INTRODUCTION

Photodeformable polymers have received increasing attention in virtue of their potential in various applications such as artificial muscles, smart robots and locomotion systems.[1,2] Current efforts have been concentrated on the development of photodeformable liquid crystal polymers (LCPs)[3−6] due to their unique advantages endowed by liquid crystal molecules, such as self-organization, reversible phase transition and macroscopic orientation.[7] These properties allow the material to produce large and fast photoinduced mechanical responses when photosensitive chromophores are incorporated into LCPs. As important photoreponsive chromophores, azobenzene (azo) groups provide azo-functionalized LCPs with photoresponsive properties and bring a reversible shape change upon exposure to UV or visible light because of the reversible trans-cis photoisomerization.[8,9]

The azo-containing LCPs have been mostly prepared via free radical polymerization with chemical crosslink, which limits their application for fabricating complex devices due to the incompatibility with existing industrial processing methods. In order to break through this limitation, we designed azo-containing linear LCPs (LLCPs) with excellent mechanical properties endowed by high molecular weight through ring-opening metathesis polymerization (ROMP).[10−12] Owing to the linear structure, traditional melting and solution processing methods were applicable in the processing procedure of LCPs, which was unattainable for the cross-linked LCPs. Furthermore, without the need for external orientation treatments (rubbing,[13−15] electric field,[16] magnetic field,[17,18] and others), an annealing treatment was enough to endow LLCPs with a highly ordered structure by the aid of flexible main chain and spacer.[19] These remarkable properties make these LLCPs excellent candidates for many applications, which inspire us to develop a homologous series with different molecular structures to satisfy different purposes. Their chemical and physical properties can be varied by introducing azo-derivatives, which act as both the mesogenic and photoresponsive units and have significant influence on the phase transition and photoresponsive behaviors.[20,21]

Photo-deformation of azo-containing LCP is normally derived from the photoinduced reversible phase transition between the ordered mesophase and the isotropic phase within the temperature range of mesophase. Therefore, it is important to widen mesophase temperature range for LLCPs to realize reliable photoresponsive properties in a broad temperature range. Azoester consisting of ester (−COO−)-azo (−N=N−) linkages is a useful group existing in liquid crystal molecules,[22] which is applicable for thermographic devices, optical imaging, and light emitting diodes.[23] When the phenyl is coupled with ester-azo linkage, the obtained
azoester is rigid and has a large length-diameter ratio. Compared with azo, the corresponding melting temperature and isotropic temperature of azoester increased accompanied by the widening of mesophase temperature range, which indicated the good phase stability of the compounds. Thus, the azoester is an ideal mesogen for preparing phase stable photodeformable LCLCs to enrich the diversity of LCLCs and enlarge their application field.

In this study, azoester was introduced into LLCP to improve the phase stability attributed to the broad mesophase temperature range. The mesomorphic and thermal behaviors were investigated by polarizing optical microscopy (POM) and differential scanning calorimetry (DSC). The spontaneous orientation and photoresponsive behaviors of the LLCP were also studied. Moreover, the relationship between photodeformable behaviors and influencing factors, such as UV light intensity, film thickness and temperature, was investigated in detail.

EXPERIMENTAL

General Considerations
All chemical reagents required for preparation were used as received. Solvents were dried and distilled prior to use. 1H-NMR spectra of PC11AE6 were recorded on a Bruker DMX500 NMR spectrometer using tetramethylsilane as the internal standard and chloroform-d as solvent. High resolution mass spectra (HR-MS) data were obtained by using electron spray ionization (ESI) and chloroform-d as solvent. High resolution mass spectra (HR-MS) data were obtained by using electron spray ionization (ESI) and tetramethylsilane as the internal standard.

Synthesis of monomer C11AE6

5-Succinyl-1-cyclooctene (D) was synthesized according to a literature. Intermediately, C (1.18 g, 2 mmol), D (0.54 g, 2.4 mmol) and DMAP (0.25 g, 2 mmol) were added into 200 mL of CHCl3. The solution of D (0.77 g, 4 mmol) in CHCl3 (20 mL) was then added dropwise into the reaction mixture at room temperature. After the mixture was stirred at room temperature under argon for 48 h, the crude product was purified by silica gel column chromatography using CH2Cl2 as eluent. The vacuum-evaporation and recrystallization procedures were necessary to obtain product (3.1 g, 5.27 mmol, 72.2%).

