

The Third Order Nonlinear Optical behaviour of Poly Ether Urethane

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ARTICLE INFO	ABSTRACT
Article history: Received 1 July 2017 Revised 18 October 2017 Accepted 19 October 2017 Available online 20 October 2017	Poly (ether) urethane close/open cell (PEUCC/PEUOC) has been prepared by using in situ polymerization approach. The third-order optical nonlinearities of dissolved samples in DMF were characterized by Z-scan technique at the measurement wavelength of 532 nm. The synthesized samples were also examined by optical microscopy and UV-Vis absorption spectroscopy. The nonlinear refractive (NLR) indices of the synthesized
<i>Keywords:</i> Nonlinearity Polyurethane Z-scan	samples were obtained in the order of 10 ⁻¹⁰ (cm ² /W) with negative sign. By focusing on the collected UV-Vis absorption spectra of PEUCC/PEUOC, it was found that the red shift in the absorption edge of the samples is larger in the case of PEUCC that in the PEUOC. It was revealed that introducing some materials into polyurethane (PU) spatially PEUCC could increase nonlinearity that they can be considered as a promising candidate for the application to optical limiting in the visible region.

چکیدہ	اطلاعات مقاله
۔ سلول بسته/باز پلی (اتر) پلی یورتان (PEUCC/PEUOC) از طریق روش پلیمری کردن در محل تهیه گردید. تابعیت	تاريخچه مقاله:
غیرخطی بودن نوری مرتبه سوم نمونه های حل شده در DMF به وسیله روش جاروب Z با طول موج لیزری ۵۳۲ نانومتر	دریافت: ۱۰ تیر ۱۳۹۶
مشخصه یابی گردید. نمونه های سنتز شده همچنین توسط میکروسکوپ نوری و طیف سنجی جذبی ماورای بنفش مریی	تصحيح: ۲۶ مهر ۱۳۹۶
تحت آزمایش قرار گرفتند. شاخص های شکست (NLR) نمونه های سنتز شده در مرتبه ۱۰–۱۰ سانتیمتر مربع بر وات و	پذیرش: ۲۷ مهر ۱۳۹۶
با علامت منفی به دست آمدند. با کانونی کردن طیف های جذبی ماورای بنفش / مریی PEUCC/PEUOC مشخص	
گردید که جابه جایی قرمز در لبه جذبی نمونه ها بزرگتر از حالت PEUCC می باشد. همچنین این موضوع تایید گردید	کلید واژگان:
که با اضافه کردن موادی به پلی یورتان (PU) به صورت حجمی، PEUCCمی تواند میزان پارامتر غیرخطی را افزایش	قابلیت غیر خطی
دهد و می تواند به عنوان یک کاندیدای مناسب برای کاربرد در محدودیت های نوری در ناحیه مریی در نظر گرفته شود.	پلیمر پلی یورتان
	جاروب Z

1 Introduction

PU has attracted an outstanding amount of attention because of extensive applications such as textiles, thermosetting resins, medical device, and smart actuators [1, 2]. Nonlinear optical materials due to their wide range of potential applications are always of considerable interest, such as in optical limiting and in all-optical photonics devices [1-4]. Additionally, organic polymeric materials are suitable for nonlinear optical applications due to having easy modified structure, the ability of converting into thin films, high laser damage thresholds, faster response time and other improved properties [5]. In particular, good nonlinear optical properties of some inorganic materials have stimulated great research interest in this field [6, 7].

In the present paper, we describe the synthesis of two new polyurethane foams with focusing on their optical properties such as optical microscopy imaging and UV-Vis spectroscopy. Furthermore, the third-order nonlinear optical properties of PEUCC/PEUOC was obtained by using the closed- aperture (CA) technique with a Nd:YAG laser at a 532 nm wavelength with continuouswave (CW).

2 Experimental

2.1. Materials

PU foam material composed of two commercially reactants: polyisocyanate and polyol components in the form of liquid supplied by Exxon Panah Co., Iran. The polyisocyanate employed was diphenyl methane diisocyanate (MDI, ρ =1.23gr/cm³) from Merck Co. and polyol component based on polyether polyol(ρ =1.1g/cm³), blowing agent (water in PEUOC and HCF–C in PEUCC), catalyst and surfactant were prepared from Exxon Panah Co., Ltd., Tehran, Iran.

2.2. Preparation of PEUOC/PEUCC

First of all, the amount of blowing agent, catalyst and surfactant was dissolved into close/open cell polyol part by using an electrical MS2 Minishaker IKA (Germany). This stage lasted for 20 seconds with 3000 rpm until a homogenized solution was reached. Then the MDI part was added to the prepared solution by vortex mixing with 2000 rpm for 4–5 seconds. Due to well-prepared sample and producing CO_2 gas during reaction time, the cover of the container of samples was taken off. After 10–12 seconds reaction was completed by forming of foams with the equal ratio of polyol:MDI. For the analysis purposes, the samples were kept in the stream of liquid nitrogen gas and then were cut in slices with 1mm diameter. The chemical reaction is illustrated in Fig. 1.



