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Development of an optical time-resolved measurement system under high-pressure and low-temperature with a piston-cylinder pressure cell

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To perform the femtosecond pump-probe spectroscopy under high pressure and low temperature, we constructed a measurement system with a piston cylinder type pressure cell installing an optical fiber bundle. The applied pressure was achieved to 6 kbar and the cell was cooled down to 15 K. Several demonstrations revealed that broadening and change of polarization of pulse (duration of ~120 fs) owing to the dispersions in the fiber bundle are much small indicating that those have little influence on the measurement of carrier relaxation dynamics. In the measurements of κ-(BEDT-TTF)2Cu(NCS)2 under 1.3 kbar at 43 K, we have successfully detected the polarization anisotropy of the carrier relaxation dynamics and estimated the decay time in the same way as the normal measurement. © 2016 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4945303]

I. INTRODUCTION

Femtosecond time-resolved optical spectroscopy is a powerful technique to measure relaxation of photo-excited quasi-particles (QPs) resolved in the time domain. This measurement has made a significant contribution to researches on the electronic phases, systematic time-resolved measurements with varying pressure have not been performed so far. This is because the optical measurements under pressure tends to require large effort to prepare and handle a pressure-tunable equipment. Under high pressure, various optical measurements have been performed by using a diamond anvil cell (DAC) because of optical transparency of diamond anvils.22,23 However, DAC tends to be expensive and difficult to handle because of small sample space (less than $10^{-3}$ mm$^3$). Another equipment to achieve high pressure is a piston-cylinder type pressure cell.24 The piston-cylinder pressure cell has been widely used for electrical resistivity, magnetization, and thermal measurements and has larger sample space ($10^2$ mm$^3$) than that of DAC. To use the piston-cylinder cell for optical measurements, an optical window is needed to set in it. A sapphire and optical fiber bundle was used as a window in the photoluminescence measurements25 and the infrared investigations under high pressure.26 On the other hand, an optical fiber is also suitable for the window since it enables light to propagate with little attenuation. Indeed, by building a thin fiber optics directly into the piston-cylinder cell, fluorescence of ruby has been measured.27 However, by use of the thin fiber, optical response from sample becomes often much small because the area irradiated by the beam is limited by the diameter of the fiber. An optical fiber bundle which consists of a large number of the thin fibers can make it possible to carry out the optical measurements under pressure. Since the overall diameter of the bundle is sufficiently larger than sample, the response signal comes from the area limited by the spot size of the beam, which is in the same way as the measurement under ambient pressure. In this paper, we report the construction of femtosecond pump-probe measurement system with a piston-cylinder type pressure cell and a short optical fiber bundle. Through the several demonstrations, the following points were evaluated: attainable pressure and temperature, effect of the dispersions on an optical pulse, detection characteristics of transient change of reflectivity, and polarization in the organic conductor κ-(BEDT-TTF)2Cu(NCS)2 (hereafter κ-NCS) under multiple conditions (43 K and 1.3 kbar).

II. PRESSURE AND TEMPERATURE CHARACTERISTICS OF PRESSURE CELL

To realize the time-resolved measurement under high pressure and low temperature, the piston-cylinder type hybrid pressure cell24 and optical fiber bundle were prepared. Our pressure cell was essentially the same with that in the previous report.24 Two-layered cylinder consists of a nonmagnetic Ni-Cr-Al and Cu-Be alloys. Plug and clump bolts are Cu-Be. Piston and piston-back-up are made of a nonmagnetic tungsten carbide (WC). Figure 1(a) shows a schematic of a cross-sectional view of the sample space and microscopic image of the end surface of the fiber bundle. The plug has a hole whose diameter is 3.4 mm to set the optical fiber bundle in it. The fiber bundle consists of about 50 thousands narrow fibers (12 μm of diameter of the individual cores), whose overall diameter is 3.2 mm and length is 25.4 mm (the specifying part number:♯53-839, Edmund Optics Inc.). Refractive index of core and cladding is 1.58 and 1.48, respectively. The bundle is set into the hole of plug and fixed tightly by stycast (2850FT). The stycast was poured into the space between the plug and fiber bundle to fill the

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space as evenly as possible after the removal of dissolved gas under vacuum. Sample and ruby particles are put on the end surface of the fiber in the sample space. The plug was joined to Teflon capsule with the pressure medium (Daphne 7373) and mounted in the pressure cell as shown in Fig. 1(b). In this setup, a vent hole was not constructed at the space under the clamp bolt to release pressure in it. For safety, the hole would be better to be formed. Figure 1(c) shows cross-sectional view of the pressure cell, which is connected to heat exchanger of the cryostat with the brass joint and cooled with a helium flow.

Applied pressure was calibrated by the ruby fluorescence method. For excitation, green laser light (532 nm) was generated from the diode pumped solid state laser. The beam reflected by a dichroic mirror (DM) was focused by an objective lens and irradiated to ruby particles presented on the end of the bundle. The fluorescence comes through DM and into a charge coupled device spectrometer (Ocean Optics, HR4000) via an optical fiber.

