Selective optical probing of the charge-density-wave phases in NbSe₃

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(Received 27 October 2006; published 21 March 2007)

We report a study of the ultrafast single particle dynamics in quasi-one-dimensional NbSe₃ whisker crystals using a two-color pump-probe measurement. A selective enhancement of the relaxation dynamics connected with one of the two charge-density-wave (CDW) phases was realized by tuning the probe energy to around 1.5 eV, in which the temperature dependence across the corresponding phase transition shows a pronounced anomaly. A similar selective excitation was possible in the coherent CDW oscillation.

DOI: 10.1103/PhysRevB.75.115120 PACS number(s): 71.45.Lr, 78.47.+p

I. INTRODUCTION

Photoinduced absorption and/or reflection spectroscopy using ultrafast laser pulses has become a powerful tool for studying the excitation dynamics of phase transition materials.¹⁻⁴ Here, the time-resolved data allows us to follow the nonequilibrium single-particle (SP) and collective electron dynamics that are inaccessible with other conventional techniques. Recently, this optical technique was applied to materials characterized by the coexistence of multiple phases,¹⁻³ which involves interesting phenomena such as high-$T_C$ superconductivity and quantum fluctuation effects. However, in this case, the transient signal becomes complicated by a superposition of relaxation components derived from the individual phases. Therefore, the selective probing of individual phases is essential if we are to obtain precise measurements.

In this paper, we demonstrate that the phase-selective measurement of NbSe₃ crystals is possible by using a two-color optical-pump-probe measurement, which allows us to evaluate the photoexcited SP dynamics precisely even in coincidence with different phases. The temperature dependence of the transient data shows a dramatic change in its amplitude and decay time across $T_C$. The magnitude of these changes strongly depends on the probing energy, which allows the selective enhancement of the signal of the corresponding charge-density-wave (CDW) phase.

II. EXPERIMENT

The samples studied in the present work are NbSe₃ whisker-shaped single crystals with lengths of a few mm along the conducting axis and widths of $\sim 50 \mu m$. They were synthesized by the chemical vapor transport method. NbSe₃ is the most extensively studied quasi-one-dimensional (1D) CDW compound.⁶,⁷ This compound undergoes two independent Peierls transitions to CDW phases at $T_{C1}=145 \text{ K}$ and $T_{C2}=59 \text{ K}$. These two CDWs occur on two of three types of chains with different nesting conditions in k space. The other chains remain metallic down to the lowest temperatures. Therefore NbSe₃ is regarded as a multiple phase system.

Previous pump-probe measurements were performed with a conventional lens, and by restricting the CDW compounds to those with a large single-domain size. Here, we employed a two-color pump and probe setup with an objective lens, in which the coaxial configuration between the pump and probe beams increases the spatial resolution up to the diffraction limit, and allows reflectivity changes with a high signal to noise ratio even in the thin whisker samples.

Time-resolved data for the photoinduced dynamics of CDWs serve as snapshots of reflectivity changes ($\Delta R$). For the excitation source, we used an optical parametric oscillator (OPO) pumped by a mode-locked Ti:sapphire laser (76 MHz repetition rate). The time resolution was estimated to be around 200 fs from cross correlation measurements. The probe energy can be set from 1.39 to 1.77 eV by using a fundamental pulse and from 1.00 to 1.16 eV by using the OPO. Note that these excitation energies are far greater than the CDW gaps [typically around several tens of meV (Refs. 8 and 9)]. Depending on the experimental configuration, the pump and probe beams can be generated by either the Ti:sapphire laser or the OPO. The pump and probe pulse polarizations were parallel to the conducting axis because $\Delta R$ is dominant in this direction, and the pulses were focused through an objective lens onto a single-crystal region. The pump and probe fluences were $\sim 40 \mu J/cm^2$ and $\sim 10 \mu J/cm^2$, respectively. The pulse overlap with a diameter of 10 μm on the sample surface was monitored using a charge coupled device camera, and was kept at a fixed position during the measurements. The steady state heating caused by the laser was accounted for by measuring the excitation power dependence. The sample was mounted on a helium-flow cryostat.

III. RESULTS AND DISCUSSIONS

Figures 1(a) and 1(b) show two typical examples of transient $\Delta R$ obtained at probe energies of 1.56 and 1.07 eV, respectively. Both sets of data are strongly temperature dependent, and are qualitatively consistent with previous measurements for similar quasi-1D compounds,¹⁰ in which the transient $\Delta R$ observed at temperatures well below $T_C$ consists of two components: one is exponential decay, which predominantly reflects the photoexcited SP dynamics, and the other is coherent collective oscillations. We focus first on the former component.

