Ultrafast dynamics in the high-symmetry and in the charge density wave phase of 2H-NbSe₂

A. Anikin[®],¹ R. D. Schaller,² G. P. Wiederrecht,² E. R. Margine[®],³ I. I. Mazin[®],⁴ and G. Karapetrov[®],^{5,*}

¹Department of Physics, Drexel University, Philadelphia, Pennsylvania 19104, USA

²Center for Nanoscale Materials, Argonne National Laboratory, Lemont, Illinois 60439, USA

³Department of Physics, Applied Physics and Astronomy, Binghamton University–SUNY, Binghamton, New York 13902, USA

⁴Department of Physics and Astronomy, George Mason University, Fairfax, Virginia 22030, USA

⁵Department of Materials Science and Engineering, Drexel University, Philadelphia, Pennsylvania 19104, USA

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We investigate carrier and collective mode dynamics in 2H-NbSe₂ using time-resolved optical pump-probe spectroscopy and compare the results with first-principles calculations. Broadband ultrafast reflectivity studies of 2H-NbSe₂ in a wide temperature interval covering the normal, charge density wave (CDW) and superconducting phase were performed. Spectral features observed in the transient reflectivity experiment were associated with specific optical transitions obtained from band structure calculations. Displacive excitation of coherent phonons showed CDW-associated coherent oscillations of the soft phonon mode across the whole spectral range. Temperature evolution of this coherent phonon mode in the low-excitation linear regime shows softening of the mode down to the CDW transition temperature T_{CDW} with subsequent hardening below T_{CDW} . The global fit of the broadband probe data reveals four different relaxation times associated with characteristic electron-electron, electron-phonon, and phonon-phonon relaxation processes. From first-principles calculations of electron-phonon coupling we associate the few picosecond electron-phonon relaxation time τ_2 with a specific group of phonons with frequencies around 20 meV. On the other hand, the anomalously long relaxation time of $\tau_3 \sim 100$ ps is associated with anharmonicity-driven phonon-phonon scattering. All relaxation processes result from anomalies near the second order CDW phase transition that are reflected in the temperature dependencies of the characteristic relaxation times and amplitudes of coherent oscillations. At the highest fluences we observe electronic melting of the CDW and disappearance of the mode hardening below T_{CDW} .

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I. INTRODUCTION

Layered transition metal dichalcogenides (TMDs) are a large family of quasi-two-dimensional (2D) materials that often exhibit charge density waves (CDWs) and/or superconductivity (SC). Both CDW and SC are usually a product of electron phonon coupling, but in several dichalcogenide systems more exotic interactions have been discussed. For example, TiSe₂ is still discussed as a potential excitonic insulator, although the jury is still out [1]. Lack of inversion symmetry in some TMDs leads to interesting effects such as potentially chiral CDWs [2-4] and spin-momentum locking in single-layer NbSe₂ leading to Ising superconductivity [5]. Although the dichalcogenides have been extensively studied for over 50 years, many interesting topics remain open. Novel experimental methods that focus on high spatial and temporal resolution provide further insights into the interplay between electrons and phonons that leads to the CDW and SC phases. Femtosecond optical spectroscopy provides a unique opportunity to examine collective modes with superior frequency resolution and high dynamic range [6]. In this paper we explore femtosecond optical reflectivity and coherent phonon dynamics in 2H-NbSe₂ — material isostructural and similar to 2H-NbS₂ in that it exhibits strong anharmonicity at low temperatures, but nevertheless goes through both CDW and superconducting phase transitions with both phases coexisting below 7.2 K [7,8].