The yield of monomer C11AE6 was 53%. 1H-NMR (500 MHz, chloroform-d, δ, ppm): 8.32 (d, J = 8.5 Hz, 2H), 7.96 (d, J = 8.5 Hz, 4H), 7.14 (d, J = 9 Hz, 2H), 7.02 (d, J = 8.5 Hz, 2H), 6.94 (d, J = 9 Hz, 2H), 5.67–5.59 (m, 2H), 4.88–4.83 (m, 1H), 4.10–4.05 (m, 4H), 3.97 (t, J = 13 Hz, 2H), 2.62–2.56 (m, 4H), 2.16 (m, 4H), 1.92–1.77 (m, 6H), 1.66–1.59 (m, 4H), 1.52–1.45 (m, 4H), 1.40–1.28 (m, 16H), 0.92 (t, J = 14 Hz, 3H).

Synthesis of monomer C11AE6

The polymerization of monomer C11AE6 was performed by using ROMP. C11AE6 (400 mg, 0.5 mmol) in distilled CH2Cl2 (1 mL) was placed into a Schlenk flask (25 mL). After the solution was degassed by freeze-pump-thaw cycles for three times, second generation Grubbs catalyst (0.8 mg) in 1 mL of degassed CH2Cl2 was injected into the Schlenk flask by syringe under argon. The mixture was stirred at 48 °C for 2 h. After vacuum evaporation, the reaction was terminated using ethyl vinyl ether, and a small amount of CH2Cl2 (~2 mL) was added to improve stirring. The contents were then precipitated into cold methanol, filtered, and dried under vacuum to yield the PC11AE6 (324 mg, 81%).

Synthesis of 4′-(11-hydroxyundecyloxy) hexyloxyphenyl azobenzoate (C)

Compounds A and B (Scheme 1) were prepared according to the method reported previously. The HCl and NaOH aqueous solutions in distilled water were added to adjust the pH of the reaction system. The yields of A and B were 63.33% and 60.76%, respectively. A mixture of B (3 g, 7.3 mmol) and 4-(hexyloxy) phenol (2.1 g, 10 mmol) and 4-dimethylaminopyridine (DMAP, 0.89 g, 7.3 mmol) was stirred in 500 mL of CH2Cl2 under argon. The solution of 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDC, 2.87 g, 15 mmol) in CH2Cl2 (20 mL) was then added dropwise into the reaction mixture at room temperature. After stirring for 48 h, the precipitate formed during the reaction was filtered off and the solvent was evaporated to obtain a crude product. Then, the crude product was purified by silica gel column chromatography using CH2Cl2 as eluent.

Preparation of PC11AE6 Films and Fibres

The thin films were prepared by the drop-coating method using a CH2Cl2 solution of PC11AE6 (3 wt% or 5 wt%) which was cast onto the dry and clean glass substrates. After solvent evapo-
RESULTS AND DISCUSSION

Synthesis and Characterization of PC11AE6

The azoester-containing LLCP (PC11AE6) was prepared as shown in Scheme 1. The $^1$H-NMR spectrum of obtained polymer is shown in Fig. 1. The characteristic aromatic proton signals from 6.93 ppm to 8.31 ppm were observable and proportionable, which indicated the presence of azoester rigid unit. The presence of resonance signal around 5.35 ppm corresponding to the double-bond protons “g” from the main chain indicated the accomplishment of polymerization, with the absence of resonance signal at 5.6 ppm, which corresponded to the double-bond protons from cyclooctene of monomer C11AE6. The resonance signal area ratio of “$a/h = 2$” (7.01 and 4.87 ppm) was consistent with theory, which demonstrated the completed conjunction between main chain and side chain. The molecular weight of obtained polymer was $3.0 \times 10^5 \text{g}\cdot\text{mol}^{-1}$, which was one order of magnitude larger than that of the azo-containing methacrylate polymers obtained by conventional free radical polymerization. With enhanced mechanical robustness endowed by high molecular weight, free-standing multidimensional shapes of PC11AE6, as shown in Fig. S1 (in the electronic supplementary information, ESI), were prepared by different processes, which satisfied different requirements in many application fields.

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Photochemical Isomerization of PC11AE6

The photoisomerization of azoester chromophore is at the origin of the attractive properties of polymers containing azoester mesogens. As shown in Fig. 2(a), upon firstly irradiation with 365 nm UV light (10 mW·cm\(^{-2}\)), PC11AE6 in CH\(_2\)Cl\(_2\) solvent underwent trans-cis isomerization. As time went on, the intensity of the \(n\rightarrow \pi^*\) transition band around 363 nm decreased, while the intensity of the \(n\rightarrow \pi^*\) transition around 453 nm increased slightly. This procedure took 10 s until a photostationary state was eventually achieved. The existence of isobestic points is a strong evidence of the presence of two distinct absorbing bands in equilibrium with each other, and indicates that no side reaction occurs during the photoisomerization process.\(^{28}\) As shown in Fig. 2(b), the cis-trans back isomerization was achieved through irradiating with visible light for 36 s. The reduction of content of the thermodynamically less stable cis form was confirmed by the intensity increase of the \(n\rightarrow \pi^*\) transition band around 363 nm and the intensity decrease of the \(n\rightarrow \pi^*\) transition around 453 nm. These results suggest that the azoester in polymer is able to undergo the photoisomerization in response to UV and visible light.

