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Time-resolved detection of structural change in polyethylene films using mid-infrared laser pulses

Eduard Ageev, Keisuke Mizobata, Takashi Nakajima, Heishun Zen, Toshiteru Kii, and Hideaki Ohgaki

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Time-resolved detection of structural change in polyethylene films using mid-infrared laser pulses

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Some of the vibrational modes of crystalline organic polymers are known to be sensitive to the structural change from the crystalline phase to the amorphous phase, and vice versa. Using a mid-infrared (mid-IR) pulse from a free-electron laser as a probe, we demonstrate the time-resolved detection of structural change in crystalline polymer (polyethylene) films upon laser heating by a Q-switched Nd:YAG laser. Transmittance of the resonant mid-IR pulse almost instantaneously changes before and after the Nd:YAG laser pulse if its fluence is sufficient to induce the structural change in the film. The developed technique would be useful to study the time-dependent dynamics of the structural change in various materials. © 2015 AIP Publishing LLC.

Various kinds of phase transitions have been attracting a lot of interests from the viewpoint of manifestation and control of the specific functions of materials. In the case of inorganic materials, magnetic phase transition and metal-insulator transitions are of great interest. On the other hand, in the case of organic materials, the structural phase change itself, which is of course accompanied by the change of the physical and chemical properties, is of great interest. Polymers are the commonly used organic materials for many purposes, and they are classified into two categories: amorphous polymers such as acryl and polystyrene and crystalline polymers such as polyethylene (PE) and polyethylene terephthalate (PET). We point out that the crystallinity of crystalline polymers is not 100%, and they always contain the amorphous component to some extent. Although PE is the simplest and very commonly used crystalline polymer with only carbon and hydrogen atoms as constituent, there is no paper in the literature which reports the time-resolved study on its structural phase change.

To determine the structure of organic materials, there are many ways. Among others, mid-infrared (mid-IR) spectroscopy, such as FTIR and Raman spectroscopy, nuclear magnetic resonance, x-ray diffraction, etc., are the well-established techniques. However, the available methods to detect the structural change, which is of course accompanied by the change of the physical and chemical properties, is of great interest. Polymers are the commonly used organic materials for many purposes, and they are classified into two categories: amorphous polymers such as acryl and polystyrene and crystalline polymers such as polyethylene (PE) and polyethylene terephthalate (PET). We point out that the crystallinity of crystalline polymers is not 100%, and they always contain the amorphous component to some extent. Although PE is the simplest and very commonly used crystalline polymer with only carbon and hydrogen atoms as constituent, there is no paper in the literature which reports the time-resolved study on its structural phase change.

To determine the structural change in various materials, direct detection of the structural change is one of the most useful ways. Among others, mid-infrared (mid-IR) spectroscopy, such as FTIR and Raman spectroscopy, nuclear magnetic resonance, x-ray diffraction, etc., are the well-established techniques. However, the available methods to detect the structural phase change itself, which is of course accompanied by the change of the physical and chemical properties, is of great interest. Polymers are the commonly used organic materials for many purposes, and they are classified into two categories: amorphous polymers such as acryl and polystyrene and crystalline polymers such as polyethylene (PE) and polyethylene terephthalate (PET). We point out that the crystallinity of crystalline polymers is not 100%, and they always contain the amorphous component to some extent. Although PE is the simplest and very commonly used crystalline polymer with only carbon and hydrogen atoms as constituent, there is no paper in the literature which reports the time-resolved study on its structural phase change.

In this paper, we report the time-resolved detection of structural change (crystalline → amorphous) in PE films using a mid-IR FEL developed at Kyoto University (KU-FEL). To induce and detect the structural change in PE films, we employ two laser pulses: Q-switched Nd:YAG laser pulse to heat the PE film on an NaCl crystal and mid-IR FEL pulse to probe the structural change of PE. Note, however, that PE itself is practically transparent in the visible ~near-IR range, and the laser-heating with the Nd:YAG laser would not be efficient. Indeed, the brute force irradiation of the Nd:YAG laser pulse at high fluence onto the PE film results in the material damage before the sufficient heating takes place. To overcome this problem, the PE film is doped with absorptive dye which has large absorption around 1064 nm. There are various ways to detect the phase change: through the reflectivity change by the streak camera, transmission change of the probe pulse in the visible or THz range, or the change of the Raman signal. In this work, we measure the transmission change of the mid-IR probe pulse which is resonant with a structure-sensitive vibrational mode of PE. Currently, the time resolution is limited by the response of the mid-IR detector we employ, and it is a few hundreds of nanosecond time resolution. It can easily go down to a few nanoseconds only if the mid-IR detector is replaced by a faster one.

