Anharmonic Decay of Coherent Optical Phonons in Antimony

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Anharmonic decay of coherent optical phonons in semimetal Sb has been investigated by using a femtosecond pump-probe technique. The coherent A1g mode is observed in time domain in a wide temperature range of 7 - 290 K. The decay rate (the inverse of the dephasing time) systematically increases as the lattice temperature increases, which is well explained by anharmonic phonon-phonon coupling, causing decay of the optical phonon into two acoustic phonon modes. The frequency of the A1g mode decreases with the temperature, which is interpreted to the results of both thermal expansion and anharmonic phonon-phonon coupling. The temperature dependence of the amplitude of the coherent A1g mode exhibits a decrease with the lattice temperature, which is well reproduced by considering the peaked intensity of spontaneous Raman scattering assuming a Lorentzian line shape with the linewidth controlled by the anharmonic decay, and this model can be applicable to other metallic system, like Zn.

I. INTRODUCTION

In recent years, coherent phonons have been studied by using a femtosecond pump-probe technique in various material systems, including semiconductors,1-2 metals,2,3 superconductors,4-6 organic conductors,7 and phase change materials.8,9 The coherent phonons can be generated by ultrashort laser pulses with a high degree of temporal coherence. The nature of the coherent phonons has been extensively studied in semimetals10,11 and semiconductors,12 where the main focus was the generation mechanism. Because the coherent phonons are lattice vibrations having the same phase in time, dephasing of the coherent phonon is directly monitored by optical pump-probe measurements.

The studies on the dephasing process are especially important for understanding the nature of in-phase coherent phonons. The dephasing process of coherent phonons excited by picosecond pulses has been examined using time-resolved CARS (coherent anti-Stokes Raman scattering).13,14 The phonon decay rate can be generally described as the sum of the anharmonic decay rate and the pure dephasing rate.15 The main channels for the relaxation of incoherent optical phonons in semiconductors are thought to be the dephasing originating from the phonon-phonon interaction caused by anharmonicity of the lattice potential. In this channel, the excited optical phonons decay into acoustic phonons.16,17 A direct comparison has been made between the results obtained by time-resolved CARS and Raman scattering spectroscopies.14,15 Anharmonic phonon decay of coherent phonons was first confirmed by Hase et al. in Bi by measuring the temperature dependence of the dephasing time or decay rate (the inverse of the dephasing time).18 It has been also shown that the dephasing of the coherent optical phonon is very sensitive to the density of lattice defect (vacancy).19,20 However, systematic study of the dephasing of coherent optical phonons in semimetals other than Bi excited by femtosecond laser pulses is still few.

In this paper, dephasing process of coherent optical phonons in semimetal antimony (Sb) is studied by femtosecond pump-probe reflectivity measurements in a temperature range from 7 to 290 K. The amplitude, the dephasing time, and the frequency of the coherent optical phonons have precisely been measured. The decay rate and frequency show the systematic change as a function of the lattice temperature, similar to those in Bi, and reproduced by the model based on anharmonic lattice effects. The amplitude of the coherent phonon also exhibits systematic decrease with increasing the lattice temperature, which is well fit by the model based on the phonon occupation number proposed by Misochko et al.21

II. EXPERIMENTAL TECHNIQUE

The samples used in this study was a single crystal of Sb with cut and polished with the (0001) surface. The femtosecond pump-probe measurements were carried out in a temperature range from ≈7 to 290 K using a closed-cycle cryostat. The light source used was a mode locked Ti:sapphire laser with a central wavelength of 800 nm, providing ≈20 fs pulses at the repetition rate of 87 MHz. The pump and probe beams were polarized orthogonal each other to avoid the scattered pump beam. Both pump- and probe-beams were focused onto a diameter of ≈100 µm on the sample. The average power of the pump and probe beams were fixed at 120 mW and 5 mW, respectively, from which we estimated the pump fluence to be 18.4 µJ/cm² at 120 mW. By changing the optical path length of the probe beam, the reflectivity
change (∆R/R) was recorded as a function of the delay time. This isotropic reflectivity measurement enable us to dominantly detect the fully symmetric A$_{1g}$ mode in Sb, while the non-symmetric E$_g$ mode is generally masked. Thus the determination of the phonon parameters, i.e., the amplitude, the dephasing time, and the frequency is accurate in the present study because a single damped harmonic oscillator model can fit the coherent phonon signal with smallest number of parameters.