2H-NbSe₂ is a van der Waals material belonging to the $P6_3/mmc$ space group with a trigonal Nb layer stacked between two trigonal Se layers having periodicity of double Se-Nb-Se trilayer. It remains metallic from room temperature down to its superconducting transition at $T_c = 7.2$ K [9] with three hole and two electron Fermi surfaces [10], with different electron phonon coupling, leading to a two-gap superconductivity [11] and to switching of the Hall conductivity from p type at $T_{\rm H} > \sim 18$ K to *n* type below $T_{\rm H}$ [12]. At temperatures below 33.5 K an incommensurate CDW sets in, corresponding to simultaneous condensation of three CDWs with the wave vectors $q\{1, 0, 0\}, q\{0, 1, 0\}, q\{1, 1, 0\}$, where $q \approx 1/3$, close to a reconstruction with the 3×3 periodicity [13]. CDW-like fluctuations persist up to 100 K in bulk NbSe₂, well above the thermodynamic second order CDW phase transition temperature of 33.5 K [14,15]. Under pressure the CDW instability is rapidly suppressed, with only a minor effect on superconductivity, indicating that the latter is not due to the soft modes related to the CDW instability [16–19].

Although initially many studies linked the CDW formation to a simple Peierls instability and Fermi surface nesting, detailed band structure calculations [10,20,21] and experiments [22–24] show that electron-phonon matrix elements play the decisive role in creating the CDW instability,

^{*}Corresponding author: goran@drexel.edu

while the Fermi surface nesting occurs at a wrong wave vector [21,24]. Moreover, experimentally and theoretically determined Lindhart susceptibility function shows no peaks [24,25] and only a very broad maximum at the corresponding wave vector q_{CDW} , extending to almost half the Brillouin zone in the ΓM direction [22], which indicates a minor role played by electronic nesting. The amplitude of CDW charge supermodulation is close to 9%, resulting in a very small kink (\sim 5%) in the metallic temperature dependence of resistivity at T_{CDW} [9,26,27]. The theory of strong q-dependent electron-phonon coupling (EPC) was proposed as an attempt to explain the structural and electronic transition [20,28]. The calculations are consistent with the experimental phonon dispersion [25]. Experimentally, qdependent strong EPC was confirmed by Weber et al. [22] and further studied in detail later on [29–31].

An incommensurate CDW in NbSe₂ forms along the ΓM direction in reciprocal space with $q_{\text{CDW}} = \frac{2\pi}{3a}(1-\delta)$ with $\delta \approx 0.02$. The CDW gaps the inner Fermi surface cylinder around the *K* point of the first Brillouin zone, as first proposed theoretically [10,16,20,28] and shown experimentally [11,22,23,29]. The longitudinal optical phonon Σ branch couples to the Nb 4*d* electrons near *K* and *K'* pockets of the double-walled Fermi surface sheets, and, as we will discuss later in the paper, is instrumental for superconductivity, but not for the CDW. The charge ordering induces a negative carrier plasmon dispersion leading to a minimum at $q = q_{\text{CDW}}$ of the $\sim 1 \text{ eV}$ plasmon, which represents an electronic signature of fluctuating CDW order parameter [32,33].

Electron-phonon coupling cannot always stabilize CDW order even in materials that are structurally and electronically almost equivalent, such as NbSe₂ and NbS₂. It seems that the strength of the electron-phonon interaction has to overcome the anharmonic effects, as the temperature is lowered. In NbS₂, anharmonicity prevents condensation of CDW phase down to T = 0 K [34], although the fluctuating CDW order parameter is present at low temperatures [35]. Anharmonic effects can be reduced by making the system more two-dimensional. Reduction of dimensionality, i.e., electronic decoupling between the van der Waals layers, reduces conductivity and charge screening [18,36], as well as prevents ionic charge transfer [37] between the transition metal cation and dichalcogenide anions. Thus in isolated monolayer NbS₂ the CDW phase is stable and wins over the anharmonic effects below 5 K [38,39]. NbSe₂ in a monolayer form experiences enhancement of T_{CDW} to 145 K, which exceeds the onset temperature of the CDW fluctuations in the bulk system (~ 100 K) [40,41]. Even in bulk NbSe₂, the surface layer seems to experience CDW fluctuations as high as room temperature, as observed by surface sensitive techniques like small-angle inelastic x-ray scattering [42] or angle-resolved photoemission spectroscopy [11]. On the other hand, effects that enhance anharmonicity and reduce the anisotropy, such as hydrostatic pressure, suppress the T_{CDW} [31,43].

