Broadband Terahertz Time-Domain Spectroscopy and Low-Frequency Raman Scattering of Propylene Glycol

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Abstract. Two different THz-TDS systems were used for the accurate study of broadband terahertz dynamics between 0.1 and 5.0 THz. These systems enabled to observe the whole spectral structure of the broad peak at 1.5 THz in propylene glycol. At high-frequency side of the broad peak in the dielectric spectrum, the small shoulder peak at 3.6 THz was also observed. It is assigned to the inter-molecular vibration, which is supported by the precedent report and the Gaussian calculation. In contrast, the low-frequency side of the broad peak includes the contribution of the relaxation processes which exist in the MHz and GHz region. Furthermore, we compared the dielectric spectrum with the Raman spectrum and concluded that the mode at 3.6 THz is IR active while Raman inactive.

Introduction

The THz dynamics have been intently investigated for various kinds of materials such as polymers [1], pharmaceuticals [2], ferroelectrics [3], and so on. Furthermore, there has been the significant improvement of the methods for measuring not only the crystalline state, but also the amorphous and liquid states of materials. Most of the glassy and liquid states materials have a broad peak in THz region, and thus it should be discussed with a broadband THz spectrum; however, unfortunately for the liquid states, the investigation in the THz region reached at most 2.5 THz even in the recent studies due to the high absorption of the liquids [4,5]. Therefore, those of the higher frequency THz dynamics remain unsolved. Recently, new THz-TDS measurement technique has been established using high-speed asynchronous optical sampling (AOS) technique, and enables to measure the high frequency region [6].

Infrared and Raman scattering spectroscopy observe the molecular dynamics through the change of dipole moment and polarizability, respectively. Dielectric loss $\varepsilon''$ and the imaginary part of Raman susceptibility $\chi''$ have been often compared and discussed [7]. For molecules and crystals, the selection rule holds for infrared (IR) active and Raman active modes according their point groups. In contrast for liquids, there is little difference in the frequency shift of a vibrational state, while dipolar aprotic liquids were reported that the relaxation mode of $\varepsilon''$ deviate from that of $\chi''$ [7]. It is significant to investigate a same sample by using both far-IR and low-frequency Raman spectroscopies.

Propylene glycol (PG) is a typical intermediate glass-forming molecular liquid and commonly used for studies of the glass transition which is related to the dynamics of super-cooled liquids [8,9]. Furthermore, the PG esters with no hydroxyl group, which are the modification of PG, have been used for the pharmaceutical application recently [10]. Therefore, the investigation of the relaxation process in the unsolved THz region for a simple glass-forming PG is important.

This work is aimed at the investigation of the broadband THz dynamics using two types of terahertz time-domain spectroscopy (THz-TDS) systems, and the accurate analysis of THz dynamics with the computational calculation using the Gaussian. Furthermore, the combination the results of THz-TDS with those of Raman scattering enable us to discuss the discrepancy between $\varepsilon''$ and $\chi''$. 
Experimental & Calculation Methods

In the analysis of THz-TDS spectra, the observed spectra of two THz-TDS systems are combined to cover a broad THz region [1,11]. The low frequency sides of THz transmission spectra were measured in the frequency range from 0.1 to 3.0 THz using a standard THz-TDS equipment (RT-10000, Tochigi Nikon Co.) with low temperature-grown GaAs photoconductive antennas for both the emitter and detector. The photoconductive antennas were triggered by a mode-locked Ti:sapphire pulsed laser with a wave-length of 780 nm, a pulse width of less than 100 fs, and a repetition rate of 80 MHz [1,11]. The high frequency sides of THz transmission spectra were measured in the frequency range from 1.0 to 5.0 THz using an recently developed THz-TDS equipment (TAS7500SU, ADVANTEC Co.) with a Cherenkov type THz generator and the high speed AOS technique [1,11]. Since the difference in the spectra observed by these systems was at most 5%, we smoothly connected the absolute value of the complex dielectric constants of the samples by multiplying a factor in the data analysis.

We performed three kinds of measurements, which are background, liquid cell (window) only, and liquid sample inserted in the cell as shown in Fig. 1. The Z-cut quartz was used for the window in the measurement of low frequency sides (RT-10000) and high density polyethylene was used for the window in the measurement of high frequency sides (TAS7500SU). The thicknesses of the window and the sample are 5.0 mm and 0.296 mm, respectively in the measurement of RT-10000, and 3.0 mm and 0.200 mm, respectively in the measurement of TAS7500SU. Figure 2(a) shows the measured time-domain THz E-field waveforms transmitted through the air (background), window, and PG observed in the THz-TDS system of RT-10000, and Fig. 2(b) shows the frequency-domain power spectra. Figure 3 shows those observed in the THz-TDS system of TAS7500SU. The absorption dip observed at 3.8 THz in Fig. 2(b) is due to the E_1 mode which is the deformed vibration of silicon atoms in the quartz [12,13]. Figure 3(b) also showed the absorption dip at 2.2 THz, and it is originated from the B_{1g} lattice mode in high density polyethylene [14,15]. From the comparison between the frequency-domain power spectra shown in Fig. 2(b) and Fig. 3(b), it can be clearly seen that the spectra measured by RT-10000 supports the low frequency side of those by using TAS7500SU. To calculate the frequency dependent complex dielectric constant \( \varepsilon = \varepsilon' + i\varepsilon'' \) from the measured time-domain waveforms, we have used following equation,

\[
\frac{E_s(v)}{E_w(v)} = t_{W}t_{SW} \frac{\exp\{i(n_s-1)\alpha d_s / c\}}{1-r_{SW}^2 \exp(i2n_s \alpha d_s / c)}, \tag{1}
\]

where \( E_s(v) \) and \( E_w(v) \) are the amplitude spectra of THz pulse transmitted the sample and the window, respectively. \( c \) and \( d_s \) are the light velocity and a thickness of sample, respectively. \( t_{ij} = 2n_i/(n_i+n_j) \) and \( r_{ij} = (n_i-n_j)/(n_i+n_j) \) are the complex Fresnel’s transmission and reflection coefficients at the interface between regions \( i \) and \( j \), respectively. The subscripts \( i \) and \( j \) stand for \( W, S \), representing the window and sample, respectively. \( n_i \) is the complex refractive index of region \( i \). Then, the complex dielectric constant is obtained from the relation of \( \varepsilon = n^2 \).