Since the pump pulse at near-infrared energy can excite the SPs into higher-energy states far above the CDW gap, a quick relaxation to states near the band edge results in an
abrupt change of $\Delta R$. The subsequent exponential decay reflects the transient density change of the SPs accumulated in the vicinity of the gap. When approaching $T_{C1}$’s from below, this component shows an extended decay time with a lower amplitude. This is known as precursor behavior related to gap closing, which occurs due to the efficient reabsorption of nonequilibrium phonons across the gap.\(^\text{(1)}\) Above $T_{C1}$, i.e., above the temperature at which all chains are metallic, $\Delta R$ becomes very small, thus suggesting that the metallic chain contributions are negligible. When we consider the SP relaxation process in each CDW phase separately, the transient $\Delta R$ in NbSe\(_3\) is described by

\[
\Delta R(t, T, \omega_{pr}) \propto \begin{cases} 
S_1(T, \omega_{pr})e^{-\dot{\tau}_1(t)} + S_2(T, \omega_{pr})e^{-\dot{\tau}_2(t)} & (T \leq T_{C2}), \\
S_1(T, \omega_{pr})e^{-\dot{\tau}_1(t)} & (T_{C2} < T \leq T_{C1}),
\end{cases}
\]

where $S_n$ and $\dot{\tau}_n$ are the amplitudes and the relaxation times of the SP component, respectively, for each CDW phase ($n=1$ for $T_{C1}$ and $n=2$ for $T_{C2}$). Note that the $S_n$ is mainly determined by the optical transition probabilities at the probe energy ($\omega_{pr}$). On the other hand, the pump-induced SPs for above-gap excitation average their distribution over the near-gap states during the instantaneous relaxation, resulting in a minor contribution to $S_n$. In addition, we mainly focus on the fraction of magnitude $S_n/S_m$ ($n \neq m$). In this case, the effect of the number of SPs, which is slightly changed in accordance with the pump energy, is cancelled out. Indeed, we found that the outline of the transient $\Delta R$ of the 1.07/1.56 eV (pump-probe) excitation was almost identical to the results for 1.56/1.56 eV (not shown). Therefore we define the characteristic amplitude as a function of $\omega_{pr}$.

The two data sets in Fig. 1 show qualitatively the same temperature dependence but differ quantitatively. In particular, in the temperature range just below $T_{C2}$ where the $T_{C1}$ and $T_{C2}$ phases coexist, $\Delta R$ in (a) has a much longer decay time and a larger amplitude reduction than $\Delta R$ in (b). These observations indicate that we can selectively probe the photoexcited SP dynamics in the $T_{C2}$ phase as described in detail below.

In Fig. 2, we plot the effective decay time ($\tau_{\text{eff}}$) and the amplitude ($S_{\text{eff}}$) at probe energies of (a) 1.56 eV and (b) 1.07 eV, in which we have fitted the exponential function to the data without distinguishing the components given in Eq. (1). In the $T_{C2}$ to $T_{C1}$ temperature range, where $S_{\text{eff}}$ and $\tau_{\text{eff}}$ are actually identical to $S_1$ and $\tau_1$, respectively, the results show similar temperature dependences for (a) and (b) [compare the inset of (a) with (b)]. As the temperature approaches $T_{C1}$ from below, $S_1(T)$ decreases suddenly and $\tau_1(T)$ increases divergently, suggesting a reduction in the gap energy of the $T_{C1}$ phase. In contrast, below $T_{C2}$, there is a significant difference between (a) and (b). $S_{\text{eff}}$ in (a) shows a great reduction at $T_{C2}$ and its magnitude is 5 times greater than that at $T_{C1}$, while the magnitudes of $T_{C1}$ and $T_{C2}$ in (b) are almost identical. Correspondingly, one can see that $\tau_{\text{eff}}$ in (a) diverges up to almost 10 ps around $T_{C2}$ while $\tau_{\text{eff}}$ in (b) reaches only 1 ps. These results clearly indicate that $S_2 \gg S_1$ is achieved in (a) while $S_1 \approx S_1$ in (b). Therefore $\Delta R(t)$ is approximately described by $S_1(T)e^{-\dot{\tau}_1(t)}$ at $\omega_{pr}=1.56$ eV. Note that at $\omega_{pr}=1.07$ eV, where $\Delta R(t) \cong S_1(T)e^{-\dot{\tau}_1(t)} + S_2(T)e^{-\dot{\tau}_2(t)}$, the divergence of $\tau_{\text{eff}}$ can be assumed to be due to the relatively large $S_1(T)$ contribution around $T_{C2}$. In other words, the strong divergence of $\tau_{\text{eff}}$ observed at $\omega_{pr}=1.56$ eV reflects $\tau_1(T)$ more precisely. Also note that the degree of divergence $\tau_1(T= T_{C1})$ is much smaller than that of $\tau_2(T=T_{C2})$, which will be discussed later.