The inset of Fig. 2 shows spectra of the ruby fluorescence for 0 and 2 ton. Both the $R_1$ and $R_2$ peaks are shifted
The plot of pressure versus the load. The solid line indicates the linear fit. The inset shows ruby fluorescence spectra.

FIG. 2. The plot of pressure versus the load. The solid line indicates the linear fit. The inset shows ruby fluorescence spectra.

clearly to the long-wavelength side with increasing load. The applied pressure is estimated by the R₁ shift $\Delta R_1$ with the simple expression $P = 2.740 (\text{kbar}/\AA) \times \Delta R_1$. The estimated pressure values are plotted as a function of the load in Fig. 2. The pressure was linearly applied up to 6 kbar and suddenly reduced. This reduction was caused by disaggregation between the bundle and plug. An optimal size of the space between the bundle and plug remains unclear although it can be important to get high pressure. If the space is too large, the bundle can be fixed tilting with respect to the plug. This may cause breaking of the bundle. If the space is too small, amount of the stycast will reduce, leading to disaggregation between them. By improving condition of a diameter of the hole and adhesion between the bundle and plug, pressure is expected to increase more than 6 kbar.

Temperatures were measured by two thermometers which were attached to the end of the cell and exchanger as shown in Fig. 2(c). The cell was cooled down to $T_{\text{cell}} \sim 15 \text{ K}$ and $T_{\text{he}} \sim 4.2 \text{ K}$, respectively, without any trouble.

III. EXPERIMENTAL SETUP OF TIME-RESOLVED SPECTROSCOPY

In the time-resolved spectroscopy, two optical pulses, namely, pump and probe pulses, are used. First pump pulse excites carriers in a solid to non-equilibrium high-energy states. The excited QPs relax immediately to states near the Fermi surface by electron-electron and electron-phonon states. The excited QPs relax immediately to states near excites carriers in a solid to non-equilibrium high-energy states, namely, pump and probe pulses, are used. First pump pulse.

IV. PROPERTIES OF OPTICAL PULSE PASSING THROUGH A FIBER BUNDLE

When an ultrashort optical pulse propagates in a fiber optics, broadening of the pulse, which will lead to reduction of time resolution on the measurements, is caused by the dispersions. To investigate the broadening effect on the transient measurement, we measured transient change of reflectivity $\Delta R/R$ by using the fiber bundle at room temperature. Figure 3(a) shows $\Delta R/R$ normalized by the maximum value $\Delta R_{\text{max}}$ of $\kappa$-NCS obtained from the measurement with the fiber bundle and the normal measurement. Decay times $\tau$ were estimated by fitting the data with a single exponential function $\exp(-t/\tau)$. The fits yield $t_1 \sim 0.88 \text{ ps}$ for the measurement with the bundle which is comparable with $\tau_{\text{w}} \sim 0.89 \text{ ps}$ for that without the bundle. To check the influence of pulse broadening more precisely, we also compare the initial rise of $\Delta R/R$. For simplicity, the rising times $t_1$ and $\tau_{\text{w}}$ were defined as a time between $1/2 \Delta R_{\text{max}}$ and $\Delta R_{\text{max}}$ as shown in Fig. 3(a). As a result, $t_1$ was $\sim 0.25 \text{ ps}$, which agrees with the $t_0$ value $\sim 0.23 \text{ ps}$ within experimental error. These results indicate that the pulse broadening has almost no effect on the relaxation measurements.

Here, we roughly estimated the broadening of the pulse caused by the material dispersion. By assuming Gaussian pulse and using an input pulse width of 120 fs and the group velocity dispersion of 36 fs²/μm on silica glass at 800 nm, the width of pulse passed through fiber was estimated as $\sim 122 \text{ fs}$. Such a small broadening will be unmeasurable, which was consistent with the above results. We have to note that the spatial mode dispersion of the fiber bundle would also broaden the pulse width. In the case of much shorter femtosecond pulse, the broadening could be larger than the theoretical estimation.

To realize the Raman-like analysis on the QP dynamics, we employ the polarization-resolved measurement. Since polarization of the optical pulse can be modulated by the polarization mode dispersion, the degree of polarization (DOP) of the pulse propagated through the fiber was evaluated.
FIG. 3. (a) $\Delta R/R$ transients obtained from the measurement with the fiber optics for $F \sim 1100 \mu J/cm^2$ and the normal measurement under ambient pressure for $F = 93 \mu J/cm^2$ at a room temperature. The dashed lines indicate the results fitted by a single exponential function. The data are normalized by the maximum values of $\Delta R$ and shifted for clarity. (b) Degree of polarization (DOP) as a function of length of the optical fiber bundle. The value at 0 mm corresponds to DOP of ambient light.

using Stokes parameters, which were obtained by combining a quarter wave plate and a polarizer. Theoretically, DOP has a value in the range from 0 to 1. DOP = 0 and 1 mean that light is unpolarized and completely polarized, respectively. Figure 3(b) shows the fiber-length dependence of DOP. The DOP value of ambient pulse presented at 0 mm is $\sim 0.978$, indicating that the pulse is mostly polarized. For the pulse propagated through the fiber bundle, the DOP values are nearly equal to those of the ambient pulse at any fiber-lengths. The results indicate that the polarization of pulse undergoes little change even with the use of the fiber bundle.