The Mesomorphic and Thermodynamic Properties of PC11AE6

As shown in Fig. 3(a), because of the linear structure and flexible main chain, PC11AE6 had a lower glass transition temperature (\(T_g = 48 \, ^\circ\text{C}\)). The polymer PC11AE6 possessed mesophase transition at 83 \(^\circ\text{C}\) and reached isotropic state at about 182 \(^\circ\text{C}\). The POM results indicated the existence of birefringence as temperature decreased to 25 \(^\circ\text{C}\) (Fig. 3b). Thus, the mesophase temperature interval of PC11AE6 was at least from 25 \(^\circ\text{C}\) to 182 \(^\circ\text{C}\), which was wider than that of previously reported LLCP with azo in the side chains.\(^{11}\) The temperature-varied 1D-WAXD data are shown in Fig. 3(c) and Fig. S2 (in ESI). In the low-angle region, two couples of diffraction peaks existed in 1D-WAXD curves (marked as \(2\theta_1\) and \(2\theta_2\) from 80 \(^\circ\text{C}\) to 160 \(^\circ\text{C}\), \(2\theta_2\) and \(2\theta_0\) from 20 \(^\circ\text{C}\) to 80 \(^\circ\text{C}\)), with a ratio of scattering vector \(q\) (\(q = 4\pi\sin\theta/\lambda\), with \(\lambda\) the X-ray wavelength and 2\(\theta\) the scattering angle) of 1:2. This result indicates PC11AE6 formed two types of long-range ordered lamellar structures at different temperature ranges, with a \(d\)-spacing of \(-4.60 \, \text{nm} (2\theta_0 = 1.92^\circ)\) and \(-5.04 \, \text{nm} (2\theta_2 = 1.75^\circ)\) at high and low temperature, respectively.\(^{29}\) In the high-angle region, the scattering halo with the maximum at \(2\theta_0\) of \(-20.5^\circ\) was more diffusive when the temperature was above the phase transition temperature (about 80 \(^\circ\text{C}\)), suggesting the molecular packing was less ordered at high temperature.

The Spontaneous Orientation of PC11AE6

The preferential orientation of mesogens is of vital importance for LCPs to deform in the anisotropic manner under external stimuli.\(^{30-32}\) The spontaneous orientation of the LCPs should be the consequence of molecular cooperative effect of the rigid azoester mesogens. The flexible backbones and long spacers increase the molecular flexibility which facilitate the azoester mesogens to generate a highly ordered structure.\(^{11}\) To determine the spontaneous orientation in PC11AE6, 2D-WAXD characterization was used. As investigated in Figs. 4(a) and 4(b), the split diffractions in the low- and high-angle regions demonstrated spontaneous formation of uniform mesogen orientation. Besides, the orientation of mesogens in fibre was similar to that of film. The diffraction arcs in the low-angle region appeared on the equator line in Figs. 4(a) and 4(b), implying the formation of the smectic phase with normal directions of the smectic layer perpendicular to the fibre axial direction or the film plane. In the high-angle region, two diffraction arcs on the meridian demonstrated the smectic A phase with out-of-plane orientation of the mesogens. Orientation parameter of fibre (\(S = 0.65\)) measured by WAXD was larger than that of film (\(S = 0.57\)), which may result from induced orientation produced by main chain extending process under fibre drawing. The schematic diagram of mesogen orientation both in PC11AE6 fibre and film is shown in Fig. 4(c). Out-of-plane orientation of mesogens was maintained in radial direction for fibre or thickness direction for film. Furthermore, when it exceeded the phase transition temperature, the mesophase changed from smectic A to smectic C with a tilting angle of 53\(^\circ\), which remained out-of-plane orientation of mesogens (Figs. 53 and S4 in ESI). The zigzag tilting of mesogens was generated in the lamella of smectic C. These results indicate that the anisotropic structures in the PC11AE6 fibre and film were formed, which established an important foundation for their photoresponsive deformation.