To clarify the energy dependence of the selectivity on the SP dynamics in the $T_{C2}$ phase, Fig. 3(a) shows a series of transient $\Delta R$ obtained at a temperature just below $T_{C2}$, where $\tau_2$ exhibits an extended decay. Because we used an OPO as the incident light source, the pump energy is also changed by tuning the probe energy, which may modify the number of photoexcited SPs. To compensate for this effect, we compare $\tau_{\text{eff}}$ rather than $S_{\text{eff}}$ in Fig. 3(b). This figure indicates that the longer $\tau_{\text{eff}}$ becomes, the more $S_{\text{eff}}$ includes the contribution of the $T_{C2}$ phase. The spectrum shows a slowly varying distribution, suggesting the energy dependent enhancement of the contribution of the $T_{C2}$ phase. The maximum value of $\tau_{\text{eff}}$ is located at $\omega_{pr}=1.56$ eV. Thus we conclude that selective
probing of the $T_{C2}$ phase is achieved in this energy region.

Let us discuss the degree of divergence of $\tau_n$ in each CDW phase. On the basis of the theoretical framework developed by Kabanov et al.,11 the SP dynamics reflects the Bardeen-Cooper-Schrieffer (BCS)-type temperature dependence of the gap $|\Delta_n(T)|$. In the simplest approximation, the model yields a $\tau_n$ value that is proportional to the inverse of $\Delta_n(T)$. In this case, $\tau_n$ diverges to infinity when the temperature is increased close to $T_{Cn}$. In a real system, $\tau_n$ is saturated by the thermal-phase fluctuations. The effects of thermal fluctuation are marginal since here we studied the SP relaxation on a picosecond time scale. The difference in the maximum value of $\tau_n$ in the $T_{C1}$ and $T_{C2}$ phases can therefore be explained by the difference in phase fluctuations at $T_{Cn}$. In this case, the suppressed (enhanced) divergence of $\tau_1$ ($\tau_2$) indicates the suppressed (enhanced) phase coherence for three-dimensional (3D) CDW ordering, which is consistent with the difference in the modulation wave vectors, namely $Q_1=(0, 0.241, 0)$ for the $T_{C1}$ phase and $Q_2=(0.5, 0.260, 0.5)$ for the $T_{C2}$ phase. We note that the maximum of $\tau_1$ is comparable with the results obtained for the other CDW compounds: $\tau_{eff}$ in 1T-TaSe$_2$ and 2H-TaSe$_2$ reaches only 1 ps ($T_c=170$ K) and 0.4 ps ($T_c=140$ K), respectively, and K$_{0.3}$MoO$_3$ reaches almost 1 ps at $T_c=180$ K. Therefore it is possible to consider that the divergence of $\tau_2$ is extremely large. One possibility for such an enhancement of the maximum $\tau_2$ is the phase locking of the coexisting CDWs below $T_{C2}$, which has been predicted for the zero-modulation wave vectors in the superposition of two CDWs.12

Although these explanations are still speculative, we emphasize that the considerable difference in $\tau_n$ indicates a strong sensitivity to the phase transition properties and thus provides a useful parameter for investigating the dynamics and interactions of coexisting CDWs. In this sense, our results connect with recent reports on the sliding properties of NbSe$_3$ CDWs.13,14 Indeed, a motional 1D coherence accompanied by a loss of transverse correlation above $T_{C2}$ and a sliding induced decoupling of coexisting CDWs have been observed, both of which are consistent with our results.

We note a difference in the $\tau_{eff}$ saturation behavior was also found in a comparison of topological NbSe$_3$ crystals, where the divergence of $\tau_2$ in the ring NbSe$_3$ is clearly suppressed even in the vicinity of $T_{C2}$. In accordance with the power-law decay of the transient $\Delta R$ around $T_{C2}$, we have concluded that the results can be explained in terms of the phase fluctuation enhanced by closed-loop topology.3,15,16

We now investigate the selectivity of the $T_{C2}$ phase in the lowest temperature regime ($T \ll T_{C2}$). For this purpose, we consider the coherent oscillation part of $\Delta R$, in which one of the CDW oscillations (amplitude mode, AM) clearly exhibits energy dependence. Since coherent AM oscillations are directly connected with optical transitions involving the CDW gap states, the excitation energy dependence of the AM allows us to evaluate the corresponding CDW phase separately.