On the measurement using the fiber optics, power loss of pulses due to the non-linear effects caused by high-intensity light may be concerned. Since the non-linear effects depend on length of the fiber optics, we measured power $p_b$ of pulse passing through the bundle and compare it with $p$ which was measured before entering the bundle with four kind of bundles whose lengths are 25.4, 76.2, 152, and 305 mm. As a result,

FIG. 4. ((a) and (b)) Transient changes of reflectivity $\Delta R/R$ for $\theta = 0^\circ$ and $90^\circ$ and intensity plots of $\Delta R/R$ as a function of probe-polarization-angle under ambient pressure at 50 K for $F = 121 \mu J/cm^2$ and 1.3 kbar at 43 K for $F \sim 1300 \mu J/cm^2$, respectively. The dashed lines indicate the results fitted by a single exponential function. ((c) and (d)) Polar plots of the maximum values of $\Delta R/R$ under ambient pressure at 50 K and 1.3 kbar at 43 K, respectively. The solid lines indicate the results fitted by a double-angle function.
for all the bundles, the ratios between them were $\frac{p_b}{p} \sim 0.4$, which corresponds to 35%–45% efficiency of transmission in the catalog. This means that the power loss was caused by reflection at an end face of the bundle. It is consistent with the fact that the laser power used in our experiments is less than or comparable to ~0.5 mW, which will be too small to cause the non-linear effects.

V. DEMONSTRATION UNDER HIGH PRESSURE AND LOW TEMPERATURE

Figure 4(a) shows transient change of reflectivity $\Delta R/R$ for probe polarization $\theta = 0^\circ$ and $90^\circ$ and intensity plots of $\Delta R/R$ as a function of $\theta$ under ambient pressure at 50 K. To obtain clear signal, the pressure measurements were carried out with higher pump fluence $F \sim 1300 \mu J/cm^2$ as compared to 121 $\mu J/cm^2$ in the normal measurements. We found that $\Delta R/R$ clearly depended on the probe polarization angle. Figure 4(c) shows a polar plot of the maximum values of $\Delta R/R$. The signal was enhanced for $\theta \sim 0^\circ$ and $180^\circ$, which correspond to the directions along the $b$ axis. This polarization anisotropy can be attributed to spatial symmetry breaking due to the glass-like structural transition below 70 K.35 Figures 4(b) and 4(d) show the polarization angular dependences of $\Delta R/R$ under 1.3 kbar at $T_{coh} = 43$ K, which were quite similar to those in the normal measurement. The slight difference of direction of the anisotropy between them can be due to the misalignment between the crystal axis and polarization of pulse. The conservation of polarization can be an important consequence for optical measurements in a pressure environment. Generally, birefringence of the window can be caused by the stress of the applied pressure, leading to non-conservation of polarization. Therefore, these results suggest that polarization is conserved even in our pressure environment and the polarization properties of the QP relaxation dynamics are measurable in the same way as the normal measurement.

Although the polarization anisotropies are comparable, we found the difference between the decay times obtained from the pressure and normal measurements. Under 1.3 kbar, the decay times are ~1.58 and 1.03 ps for $\theta = 0^\circ$ and $90^\circ$, respectively, while under ambient pressure, $\tau \sim 0.95$ and 0.25 ps, respectively. The difference of the pump fluence between the measurements does not influence the decay time above the saturation condition ($F_{th} \sim 40 \mu J/cm^2$). Generally, application of pressure modulates the $e$-$ph$ interaction. Since the carrier relaxation rate $1/\tau$ is proportional to the strength of the $e$-$ph$ interaction, the applied pressure may lead to modulation of the decay time of the QP relaxation.

In the demonstrations, the pressure was calibrated at room temperature. In the previous report,36 the applied pressure is reduced for the Daphne 7373 pressure medium as temperature decreases, meaning that the pressure value at low temperatures can become smaller than 1.3 kbar. However, it is difficult to determine pressure accurately at low temperature by the ruby fluorescence method since the $R_1$ and $R_2$ lines of the ruby fluorescence are shifted depending on temperature.37 One of the solutions is that temperature dependence of the $R_1$ shift $\Delta R_1$ is investigated previously in the Daphne 7373 under 0 GPa. Because amount of $\Delta R_1$ is almost unchanged even at low temperatures,38 a pressure value may be calibrated from the pre-investigations.

VI. SUMMARY

We developed the measurement technique of femtosecond optical pump-probe spectroscopy under high-pressure and low-temperature by using the piston-cylinder type pressure cell and optical fiber bundle. The series of demonstrations revealed that our measurement system is available for the pump-probe measurement and provides reliable results in terms of the decay time and polarizations of the relaxation dynamics. The technique will be applicable to other optical measurements and become a key part of the time-resolved spectroscopy under multiple conditions including high magnetic field, opening a new research field.

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