Fig 1. Schematic of chemical reaction between polyol and isocyanate.

2.3. Characterization

Optical micrographs were taken using an optical microscope (Nikon; TE 2000-S) in transmission mode with ×40 magnification. The absorption spectra of samples have been analysed by collecting the UV-Visible-Near-IR spectra and using a double beam optical spectrometer model T80+ manufactured by PG Instrument. The experimental Z-scan set-up of the proposed technique is shown schematically in Fig. 2.



Fig. 2. Experimental set-up for the closed aperture Z-scan technique.

According to typical preparation for this method, PU have dissolved N. foams been into N-Dimethylformamide (DMF) in 60°C until a solution with 0.01 M was achieved. For Z-scan method, an incident laser beam from a CW Nd:YAG laser at its second harmonic frequency of 532 nm propagating with TEM_{00} Gaussian profile and focused by a lens (f = 5 cm) onto the sample was used. The radius of the beam waist was determined as 37 μ m. The sample is moved along the z axis in the vicinity of the focal length lens (the +z direction is the laser propagation direction). The transmitted intensity is collected and recorded by a photo diode detector.

3 Results and discussion

3.1. Microscopic evaluation

The cell size of PEUCC/PEUOC can be estimated by optical microscopy. For this aim, thin layers of pure PEUCC/PEUOC with thicknesses of about 1mm were cut perpendicular to the foams rising direction with a sharp razor blade into circular slices with 5 mm radius and 1mm thickness. The liquid nitrogen was used for freeze–fractured surfaces of samples and after that they were examined.

Figure 3 shows the transmitting optical microscopy images for PEUCC/PEUOC. Accordingly, by focusing on micrographs of samples, it is clear that PEUOC foams have smaller cell sizes in comparison with PEUCC.



Fig. 3. Microstructures of: (a) PEUCC, (b) PEUOC.

This finding is reasonable because the produced CO_2 gas during reaction time in PEUCC remains in the cells and the mean cell sizes of PEUCC will be larger in compare with PEUOC foams.

3.2. The UV-Vis spectra

Figure 4 shows the optical absorption spectrum recorded in the spectral region of 190 to 650 nm for PEUCC and PEUOC foams dissolved in N, N Dimethylformamide.

The absorbance values in PEUCC are more than PEUOC foams. It can be understood that PEUCC show

red shifts in the absorption edges relevant to PEUOC. In addition, the cut-off wavelength was measured to be at around 300-350 nm wavelength. The energy band gap of PEUCC/PEUOC was calculated by using cut-off wavelength. The obtained values for direct band gap transition of PEUCC and PEUOC are 3.2 and 3.7eV, respectively.



Fig. 4. UV/Vis absorption spectra of PEUOC and PEUCC.

Since the ratio of polyol and isocyanate is equal, the amount of polyol which was involved into closed cell of PEUCC, its equal amount of isocyanate will remain in the interaction container. Since polyol has no double or π bonding and isocyanate has π bonding and the π -electron delocalization in organic materials is responsible for high tendency to absorb UV radiation; then the UV absorption of PEUCC will be stronger than PEUOC [2, 8].

3.3. Nonlinearity capability

3.3.1. Z-scan theories on nonlinear process

The linear absorption in low incident powers was calculated by the following "Eq. (1)" that P and P_0 are related to the output power with and without sample, respectively [9-11]

$$\alpha = -\frac{1}{L} Ln \left(\frac{P}{P_0} \right). \tag{1}$$

The phase shift incurred by an optical beam travelling a distance L in the medium is [7, 9-11]:

$$\left|\Delta\varphi_{0}\right| = -\left(\frac{2\pi}{\lambda}\right)\Delta n L_{eff} . \tag{2}$$

Where the L_{eff} is an effective sample thickness, $L_{eff} = (1 - e^{-\alpha L})/\alpha$, and Δn is referred to NLR index [10, 11]. The usual relation about nonlinear refractive index is $n = n_0 + n_2 I_0$, in which n, the total refractive index, n_0 , linear refractive index, n_2 , nonlinear refractive index and $I_0 = 2P_{in}/(\pi \omega_0^2)$, is the incident intensity at focal point that is 2493.7 W/cm² and P_{in} is the laser power that is 55 mW in this work [12]. Also ω_0 is the radius of the waist of the illumination beam inside the sample was detected by edge scan method. According to these formulas, the equivalent relation for "Eq. (2)" is [10, 11]:

$$\left|\Delta\varphi_{0}\right| = -\left(\frac{2\pi}{\lambda}\right)n_{2}I_{0}L_{eff} . \tag{3}$$

In CA scheme, an aperture placed at the far field position by monitoring the transmittance change through a small aperture one which is able to determine the amplitude of the phase shift. By using CA scheme, the value of distance between the normalized transmitted peak and transmitted valley is nominated by ΔT_{p-v} which it related to $|\Delta \varphi_0|$ with the following expression [10, 11]:

$$\Delta T_{p-\nu} = f \left| \Delta \varphi_0 \right| \qquad for \quad \Delta \varphi_0 \prec \pi.$$
⁽⁴⁾