In this paper we explore anharmonic processes in NbSe₂ using transient optical reflectivity. We explore the temperature evolution of the anharmonicity in order to understand the electronic processes taking place near the normal to incommensurate CDW second order phase transition in 2H-NbSe₂. We find that the femtosecond optical pulse leads to

excitation of coherent phonon with A_1 symmetry [44] that corresponds to the two-phonon peak observed in Raman spectroscopy at 5.4 THz (180 cm^{-1}) at 300 K [42,45–49]. We observe strong damping of the coherent phonon and correlate it with the broad spectral width of the Raman two-phonon peak. As temperature is lowered towards T_{CDW} , the coherent phonon softens and the damping enhances, in agreement with expected strong electron-phonon coupling near the CDW transition in this system. Below the T_{CDW} the coherent phonon hardens and dephasing time increases, effects that have not been previously observed with optical probes. We believe that blueshifting of the coherent phonon below T_{CDW} is a result of strengthening of the CDW order parameter as temperature is lowered away from T_{CDW} . The contribution to the Raman tensor of the phase modes, which are normally the acoustic modes of the incommensurate CDW state, is raised to a finite frequency if the inversion symmetry is broken by the layer stacking [50], as in single unit cell thick NbSe₂ [40]. But in bulk NbSe₂, the CDW phase mode in 2*H*-NbSe₂ is inaccessible by Raman since there is no breaking of inversion symmetry. As the fluence of the pump pulse increases, nonthermal melting of CDW takes place and the blueshift of the coherent phonon disappears below T_{CDW} . This confirms that the CDW mode might be involved in the low-temperature energy shift of the coherent two-phonon peak as has been observed in other CDW materials [51].

II. EXPERIMENTAL AND COMPUTATIONAL DETAILS

High quality single crystals of 2H-NbSe₂ were grown by the chemical vapor transport method with iodine as the transport agent. Resistivity measurements revealed $R(300 \text{ K})/R(8 \text{ K}) \approx 100, T_c = 7.2 \text{ K}, \text{ and } T_{CDW} = 33.5 \text{ K},$ values similar to the highest quality crystals (see the Supplementary Material [52]) [53–56]. A freshly cleaved surface of the single crystal was exposed before introducing the sample into the optical cryostat (Janis Model PTSHI-4 (Janis Research Co., LLC, USA) equipped with the cryocooler Model SRP-062B (Sumitomo Heavy Industries, Ltd, Japan). Pump-probe reflectivity measurements were conducted using a linearly polarized pump laser at 800 nm (1.55 eV) operating at a repetition rate of 2 kHz and a pulse width of \sim 35 fs [full width at half maximum (FWHM)] (Fig. 1). Part of the 800-nm output was focused on CaF₂ crystal to produce white light probe pulses that were variably delayed relative



FIG. 1. Experimental setup: split output from an 800-nm 35-fs Ti:sapphire laser produces 800-nm pulses at 2 kHz, part of which are focused on a CaF₂ crystal to produce white light supercontinuum probe pulses. The chopped 800-nm pump beam was directed at the sample at $\sim 20^{\circ}$ incidence.

to the pump pulse arrival time at the sample. The pump pulses were chopped at 1 kHz to enable a probe signal with and without photoexcitation to be collected and compared. The probe signal was monitored using a complementary metal-oxide semiconductor camera. Spectra were corrected for chirp using a rising edge fit spanning across multiple wavelengths and applying the quadratic relation $-t = a\lambda^2 + b^2$ $b\lambda + c$. The supercontinuum generation introduces the group velocity dispersion that negligibly increases the FWHM of the cross-correlation function from 35 to 35.3 fs, i.e., by less than 1%. The measured pump beam diameter is approximately 600 μ m. Calculated pulse penetration depth is about 30 nm. To probe the dynamics, a linearly polarized white light probe beam was used with diameter of 180 μ m. The measurements were taken at temperature intervals from 3 to 150 K at several pump fluences. Data analysis of the changes in optical density were extracted using the open-source software package GLOTARAN [57].