\[
\frac{\Delta R(t)}{R} = A \exp\left(-\frac{t}{\tau}\right) \cos(\omega_{A_{1g}}t + \phi) + B \left[ \exp\left(-\frac{t}{\tau_1}\right) - \exp\left(-\frac{t}{\tau_2}\right) \right],
\]

where $A$ is the amplitude, $\omega_{A_{1g}}$ is the frequency, $\tau$ is the dephasing time, and $\phi$ is the initial phase of the coherent A$_{1g}$ mode. The second term arises from the photoexcited carriers. Here $B$ is the amplitude, and $\tau_1$ and $\tau_2$ are the relaxation time, and the rising time of the electric component, respectively. The fit results in Fig. 1 are satisfactory good and thus we obtain the phonon parameters. The initial phase ($\phi$) obtained is nearly zero, meaning the coherent oscillation is cosine-like. This phase value matches with the previous results in Sb that the generation mechanism of the coherent A$_{1g}$ mode is governed by the displacive excitation of coherent phonon (DECP) model or Raman scattering finite lifetime (RSFL) model, the latter of which took both the finite lifetime of the coupled charge carrier density (a nature of DECP) and stimulated Raman scattering into account. The amplitude of the electric component, $B$, exhibits increase by $\sim 15 \%$ as the temperature is lowered as seen in Fig. 1, suggesting the photoexcited carrier density increases with decreasing the lattice temperature. We will discuss the temperature dependence of the parameter $B$ together with the amplitude of the coherent A$_{1g}$ mode ($A$) in the later sections. The values of the relaxation time, $\tau_1$ and $\tau_2$, do not systematically change with the temperature, and therefore we would not discuss the time constants of the electric component in detail in the present paper. Hereafter, the temperature dependences of the decay rate, the frequency, and the amplitude of the coherent optical phonons are mainly focused and explored.

In Fig. 2 the decay rate (the inverse of the dephasing time) of the A$_{1g}$ mode at different temperatures obtained by the time-domain measurement are plotted. The frequency of the coherent A$_{1g}$ mode is also plotted in the inset. The frequency of the A$_{1g}$ mode shifts from 4.67 THz to 4.52 THz when the temperature increases from 7 to 290 K. The decay rates of the A$_{1g}$ mode monotonically increases as the lattice temperature rises, which is comparable to the results in Bi films. This behavior is attempt to fit by an anharmonic decay model, in which the optical phonon decays into two acoustic phonons with half the frequency of the optical mode ($\omega_{A_{1g}}/2$) and with opposite wavevectors.

\[
\Gamma_{A_{1g}} = \Gamma_0 + \Gamma(1 + n_{A_{1g}}/2),
\]
where $\Gamma_0$ is a background contribution due to impurity and defect scattering, $\Gamma$ is the anharmonic coefficient, $n_{A_{1g}/2} = [\exp(h\omega_{A_{1g}}/2k_B T) - 1]^{-1}$ is the Bose-Einstein factor, and $k_B$ is the Boltzmann constant. The fitting of the time-domain data to Eq. (2) is shown in Fig. 2, where we obtain $\Gamma_0 = 0.1 \text{ ps}^{-1}$ and $\Gamma = 0.06 \text{ ps}^{-1}$ for Sb. Thus, the temperature dependence of the decay rate of the coherent $A_{1g}$ mode is well described by Eq. (2), indicating that the dephasing of the coherent $A_{1g}$ mode is governed by the anharmonic phonon-phonon coupling. In the same manner, the frequency softening of the $A_{1g}$ mode is expressed by taking into account contributions from the thermal expansion and anharmonic coupling, both of which are related to the cubic anharmonic term.

$$\omega_{A_{1g}} = \omega_0 + \Delta_0(T) + \Omega(1 + 2n_{A_{1g}/2}), \quad (3)$$

where $\omega_0$ is the harmonic frequency at the lowest temperature limit, $\Omega$ is an anharmonic constant, and $\Delta_0(T)$ represents the frequency shift due to the thermal expansion of the lattice, which is given by

$$\Delta_0(T) = -\omega_0 \gamma \int_0^T \alpha_{//}(T') + 2\alpha_{\perp}(T') \text{d}T', \quad (4)$$

where $\gamma$ is the Grüneisen parameter, $\alpha_{//}(T')$ and $\alpha_{\perp}(T')$ are the linear thermal expansion coefficients along directions parallel and perpendicular to the c-axis, respectively. In order to calculate $\Delta_0(T)$, we took the experimental linear thermal expansion coefficients and the Grüneisen parameter ($\gamma = 1$) measured by White and thus model fitting based on Eqs. (3) and (4) was carried out. The fitting result of the time-domain data to Eqs. (3) and (4) is shown in the Fig. 2 inset, where we obtain $\omega_0 = 4.75 \text{ THz}$ and $\Omega = -0.087 \text{ THz}$, the latter of which is comparable to the result ($\Omega = -0.056 \text{ THz}$) in Bi films.