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Photoinduced Reversible Deformation of PC11AE6

With the anisotropic structure, photoresponsive azoester-containing PC11AE6 exhibited reversible macroscopic deformation under illumination (Fig. 5). As shown in Fig. 5(a), upon UV irradiation (365 nm, 40 mW·cm\(^{-2}\)), the PC11AE6 films and fibres bent away from the light source to reach different bending angles. Thanks to the out-of-plane orientation of mesogens, the trans-cis photoisomerization caused the expansion on materials surfaces of PC11AE6, which led to the bending behaviors (Fig. 5a and Movies S1 and S2 in ESI). The responsiveness of films with different thicknesses was investigated, as shown in Fig. 5(a) and Fig. S5 (in ESI). It should be noted that the bending angle of the thinner film (10 μm, 68°) was larger than that of the thicker one (20 μm, 21°). This is probably because that trans-cis photoisomerization reactions of azoester occurred only at the surface region of the films due to the large extinction coefficient of azoester moieties at about 360 nm when the PC11AE6 films were irradiated with UV light.
Fig. 5 The reversible photo-induced deformation behaviors of PC11AE6 films and fibres upon irradiation with 365 nm UV light and 530 nm visible light at room temperature. (a) The photodeformation of films and fibres upon irradiation from left. The size of films used here is 5 mm × 1 mm × 10 μm, while the diameter for fibres is 40 μm. The scale bar is 1 mm. (b) The relationship between bending angle and irradiation time of PC11AE6 films (10 μm) and fibres (40 μm) upon UV irradiation with different intensities. The visible light intensity was always kept at 50 mW·cm⁻².

Fig. 6 The infrared imageries of the reversible photo-induced deformation of PC11AE6 films at about 60 °C. (a) The initial state of films at about 60 °C in oven. (b) The bending of films under UV irradiation from right to left. The UV light intensity was always kept at 40 mW·cm⁻². The size of films used here is 5 mm × 1 mm × 10 μm. The scale bar is 2 mm. The transient temperatures of films marked by a white cross were recorded and showed in the upper left corner of imageries.

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from one side. The resistance existing intrinsically in the thicker film was larger than that of the thinner one, which brought small bending angle under the same irradiation condition. However, fibre with a diameter of 40 μm exhibited better photoresponsibility compared with film with a diameter of 10 μm under the same irradiation condition because of the increased order parameter (Movie S3 in ESI). Besides, as shown in Fig. S5(b) and Fig. S6 (in ESI), with UV light intensity increasing, films and fibres showed larger bending angles, which was caused by the more trans-cis photoisomerization driven by high photon energy. The average angular speed was used to quantify the photodeformation velocity. An average angle speed of 1.25 (°)·s⁻¹ was achieved for film and 3.67 (°)·s⁻¹ was reached for fibre under the same irradiation. After visible light irradiation with constant intensity (50 mW·cm⁻²), the bent PC11AE6 films and fibres recovered in 30 s, indicating that the deformation of PC11AE6 films was completely reversible.

The PC11AE6 film was used to investigate the photoinduced deformation of PC11AE6 at an elevated temperature, as shown in Fig. 6. When the temperature was around 60 °C, which was in the temperature range of smectic A phase, film bent away from the light source in 5 s under UV irradiation, and then it recovered immediately after removing the UV light (Movie S4 in ESI). This spontaneously recovering phenomenon may be caused by the on-going cis-trans back isomerization which was accelerated by heating because of the thermodynamic instability of cis form.

CONCLUSIONS

We designed an azoester-containing linear liquid crystal polymer PC11AE6 with excellent mesophase stability and photoresponsivity. The polymer exhibited wide mesophase temperature range, which possessed two mesophases in the process of heating or cooling and a high isotropic temperature (182 °C). With out-of-plane orientation of mesogenic, the PC11AE6 films bent away from the UV source and returned to the original shape with a higher order parameter (Movie S3 in ESI). Besides, the resistance existing intrinsically in the thicker film was larger than that of the thinner one, which brought small bending angle under the same irradiation condition. However, fibre with a diameter of 40 μm exhibited better photoresponsibility compared with film with a diameter of 10 μm under the same irradiation condition because of the increased order parameter (Movie S3 in ESI). Besides, as shown in Fig. S5(b) and Fig. S6 (in ESI), with UV light intensity increasing, films and fibres showed larger bending angles, which was caused by the more trans-cis photoisomerization driven by high photon energy. The average angular speed was used to quantify the photodeformation velocity. An average angle speed of 1.25 (°)·s⁻¹ was achieved for film and 3.67 (°)·s⁻¹ was reached for fibre under the same irradiation. After visible light irradiation with constant intensity (50 mW·cm⁻²), the bent PC11AE6 films and fibres recovered in 30 s, indicating that the deformation of PC11AE6 films was completely reversible.

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Electronic Supplementary Information

Electronic supplementary information (ESI) is available free of charge in the online version of this article at http://dx.doi.org/10.1007/s10118-020-2383-0.

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