First-principles calculations were performed with density functional theory (DFT) using the Quantum ESPRESSO (OE) [58] code. We employed optimized norm-conserving Vanderbilt (ONCV) pseudopotentials [59,60] with the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional in the generalized gradient approximation [61], where the Nb $4s^24p^64d^35s^2$ and Se $4s^24p^4$ orbitals were included as valence electrons. All calculations were performed for the experimental lattice parameters at ambient pressure [22] with relaxed internal coordinates. We used a plane wave kinetic-energy cutoff value of 80 Ry, and the electronic and vibrational Brillouin zones were sampled using $24 \times 24 \times 8$ and $6 \times 6 \times 2$ points, respectively. A Methfessel and Paxton smearing [62] width of 0.025 Ry was used in order to resolve the CDW instability. The electron-phonon calculations were performed with the EPW code [63,64] of the QE distribution, in conjunction with the WANNIER90 library [65,66]. We used 22 maximally localized Wannier functions (five d orbitals for each Nb atom and three p orbitals for each Se atom) and a uniform Γ -centered $12 \times 12 \times 4$ electron-momentum grid. The linewidth associated with the electron-phonon interaction $\gamma_{\nu,q}$ of mode ν with momentum **q** were evaluated on a uniform $240 \times 240 \times 60$ k-point grid. The isotropic Eliashberg spectral function and the phonon linewidth spectral function were evaluated on uniform $80 \times 80 \times 40$ k- and $40 \times 40 \times 20$ qpoint grids. The Dirac deltas were replaced by Lorentzians of width 25 meV (electrons) and 0.05 meV (phonons).

III. RESULTS AND DISCUSSION

Figure 2 displays the transient change in reflectivity *R*, induced by the pump pulse, defined as signal $\Delta A(\omega, T) =$ $-\log_{10}[R_{\text{pump}}(\omega, T)/R_{\text{no pump}}(\omega, T)]$. The data were collected at 3 K with 0.5 mJ/cm² pump fluence. The positive values on the color map correspond to a decrease in reflectivity (increased optical density) due to the pump pulse, while the negative values correspond to an increase in reflectivity (decreased optical density) induced by the pump pulse.

The physics of the transient response is, in the first approximation, rather simple. First, the laser pump excites electrons, by triggering electronic transitions with the energy equal to the laser photon energy. On a rather short time



FIG. 2. (a) Transient reflectivity data taken at 3 K with the pump fluence of 0.5 mJ/cm^2 . Arrows indicate the coherent oscillations within the first picosecond. (b) The kinetic trace at $\lambda = 500 \text{ nm}$ showing the coherent oscillations in the inset. The experimental points are connected with a spline (black line).

scale, $\tau_1 \sim 0.3$ ps, these electrons thermalize among themselves via electron-electron scattering. Typically, the resulting electronic temperature is $T_{\rm e} \sim 1000 \, \text{K}$. This, first of all, leads to a modification of the electronic absorption $\varepsilon_2(\omega) =$ const $\sum_{k,i,j} (f_{k,i} - f_{k,j}) \delta(E_{k,j} - E_{k,i} - \omega)$, where f(T) are Fermi occupation factors, and we have absorbed the average dipole matrix elements into the prefactor. Taking the derivative with respect to T, we immediately see that summation is now over the product of the two δ functions, $\sum_{k,i,j} \delta(E_{k,i} - \mu) \delta(E_{k,j} - \omega)$, where μ is the Fermi energy. This is equivalent to integrating over the line where the two isoenergetic surfaces intersect (Fig. 3), with the weight equal to the inverse cross product of the relevant electron velocities [67]. The sign does not depend on whether the transition is to or from the Fermi level, but on the details of the joint density of states and matrix elements.

Three features can be seen in Fig. 2. The strongest is at 485 nm (2.6 eV) accompanied by a weaker one at 570 nm



FIG. 3. (a) Electronic structure of NbSe₂. (b)–(d) The Fermi surfaces are depicted as opaque, the isosurfaces corresponding to the energies $E_1 = 1.95 \text{ eV}$ (635 nm) (b), $E_2 = 2.18 \text{ eV}$ (570 nm) (c), and $E_3 = 2.56 \text{ eV}$ (485 nm) (d) as semitransparent. The colors correspond to the first panel (a). Vertical lines in the first panel show examples of the corresponding transitions.