It should be mentioned here that the contribution from the thermal expansion to the frequency softening becomes effective at $T \geq 100 \text{ K}$ because the linear thermal expansion coefficients are rather small at $T \leq 100 \text{ K}$, while they become larger and nearly constant at $T \geq 100 \text{ K}$. As shown in Fig. 3, the amplitude of the coherent $A_{1g}$ mode increases with decreasing the temperature. The ratio of the amplitude of the coherent $A_{1g}$ mode obtained at 7 K to that obtained at 290 K is $\approx 1.3$. A similar temperature dependence of the coherent phonon amplitude was observed for the $E_g$ mode in a single crystal Sb by Ishioka et al., while they observed rather flat behavior for the $A_{1g}$ mode. The difference in the amplitude of the coherent $A_{1g}$ mode between the two measurements might partly come from the different temperature dependence of the electric component, since the amplitude of the $A_{1g}$ mode is proportional to the photoexcited carrier density, $A = \kappa B$, where $\kappa$ is an electron-phonon coupling constant, under the DECP model. In fact, the one observed the increase of the electronic transient between 100 and 200 K by the isotropic reflectivity measurements while we observed rather decrease of the electric component with the temperature by the isotropic reflectivity measurements (see Fig. 3), as has been observed also in simple metals. We fitted the temperature data of the phonon amplitude in Fig. 3 to the expected temperature dependence of the peaked intensity of spontaneous Raman scattering assuming a Lorentzian line shape with the linewidth controlled by anharmonic decay. By using the well-known property of a Lorentzian, the temperature dependence of peak intensity of the Raman-active $A_{1g}$ mode is proportional...
in terms of the quasiparticle density. Zn, in which the temperature dependence was examined coherent optical phonon was observed in the simple metal the materials. A strong temperature dependence of the amplitude of the coherent optical phonon with Eq. (5) found in Fig. 3 suggests the RSFL mechanism would predominate the generation of the coherent $A_{1g}$ mode over the DECP mechanism. This interpretation is further supported by the fact that the temperature dependence of the electronic amplitude ($B$) exhibits a significantly different nature from that of the coherent $A_{1g}$ mode ($A$) as shown in Fig. 3, and consequently the DECP model ($A = \kappa B$) cannot solely account for the observed results.

IV. DISCUSSION

It is to be noted that in case of coherent phonons of Bismuth, the $E_g$ mode was also observed with $\approx 1/10$ Fourier transformed (FT) spectral intensity at low temperatures even with the isotropic detection scheme. In Sb, there seems to be no $E_g$ component at 7 K in Fig. 1, which was also confirmed by the FT spectra (not shown). A possible reason why the difference in the appearance of the $E_g$ mode between these two different atoms was found is polarization dependence of Raman tensor for the $E_g$ mode. In order to observe the $E_g$ mode by the isotropic detection one would need to use highly-oriented single crystal sample and to precisely rotate the pump-polarization in $x - y$ plane to match the appropriate crystal axis, although the study of pump-polarization dependence of the $E_g$ mode using such a highly-oriented single crystal is beyond the scope of this paper.

Hereafter we rather focus on the temperature dependence of the amplitude of the coherent phonon, referring other metallic system, since it will give a new insight into the physics of the coherent phonons in a wide range of the materials. A strong temperature dependence of the coherent optical phonon was observed in the simple metal $Zn$, in which the temperature dependence was examined in terms of the quasiparticle density. Fig. 4 shows the comparison of the fitting models for the temperature dependence of the amplitude of the coherent optical phonon observed in $Zn$.

It is found that the current model, Eq. (5) in the present study, rather well describes the overall data in Fig. 4, although the previous model based on the density of the quasiparticles also could fit the data at $T \geq 80$ K while some deviation was found at lower temperature than 80 K. Consequently we conclude that the overall behavior of the coherent phonon amplitude as a function of the lattice temperature is well modeled by Eq. (5) in a wide range of metallic systems (Bi, Sb, and Zn) or in the wide range of the Debye temperature, $\Theta_D \approx 120$ K for Bi, $\Theta_D \approx 210$ K for Sb and $\Theta_D \approx 330$ K for Zn. Note that a plausible reason why the deviation was observed at $T \leq 80$ K for Zn in Fig. 4 between the model in Ref. [4] and the data is a classical nature of the simple model in Ref. [4], or $A \propto T'_c - T_c$, where $T'_c$ and $T_c$ are the final and initial electron temperature, respectively. The current model, Eq. (5), is rather based on quantum effect described by the Bose-Einstein factor.

V. CONCLUSION

In conclusion, temperature dependence of the dynamics of coherent optical phonons in antimony single crystal has been studied by a femtosecond pump-probe technique at various lattice temperatures. The agreement of the decay rate and the frequency of the coherent $A_{1g}$ mode with the anharmonic model indicates that the dephasing of coherent phonons in antimony is dominated by anharmonic decay (energy relaxation), and that the frequency softening is due to cubic term of the anharmonicity, in which thermal expansion and anharmonic phonon-phonon coupling play important roles. The coherent phonon amplitude of the $A_{1g}$ mode shows the decrease with the lattice temperature, which is well modeled by the use of the peaked intensity of spontaneous Raman scattering, and which also reproduces the amplitude of the coherent $E_g$ mode in Zn satisfactory. Thus, in a wide range of metallic systems anharmonic lattice effect dominates the temperature dependence of the decay rate, the frequency,
and the amplitude of the photo-excited coherent optical phonons.

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