Depolarized VH Raman scattering spectra were measured in the frequency range from 0.3 to 5.0 THz in the backward geometry with a scattering angle \( \theta = 180^\circ \), using a single frequency green-YAG laser with wavelength 532 nm. The observed Raman intensity spectra \( I_{raw}(v) \) were converted to the imaginary part of the Raman susceptibility \( \chi''(v) \) by the following equation,

\[
\chi''(v) = \frac{I_{raw}(v)}{n_B(v) + 1}, \tag{2}
\]

where \( n_B(v) = (\exp(hv/k_B T))^{-1} \) is the Bose-Einstein distribution function. The sample of PG was purchased from Sigma-Aldrich, and studied without further purification, and geometry optimizations and the frequency calculations of PG was performed by density functional theory (DFT) using the Gaussian09 B3LYP/6-31G+(d, p) level of the theory [16,17].
Results & Discussion

Figure 4 shows the complex dielectric constants and the χ” in the broad THz region of PG. The combination of two THz-TDS systems enable to observe the broadband dielectric constants spectra and the whole structure of the broad peak at 1.5 THz for ε” in PG. The dielectric spectrum of PG consists of the broad peak at 1.5 THz and the small shoulder at 3.6 THz, while only one peak at 1.5 THz seems to be observed in the χ” spectrum. The peaks at 1.5 THz of alcohols have been discussed for some kinds of vibrational modes. Yomogida et al. observed the peak at 1.5 THz of pentanol isomers and suggested to be the vibrational mode which is originated by the location of the

Figure 1. (a) The schematic diagrams of three kinds of THz transmittance measurements which consists of the transmittance of (a) air (background), (b) window (high density polyethylene or quartz), and (c) sample.

Figure 2. (a) Time-domain waveforms and (b) frequency-domain power spectra of background (black solid lines), window (blue dots lines), and PG (red solid lines) measured by RT-10000 (Tochigi Nikon corp.) at room temperature.

Figure 3. (a) Time-domain waveforms and (b) frequency-domain power spectra of background (black solid lines), window (blue dots lines), and PG (red solid lines) measured by TAS7500SU (ADV ANTEST corp.) at room temperature.
OH groups and the carbon chain structure [18,19]. They fitted the peak at 1.5 THz by using the damped harmonic oscillator (DHO) model, which is commonly used for fitting a vibrational mode.

The shoulder at 3.6 THz observed in the dielectric spectrum of PG has never been reported so far. The Gaussian calculation for PG monomer shows no vibrational mode below 4.5 THz as shown in Fig. 5. The precedent researches of glycerol reported two kinds of vibrational modes at around 3.0 THz for cyclic glycerol trimers. One is the collective motions of the OH groups and the other is the translational motions of the CH₂ and CH groups [20,21]. These experimental and calculation results suggest that the observed small shoulder at 3.6 THz can be attributed to not intra- but inter-molecular vibrations.

The low frequency side of ε” suggests a contribution from the GHz region. The precedent work of the dielectric measurement of PG reported the α- and fast processes exist in the MHz-GHz region, and the high frequency sides of the processes remain in the THz region at room temperature [8]. The comparison between the ε” and χ” doesn’t seem to show the relaxation process in the low-frequency region of PG. We have reported the wide frequency range from GHz to THz for PG of χ” using Raman scattering and Brillouin scattering [22]. This result showed the relaxation processes of the α- and fast processes; however, the relaxation strength of these relaxation process of χ” is much smaller than those of ε” measured by Kohler et al [8]. The similar results were reported by Fukasawa et al. [7], and they suggested that the relaxation arising from the long range hydroxyl bond mediated dipole-dipole interactions gives the main contribution to the dielectric spectrum, and this process is Raman inactive.

Figure 4. (a) The ε’ and (b) the comparison between the ε” and the χ” (blue solid line) of PG at room temperature. In the complex dielectric constants spectra, the orange circles show the result from RT-10000 and the red dots show the result from TAS7500SU.
Conclusion

We have investigated the broadband terahertz dynamics (0.1 to 5.0 THz) of the liquid state of PG by using two different THz-TDS systems and Raman scattering equipment. Two THz-TDS systems enable to observe the whole spectral structure of the broad peak at 1.5 THz in PG. The peak at 1.5 THz is observed both in $\varepsilon''$ and $\chi''$, while the small shoulder peak at 3.6 THz is only observed in the $\varepsilon''$ spectrum. In contrast to the high frequency region, the low frequency side of the $\varepsilon''$ spectrum of PG includes the contribution from the GHz region, and the precedent report of the GHz dielectric measurement supports this suggestion [8]. From the comparison between $\varepsilon''$ and $\chi''$, the line-shape of the spectra is clearly different, and it suggested that it relates to the motion of the OH groups which cause the hydrogen bond mediated dipole-dipole interactions.

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