(2.18 eV) and an even weaker one, of the opposite sign, at 635 nm (1.95 eV). These wavelengths correspond to features that have been observed in static reflectivity [68–70]. The calculated reflectivity together with the optical absorption $\varepsilon_2(\omega)$ are shown in Fig. 4. The energy is scaled by 4% to ensure better agreement with the experiment [67–69]. This is not uncommon in optical calculation, reflecting renormalization of the bandwidth due to strong interaction. For instance, in Ref. [71] the best agreement of the calculated optical absorption was found when all features were redshifted by ~10%. This was ascribed to the bandwidth renormalization due to magnon exchange. The effect is, apparently, considerably weaker in NbSe₂, consistent with the presence of spin fluctuation there [72], albeit, of course, much weaker than in CrO₂. We can clearly see a very strong and sharp peak



FIG. 4. Calculated reflectivity (red) and the imaginary part of the dielectric function (green). Three arrows indicate three frequencies at which experimental features are observed.

exactly at the frequency of our strongest feature at 2.6 eV and a much weaker one at 2.18 eV. We also see a tiny shoulder corresponding to our third feature. Band analysis reveals that while the features appear as combinations of several interband transitions, each one is dominated by the one transition, depicted in Fig. 3.

Another interesting piece of information that can be extracted from our data is the oscillatory behavior of the transient response. This is typical for displacive excitation of coherent phonons (DECP), whereupon nearly instantaneous heating of the electron subsystem to $T_{\rm e} \sim 1000 \, {\rm K}$ shifts the equilibrium positions of the ions that are not fixed by symmetry (in our case, the z position of Se), and then the ions oscillate around the new position with a pattern consistent with the A_{1g} symmetry. We see no signature of the actual A_{1g} phonon, which exists at 6.9 THz ($\sim 230 \,\mathrm{cm}^{-1}$), presumably, because this phonon does not modify the dielectric response at 400-700 nm. However, we see a broad frequency range of oscillations centered at about 3.5-4.5 THz $(120-150 \text{ cm}^{-1})$ [Fig. 5(c)], exactly matching the well-known broad feature in the A_{1g} Raman spectra [42,45–48,73], usually interpreted as a composite excitation of two phonons propagating in the opposite direction (a standing wave) [74,75]. Generally, such a feature would be weak compared to a single A_{1g} phonon, but if such a wave modulates the dielectric function in the relevant energy range, which is highly possible (because, as opposed to the A_{1g} mode, it involved Nb displacements), it will be the only one observed.

Let us analyze this feature in more detail (Fig. 5). Highly damped coherent oscillations can be resolved within the first picosecond across the whole measured spectral interval. Oscillations were detected at all sample temperatures which allows us to study their temperature evolution. Since the oscillation amplitudes are relatively weak, we averaged the kinetic traces from within ± 5 nm around the central values of 500 nm [Fig. 5(a)] and 560 nm [Fig. 5(b)] wavelengths. The traces



FIG. 5. (a), (b) Kinetic traces near 500 and 560 nm showing the coherent oscillations (yellow circles) and the fits using exponentially modified peak function and damped sine function (red lines). (c) Extracted temperature dependences of oscillation frequencies and dephasing rates (d).

were fitted by exponentially modified peak function convoluted with damped sine function to capture the oscillations [Figs. 5(a) and 5(b)]. Extracted temperature dependence of the frequency and dephasing rate are shown in Figs. 5(c) and 5(d).

We can clearly see softening of this composite excitation from 150 K down to T_{CDW} . At the same time, the dephasing rate gets slightly shorter. Below T_{CDW} this excitation hardens and the damping rapidly decreases. However, the frequency remains finite at $T = T_{CDW}$. One can understand that by looking at the calculated phonon dispersion near $q = q_{CDW}$. Our calculations (Fig. 6), consistent with previously published results by Weber et al. [23], indicate that the soft mode frequency is minimal at q_{CDW} and then increases quadratically as one moves away from q_{CDW} , and reaches a maximum of about 3 THz (12 meV) at some $q = q_{\text{max}}$, (approximately halfway between Γ and *M*) which is temperature independent. The minimum at q_{CDW} , as measured by inelastic x-ray scattering [22], shifts with temperature in a mean-field manner as $\sqrt{\frac{T}{T_{\text{CDW}}}}$ -1 and rises about twice that fast again below T_{CDW} .