Figure 4(a) shows a series of such oscillating data at various probe energies after subtraction of the exponential decay. For clarity, the corresponding fast Fourier transform (FFT) spectra are shown in Fig. 4(b). Two characteristic peaks are dominant in these spectra. The lower peak at 1.13 THz shows a softening behavior with increasing temperature and completely disappears above $T_{C2}$, indicating AM associated with the $T_{C2}$ phase. The higher peak at 1.22 THz is identified as a coherent phonon because of its less pronounced temperature dependence.

It is important to note that at $T \ll T_{Cn}$, the SP dynamics from each CDW phase exhibit almost the same decay time ($\tau_1 \approx \tau_2$) mainly due to the completion of the CDW gap (see the bottom two curves in Fig. 1). Consequently, we cannot distinguish the contributions of the different $T_{Cn}$ phases in the exponential part of $\Delta R$. On the other hand, the AM os-
oscillation shows a significant energy dependence as shown in Fig. 4(b). Therefore we consider the AM oscillation amplitude to be an alternative parameter with which to evaluate the contribution of the $T_{C2}$ phase at $T \ll T_{C2}$.

Figure 4(c) shows plots of the FFT amplitude of AM as a function of probe energy. In this case, the more intense the AM is, the more the signal includes the contribution of the $T_{C2}$ phase. As seen in Fig. 4(b), the spectrum exhibits a distribution, where AM becomes dominant by tuning the probe energy to the lower limit of the laser, suggesting an energy dependent contribution of the $T_{C2}$ phase.

We now discuss the mechanism by which the selective excitation for one of two CDW phases was achieved in the transient reflectivity measurements. Previous optical experiments on the NbSe$_3$ whisker have shown that there are several small but distinct resonances originating from interband transitions above the plasma edge. Theoretically, the band electronic structure based on tight-binding calculations also predicts the number of higher excited states with flat dispersions parallel to the CDW band where we can expect a large transition probability for the resonant optical excitation. As mentioned by Demsar et al., the probe signal for above-gap excitation reflects the interband transitions from the band edge to an appropriate higher excited band. Therefore, the excitation energy dependence in our present results suggests a transition to the excited state with a large transition probability that increases the contribution of the corresponding CDW phase in the signal [see Fig. 5(a)].

We comment on the difference between the resonance energies in the two experiments at different temperatures. Although the exact spectrum peak is obscured by the laser limit in Fig. 4(b), the energy corresponding to the efficient CDW oscillation shifts toward a lower energy than the SP decay spectrum [Fig. 3(b)]. This shift can be explained by the transition schemes summarized in Figs. 5(a) and 5(b). The SP decay was evaluated just below $T_{C2}$, at which we observed the opening of $2\Delta_{C2}$. On the other hand, we obtained the CDW oscillation at the lowest temperature where the complete gap exists. When we consider the transition from the CDW band edge as the dominant process for the probe signal, the resonance energy should be shifted by the amount of gap energy. $2\Delta_{C2}$ is approximately 65–100 meV, which is consistent with the difference in the resonance. Note that the probe signal also reflects the upper edge of the occupied electron band. Figure 4(c) also shows a small peak around 1.56 eV, which may account for this resonance condition.

In summary, we have performed energy-selective pump-probe reflectivity measurements on whisker NbSe$_3$ with a micron size spatial resolution. The SP dynamics clearly shows two anomalies in the temperature dependence, which reflects the CDW phase transitions. The magnitudes of the anomalies in the SP decay constant and the SP excitation amplitude are changed by changing the excitation energies, indicating selective probing of the CDW phases. We have also demonstrated that the AM oscillation can be employed as another selective probe at the lowest temperature. The results allow us to identify the transient signals in each CDW phase and to clarify the differences in the phase transition properties. In addition, the temperature dependence of selectivity could help to verify the selective excitation mechanism in terms of the optical transitions connected with the CDW gap states. These selective excitation experiments have great potential as regards undertaking precision measurements of nonequilibrium dynamics characterized by the coexistence of multiple phases.

ACKNOWLEDGMENTS

The authors sincerely thank P. Monceau, M. Oda, R. Morita, and K. Inagaki for fruitful discussions. This work was partially supported by a Grant-in-Aid for the 21st Century COE program “Topological Science and Technology.”

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