In this relation f is an experimental constant and is assigned by means of a parameter as aperture linear transmission, $S = 1 - \exp(-\frac{2r_a^2}{\omega_a^2})$, that is about 0.3 (or 1) for close (or open) aperture Z-scan experiment. r_a is the radius of the beam in the aperture position. The experimental constant f is introduced with the following equation $f = 0.406(1-S)^{0.25}$, then by putting the amount of f into "Eq. (4)" the nonlinear refractive index, $n_2(cm^2/W)$ can be obtained from the following equation [10, 11]:

$$\Delta T_{p-\nu} = 0.406(1-S)^{0.25} (2\pi / \lambda) n_2 I_0 L_{eff}$$
(5)

It should be noted that the experimental results of Zscan data with an aperture is divided by those without an aperture to obtain pure nonlinear refraction.

3.3.2. Closed aperture Z-scan results

According to typical preparation for Z-scan method, PEUOC/PEUCC foams have been dissolved into DMF since the solution with 0.01 M was achieved. The sample was placed in a 1-mm-thick quartz cell. In Zscan experiment, the sample was moved forward or backward along the direction of the laser beam around the focus.

In the present work, a CW Nd:YAG laser operating at 532 nm with TEM_{00} Gaussian profile focused by a lens (f=5cm) onto a sample was used as the laser source for the Z-scan experiment. The laser beam waist set at the focal point and the radius of the waist of the illumination beam inside the sample (ω_0) was $37\mu m$ and The Rayleigh diffraction length is 8.1 mm at the focal plane. The CA (S = 0.32) scheme allowed to determine both of the sign and the magnitude of n_2 . Fig. 5 shows pure closed Z-scan data of two different foams. As it can be seen in the normalized curves of CA Z-scan exists a pair of peak and deep valley which can be identified as a selfdefocusing material [12]. Since the valley comes after the peak of the transmittance, the sign of the refractive nonlinearity of synthesized samples are negative that it shows self-defocusing effect of them. The NLR indices coefficients of the synthesized samples were obtained from "Eq. (5)" in the order of $10^{-10} (cm^2/W)$ with negative sign. As a result, PEUOC foam has higher NLR index than PEUCC foam then the nonlinear effect is more

than PEUCC foam then the nonlinear effect is more obvious in PEUOC than PEUCC foams. We considered that the values of n_2 are originated from ultrafast electronic nonlinear changes in the refractive index to be compared with the thermal nonlinear change in it [6, 13, 14]. This estimative confirms the theoretical and general estimative made by Boyd [6], that $n_{2(thermal)} \approx$ $10^{-5} cm^2/W \gg n_{2(electronic)}$ [14].



Fig. 5. Close aperture Z-scan experimental curves of PEUOC and PEUCC.

Furthermore, it can be noticed in Fig. 5 that the peak height $(T_{P}-1)$ and the valley depth $(1-T_{V})$ increase coincidentally, where T_P and T_V are the normalized peak and valley transmittance in the Z-scan drawing, respectively. Furthermore, in PEUOC the phenomenon of the peak and valley exhibited asymmetrical behaviour, the peak of it was restrained according to PEUCC while the valley was enhanced. The measurement conditions and results are listed in Table 1. According to similar reports of nonlinearity of PEU, the nonlinear refractive index of PEUCC is smaller than PEU as well [15]. It was shown that there are various physical mechanisms such as thermal and ultrafast electronic effects that can produce the NLR index change in both types of polymers. A considerable contribution to the NLR index under a CW

Table 1. Z-Scan technique and the determined n_2 value for

Sample	$I_0(W/cm^2)$	$\alpha(cm^{-1})$	$L_{eff}(mm)$	ΔT	$n_2(cm^2/W)$
PEUCC	2493.7	0.89	0.957	1.87×10 ⁻²	1.8×10^{-10}
PEUOC	2493.7	1.04	0.95	5.97×10 ⁻²	5.7×10 ⁻¹⁰

laser for sample illumination can take place due to thermal effects [9]. Thus, we expect that thermal effect overcomes the other effects.

3 Conclusions

Two poly ether urethanes (PEUOC and PEUCC) have been prepared by using in situ polymerization approach. The third-order optical nonlinearities of PEU dissolved in DMF are characterized by Z-scan technique with CW Nd:YAG laser at its second harmonic frequency of 532 nm as a light source. They exhibit good optical limiting properties at the wavelength used. The results conclude that they can be a promising candidate for the future optical device applications. The other following results can be concluded:

• By evaluating optical micrographs of the synthesized foams, PEUCC has bigger cell sizes than PEUOC foams.

• By focusing on the collected UV-Vis absorption spectra data of PEUOC/PEUCC samples, the optical band gap was measured. The red shift in the absorption edge of the samples was larger in the case of PEUCC. Therefore, PEUCC has lower band gap than PEUOC ones.

• The NLR index of the samples was obtained from close aperture Z-scan in the order of $10^{-10} (cm^2/W)$ with negative sign for both of PEU.

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