Approximating the frequency of this branch as

$$\omega(T,q) = 2A \frac{|q - q_{\rm CDW}|}{q_{\rm CDW}^2} + B \sqrt{\frac{T}{T_{\rm CDW}} - 1},$$
 (1)

where $A \sim 3$ THz and *B*, from Ref. [22], ~0.8 THz, and averaging over $|q - q_{CDW}| < q_{max}$ (where $q_{max} \approx \frac{1}{2}|\Gamma M|$), we get $\omega(T, q) = 0.6A + B\sqrt{\frac{T}{T_{CDW}} - 1} \approx 2$ THz + $0.8\sqrt{\frac{T}{T_{CDW}} - 1}$ THz. Below T_{CDW} , *B*, as extracted from Ref. [22], is about twice as large. As our results show (Fig. 6), this scenario describes our data rather well, strongly suggesting that the two-phonon mode is composed of the phonons of the same branch that is responsible for the CDW transition, just averaged over a large part of the Brillouin zone.

It is worth noting that the constants A and B above are not adjustable parameters but are estimated from either neutron scattering or first-principles calculations. A functional form very similar to Eq. (1) was proposed previously [48] based on an *ad hoc* assumption that the composite excitation is a



FIG. 6. Calculated (a) phonon dispersion, (b) phonon density of states, (c) Eliashberg spectral function $\alpha^2 F(\omega)$ and integrated electronphonon coupling strength λ , and (d) electron-phonon linewidth spectral function $\gamma(\omega)$. In (a) the size of the bars is proportional to the electron-phonon contribution to the phonon linewidth $\gamma_{\nu,q}$.



FIG. 7. Frequency of the soft mode and the dephasing rate as a function of temperature. The triangles and squares are experimental data, and the solid line is a fit of the data with the anharmonic decay model in Eq. (1).

combination of a soft phonon with $q = q_{\text{CDW}}$ and another phonon from a nonsoft branch with $q = -q_{\text{CDW}}$. However, this assumption lacks any physical justification so we do not consider it as a viable alternative. The dephasing time can be obtained by averaging the population numbers in a similar way,

$$\Gamma(T) = \Gamma_0 + \Gamma_{anh} \langle 1 + 2n(\omega) \rangle_q, \qquad (2)$$

where Γ_0 is the background contribution, Γ_{anh} is the proportionality constant, and $n(\omega)$ is the thermal occupation number of the phonon mode. As shown in Fig. 7, the experimental data are well described by the proposed anharmonic decay model [Eq. (2)].

Now we can turn to the analysis of the transient reflectivity dynamics. The transient optical response $\Delta A(\lambda)$ has been fit-

ted using the global fit approach in GLOTARAN [57] as follows:

$$\Delta A(\lambda) = \sum_{i} \text{IRF} \otimes \text{DAS}_{i}(\lambda) \ e^{-t/\tau_{i}}, \tag{3}$$

where the Gaussian instrument response function (IRF) is being convoluted (\otimes) with global exponential decays with starting magnitudes of the decay associated spectra (DAS_i) and decay constants τ_i characterizing a particular relaxation channel. Typically, there are three distinct relaxation scales: τ_{e-e} , the thermalization time of the electron subsystem due to electron-electron scattering, τ_{e-p} , the time scale at which electron thermalize with the phonons via electron-phonon coupling, and τ_{diss} , the time for the heat to dissipate away from the laser spot.

In our experiment, however, we can identify four different relaxation times, τ_i , having temperature evolutions shown in Figs. 8(a)–8(d). These time scales are on the order of ~100 fs, 1 ps, 100 ps, and 1 ns. The fastest component, τ_1 , is extracted from the sharp negative peak that appears between 520 and 680 nm at the time scale of ~200 fs. Since the probe has a pulse width of ~35 fs, we cannot determine this relaxation time with great accuracy although this peak has a significant negative amplitude. This is, obviously, the standard τ_{e-e} relaxation time.

