)].$$

The number of photons as a function of photon energy $N_{ph}(E)$ is obtained from the measured spectrum of the pump pulse [Fig. 9(a)]. The energies of valence and conduction bands $n_r$ and $n_h$ are obtained from the calculated band structure [Fig. 4(b)].

### APPENDIX D: OPTICALLY EXCITED CARRIER DENSITY

The photoexcited carrier density $\rho_{exc}$ is estimated by calculating the number of absorbed photons in the 40-nm-thick WS$_2$ film per unit area $\sigma_{abs}$ divided by the thickness of the film $d$: $\rho_{exc} = \sigma_{abs}/d$. The number of photons absorbed per unit area can be calculated with the equation

$$\sigma_{abs} = \int d\omega \, \sigma_{inc}(\omega) f_{abs}(\omega),$$

where $\sigma_{inc}(\omega)$ is the number of incident photons per unit area with photon energy $\omega$ and $f_{abs}$ is the fraction of photons absorbed in the film. $\sigma_{inc}(\omega)$ can be calculated from the spectrum of the pump pulse and the measured fluence. The fraction of photons absorbed ($f_{abs}$) is calculated using the transfer matrix method including the 40-nm-thick WS$_2$ film and the silicon nitride window [75]. The refractive indices of WS$_2$ and silicon nitride are taken from Refs. [76,77]. We conducted the experiments with fluences ranging from 6 to 30 mJ/cm$^2$, and the resulting calculated excited carrier density ranges from $3 \times 10^{20}$ to $2 \times 10^{21}$ cm$^{-3}$. The carrier density per layer is calculated multiplying the carrier density by volume with the layer thickness of 6.2 Å [78].

### APPENDIX E: SINGULAR VALUE DECOMPOSITION

The XUV TA spectra between $-25$ and 25 fs below 37.5 eV are analyzed with global fitting via singular value decomposition (SVD), where the TA signal $\Delta A(t, E)$ is written as a matrix with rows and columns indicating different time $t$ and energy $E$, respectively (Figs. 5 and 10). The TA matrix $\Delta A(t, E)$ is then decomposed with SVD into $\Delta A(t, E) = U(t)^T SV(E)$, where $U$ and $V$ are unitary matrices consisting of singular vectors $\{u_m(t)\}$ and $\{v_n(E)\}$, respectively. $S$ is a rectangular diagonal matrix, and the diagonal matrix elements $S_{mn} = s_{mn}$ are singular values ranked in descending order. The reconstruction of the TA signal $\Delta A_{rec}(t, E)$ by components up to the $n$th rank is defined as $\Delta A_{rec}(t, E) = \sum_{m=1}^{n} s_{mn} u_m(t) v_n(E)$. Note that the SVD approach is based on the assumption that the transient absorption spectra can be represented by a linear combination of components from different contributions. Here, such an assumption is valid because the carrier-induced spectral modification is the dominant contributor [Figs. 5(a)–5(c)] and the direct field-induced effects can be regarded as a minor component.

To verify whether phonon-induced dephasing contributes significantly to the decay of core excitons, we focus on the decay dynamics of the largest component in the SVD, $u_1(t)$ [Figs. 5(c) and 5(f), red dots], at negative delays. The largest SVD component is plotted in logarithmic scale in Fig. 11, and the component $\log_{10} u_1(t)$ is fitted with a quadratic function $at^2 + bt + c$. The fitted coefficients of the quadratic function $\log_{10} u_1(t)$ for core exciton A and B and core exciton C (Fig. 11).

<table>
<thead>
<tr>
<th></th>
<th>A and B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a$</td>
<td>$(6 \pm 2) \times 10^{-4}$</td>
<td>$(5 \pm 2) \times 10^{-4}$</td>
</tr>
<tr>
<td>$b$</td>
<td>$0.061 \pm 0.008$</td>
<td>$0.084 \pm 0.007$</td>
</tr>
<tr>
<td>$c$</td>
<td>$-1.21 \pm 0.06$</td>
<td>$-0.4 \pm 0.07$</td>
</tr>
</tbody>
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are listed in Table I, showing that the quadratic term $a$ is two orders of magnitude smaller than the linear term $b$. In addition, the fitted $a$ are positive rather than negative as expected for a Gaussian function. This indicates that the decay of XUV TA signal at negative time delays is exponential rather than Gaussian and the effect of phonon-induced dephasing is insignificant.

APPENDIX F: COMPARISON WITH XUV TOTAL ELECTRON YIELD OF SINGLE-CRYSTAL WS$_2$

To verify that the absorption peaks below 37 eV are not due to defect-induced color centers, the XUV absorption spectrum of the synthesized film is compared with the total electron yield (TEY) spectrum of single-crystal WS$_2$ (2Dsemiconductors USA) measured at Beamline 4.0.3 at the Advanced Light
Source. The measured TEY of the single-crystal sample is shown in Fig. 12. The spectrum is cut off at 34.5 eV due to the lack of XUV photons below 34.5 eV at the undulator beamline. The measured TEY of WS₂ shown in Fig. 12 is normalized by the measured TEY of a gold film: \( \text{TEY}_{\text{norm}} = \frac{\text{TEY}_{\text{sample}}}{\text{TEY}_{\text{Au}}}. \)

**APPENDIX G: ADDITIONAL STATIC AND TRANSIENT REFLECTIVITY MEASUREMENTS**

Here, we provide measurements of core-exciton dynamics in the presence of photoexcited carriers in reflectivity geometry. Certain materials are challenging to synthesize as thin films for XUV absorption measurements; yet the analysis of reflectivity data alone is challenging and often relies on Kramers-Kronig transforms. Therefore the data presented below can serve as a useful reference point for future studies of materials other than WS₂.

The absolute static reflectivity of WS₂ deposited on a silicon wafer, taken at 66° from normal [Fig. 13(a)], shows that while core exciton C is very visible, core excitons A and B are difficult to resolve. Nevertheless, reflectivity changes are clearly observed [shown in Figs. 13(b) and 13(c) at +10 fs delay] for each peak and share the same shape: a reduced reflectivity at the center of the exciton lineshape and a slight increase on each side of it. The comparison [Fig. 13(d)] with the transient absorption reported in the main text shows that the two observables are consistent with each other. These results display how the core-excitonic lineshapes in reflection geometry are modified by the excitation of free carriers.

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**References**