Dye-doped PE films we use in this study are fabricated in the following way. First, we dissolve a high-density PE pellet (Sigma-Aldrich, CAS: 9002-88-4, mp:125–140 °C) into the o-Dichlorobenzene solution (Wako special grade, CAS: 95-50-1) at 170 °C on a hot plate for a few hours to obtain the solution with the PE concentration of 3 wt. %. Then, we preheat a NaCl crystal (25 mm diameter, 2 mm thickness) mounted on the spin-coater and a syringe tip (made of polypropylene) to 150 °C. The preheating of a NaCl crystal and a syringe tip is very important, since the use of a syringe tip at high temperature makes the PE solution more viscous and the photoirradiation in the subsequent experiments becomes more efficient.

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room temperature results in the immediate solidification of the PE solution in the tip. Recall that PE has very high chemical resistance to most of the organic solvents, and it is very difficult to dissolve PE. Similarly, a drop of the PE solution in a preheated syringe onto a NaCl crystal at room temperature results in the immediate solidification of the PE solution before spreading over the crystal surface. Just before the spin-coating, we add the appropriate amount of IR165 dye (Exciton) into the PE solution at 150 °C to minimize the thermal damage of dye molecules. The dye concentration in the PE solution is carefully controlled to be 0.5 wt. %. Using a surface profilometer, the thickness of the dye-doped PE film fabricated this way is found to be 2.0 ± 0.4 μm.

Prior to the main experiment, we scan the wavelength of the FEL beam without introducing the Nd:YAG laser pulse and measure the transmission spectra through the PE film with two energy detectors (Gentec QE8SP-I-BL-BNC) of pyroelectric type for the reference and transmission signals to minimize the influence of the shot-to-shot fluctuation of FEL pulse energy. We repeat the similar measurement with a bare NaCl crystal to account for the transmission loss by the NaCl crystal itself. Thus, the transmission signals are converted to the transmittance. In Fig. 1, we show the transmittance of the dye-doped PE film at 730 cm⁻¹ (13.7 μm) with ~10 mJ energy and ~1.5 μs duration for the macropulse at the 1 Hz repetition rate goes through the monochromator (slit width 0.5 mm) to narrow the FEL linewidth and then 2 mm aperture, as a result of which the FEL macropulse energy at the sample is less than several μJ. Each FEL macropulse contains a few thousands of micropulses with a sub-ps duration and 350 ps time interval, but our detectors do not have a time-response to resolve them. To minimize the influence of the shot-to-shot change of FEL pulse energy, we employ two photodetectors (Teledyne Judson Technologies J15D12-M204-S01M-60) of mercury cadmium telluride (MCT) type for the reference and transmission signals. Since the MCT detectors have undesired sensitivity at the wavelength of the Nd:YAG laser (Rayture systems GAIA-III, 1064 nm, 5 ns pulse duration, max. 40 mJ pulse energy) we use to heat the PE film, long pass filters are placed in front of both detectors. To synchronize the Nd:YAG pulse and the FEL pulse, we use the radio-frequency (RF) pulse for electron acceleration of FEL as a master trigger for the delay generator (SRS DG645), through which the Q-switch of the Nd:YAG laser is triggered with an appropriate delay. The timing of the Nd:YAG laser pulse with respect to the RF pulse is adjusted so that the Nd:YAG laser pulse is turned on just after the peak of the FEL macropulse. There is tens of ns jitter between the RF pulse and the peak of the FEL pulse. This is due to the shot-to-shot difference of the amplification processes. This jitter, however, is negligible for the accuracy of the present experiment. The diameter of the Nd:YAG laser pulse is slightly shrunk with a convex lens to result in 4 mm diameter (defined for 1/e² and measured by the beam profiler, Thorlabs BC106-VIS) at the sample, which is sufficiently larger than that of the FEL pulse.

![Fig. 1. Transmittance of the dye-doped PE film at 32 and 84 °C obtained by scanning the FEL wavelength. An FTIR spectrum of the same sample is also shown for comparison.](image1)

![Fig. 2. Experimental setup. M, BS, L, A, and LPF stand for the gold mirror, ZnSe beam splitter, f = 150 mm plano-convex ZnSe lens, 2 mm aperture, and long pass filter. RD and TD stand for the MCT detectors for the reference and transmission signals of FEL pulses, respectively.](image2)
In Fig. 3, we show the transmittance at 1064 nm through the dye-doped PE film as a function of Nd:YAG laser fluence. We first increase the fluence from 30 mJ/cm² up to 390 mJ/cm² after the irradiation of 300 shots at 10 Hz at each fluence. At 75 mJ/cm², we notice the slight damage on the film surface by naked eyes, and we define this fluence as damage threshold. We further increase the fluence up to 390 mJ/cm². Increase of transmittance at the higher fluence implies that more dye molecules are destroyed by the irradiation of more intense laser. After the fluence of 390 mJ/cm² is reached, we gradually decrease the fluence back to 30 mJ/cm². During the decrease of fluence, the transmittance at 1064 nm remains to be the same, as expected.