The longest relaxation time τ_4 is on the nanosecond scale and it is easily identifiable as τ_{diss} . The temperature evolution shows a slowdown of the relaxation process at T_{CDW} that is a signature of the second order CDW phase transition and is accompanied by an anomaly in elastic modulus and thermal conductivity at T_{CDW} [76]. At the highest fluence when CDW's electronic modulation is destroyed, the anomaly in τ_4 at T_{CDW} disappears.

Most interesting are the second and the third relaxation constants with 1- and 100-ps time scales. They have exponential temperature dependence, suggestive of electron-phonon and phonon-phonon scattering. In order to verify this assumption we have performed first-principles calculations of



FIG. 8. Temperature dependencies of relaxation constants (a)-(d) and associated changes in reflectivities (e)-(h).

the electron-phonon coupling (see the Supplemental Material [52]). According to Allen's theory [77], relaxation of electrons with generation of a particular phonon (ν, \mathbf{q}) of the branch ν with the wave vector \mathbf{q} proceeds with the rate $1/\tau_{e-p} \propto \lambda_{\nu,q} \omega_{\nu,q}^2 = \gamma_{\nu,q}$, where $\lambda_{\nu,q}$ is the coupling constant with this particular mode and $\gamma_{\nu,q}$ is the electron-phonon linewidth for the same mode. One may naively think that this process will be dominated by the soft mode with the large λ , but this is not the case. Rather, it is dominated by a group of phonons with frequencies of ~20 meV (4.8–5 THz) [23].

We conclude that the *second* relaxation time, $\tau_2 = \tau_{e-p}$, characterizes electron relaxation into this particular group of phonons, while the *third* one is due to (much slower) thermalization of these phonons with the rest of the phonon subsystem, via relatively slow, anharmonicity-driven phonon-phonon scattering. Thus, $\tau_3 = \tau_{p-p}$.

Both τ_2 and τ_3 decay rapidly with temperature above T_{CDW} which is a result of an increase in phonon population number with temperature that leads to faster relaxation in both channels. As the temperature is lowered below T_{CDW} we observe a decrease in relaxation times for all fluences, except for the highest one of 1 mJ/cm². This may be indirect evidence of electronic melting of the CDW state by a rapid flood of a large number of hot charge carriers, as observed in other CDW systems using pump-probe techniques [78,79].

Changes in optical density associated with each time constant are shown in Figs. 8(e)–8(h). For τ_2 , τ_3 , and τ_4 the spectral distribution has similar character—we observe features that correspond to transitions observed in static reflectivity at 485 nm (2.6 eV), 580 nm (2.15 eV), and 635 nm (1.95 eV) as described earlier in the text. At the transition near 485 nm, dramatic changes in optical density are seen when the sample enters the superconducting state. This is followed by a negligible change between 10 and 40 K within the CDW phase transition interval. Further detailed DFT calculation of the optical transitions are necessary to gain better understanding of the features in DAS(λ) amplitudes' behavior.

IV. CONCLUSIONS

We have performed a broadband ultrafast reflectivity study of 2H-NbSe₂ in a wide temperature interval covering the normal, CDW, and superconducting phase. Spectral features observed in the transient reflectivity experiment were associated with specific optical transitions obtained from band structure calculations. Displacive excitation of coherent phonons showed CDW-associated coherent oscillations of the soft phonon mode across the whole spectral range. Temperature evolution of this coherent phonon mode in the low-excitation linear regime shows softening of the mode down to T_{CDW} with subsequent hardening below T_{CDW} . A global fit of the broadband probe data reveals four different relaxation times associated with characteristic electron-electron, electron-phonon, and phonon-phonon relaxation times. From first-principles calculations of electron-phonon coupling we associate the few ps electron-phonon relaxation time τ_2 with a specific group of phonons with frequencies of \sim 20 meV. On the other hand, an anomalously long relaxation time of $\tau_3 \sim 100 \,\mathrm{ps}$ is associated with anharmonicity-driven phonon-phonon scattering. All of the relaxation processes experience anomalies near the second order phase transition that are reflected in the temperature dependencies of the characteristic relaxation times and amplitudes of optical densities.

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