Now, we tune the FEL pulse to be resonant with the absorption line of PE film at 730 cm⁻¹ and measure its transmission through the PE film with the Nd:YAG laser on at the fluence of 77 mJ/cm² to find that there is no transmission change of the FEL pulse before and after the Nd:YAG laser pulse (Fig. 4(a)). It is not until 390 mJ/cm² that we observe the clear transmission change before and after the Nd:YAG laser pulse (Fig. 4(b)). In order to ensure that this transmission change is indeed due to the structural change of the dye-doped PE film, we carry out a similar experiment at the same Nd:YAG laser fluence but with the off-resonant FEL wavelength of 712 cm⁻¹ (Fig. 4(c)). From the comparison of Figs. 4(a)–4(c), we can confidently conclude that the transmission increase of the resonant FEL pulse upon irradiation of the Nd:YAG laser pulse (Fig. 4(b)) with sufficient fluence must be due to the structural change (crystalline \(\rightarrow\) amorphous) of the PE film upon melting by laser heating. The slight modulations in the FEL signals of Fig. 4 are due to the noise from the RF source for electron acceleration of FEL, which is different from shot-to-shot, and hence, it cannot be subtracted when we plot the transmission change. The increase of transmission change in Fig. 4(b) is in the time scale of \(\sim\)μs, which is much longer than the time for the pressure-induced phase change in polytetrafluoroethylene (PTFE).\(^{15}\) By noticing that the local temperatures are different at different film depths in our case because neither Nd:YAG laser fluence nor distribution of dye molecules are uniform (see Fig. 2), we expect that the apparent response time for the transmission change would be much longer than the intrinsic response time for the structural change of a single polymer. This explains why the time scale for the phase change we find in Fig. 4(b) is \(\sim\)μs. In the time range of several μs, we only observe the transmission increase, and do not observe the decrease back to the initial value. This implies that the time needed for the film to cool down is much longer than this time scale.

Before closing the paper, we make a rough estimation of the film temperature increase upon irradiation of Nd:YAG laser pulses. This is not an easy task, since the dye
concentration is not uniform along the laser propagation axis. Recall that the Nd:YAG laser pulse at high fluence destroys some of the dye molecules as we have seen in Fig. 3. Qualitatively, however, we can say, that, at the position deeper in the film, more dye molecules survive the irradiation of intense Nd:YAG laser pulse. From the experimental fact that we can see the obvious transmission change of the resonant FEL pulse at the Nd:YAG laser fluence of 390 mJ/cm² and not below this value, we consider that the temperature at the deepest part of the film can go beyond the melting point around this fluence. Assuming that none of the dye molecules are destroyed and its concentration is nearly uniform at the film depth of 1.9–2.0 µm, which means that the Beer’s law is valid at this film depth, we can calculate the absorbed pulse energy to be about 1.3% with the aid of Fig. 3. Then, using the formula of $FA = \rho h c (\Delta T + L/c)$ where $F$ is Nd:YAG laser fluence, $A$ is absorptance, and $\rho$, $h$, $c$, $L$ are the density (0.952 g/cm³), thickness (~2 µm), heat capacity (2.3 J/g °C), and latent heat (286.2 J/g) of PE with $\Delta T$ being the temperature increase, we obtain $\Delta T = 114$ °C at the deepest part (1.9–2.0 µm) of the PE film. This agrees well with the minimum temperature increase, 110 °C, required to melt the PE film.

In conclusion, we have demonstrated the time-resolved detection of structural change in crystalline polymer (polyethylene) films upon laser heating. When we tune the mid-IR probe pulse to the structure-sensitive vibrational mode of polyethylene, transmission of the probe pulse clearly changes before and after the laser heating pulse if the fluence of the heating pulse is sufficient. The experimental technique we have developed in this work would be useful to study the time-dependent dynamics of the structural change in various polymer materials. For the more detailed study of the dynamics, it is essential to fabricate a thinner (<µm thickness) polymer film with better quality so that we can see the structural change without damaging the dye molecules and carry out more refined experiments to clarify not only the heating but also the cooling processes to understand the re-crystallization dynamics, which is underway in our group.

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