Effect of the Concentration Gradient on Molecular Alignment by Scanning Wave Photopolymerization

Yoshiaki Kobayashi1,2, Ryo Taguchi1,2, Norihisa Akamatsu1,2, and Atsushi Shishido1,2*

1 Laboratory for Chemistry and Life Science, Institute of Innovative Research, Tokyo Institute of Technology, 4259 Nagatsutacho, Midori-ku, Yokohama 226-8503, Japan
2 Department of Chemical Science and Engineering, School of Materials and Chemical Technology, Tokyo Institute of Technology, 2-12-1 Ookayama, Meguro-ku, Tokyo 152-8552, Japan
*ashishid@res.titech.ac.jp

Macroscopic and precise alignment control of functional molecules can provide increased functionality and enhanced device performance. However, even the most powerful current method has some limitations, such as the need for polarized light and photoresponsive dye molecules. We reported an alignment process based on a new concept of scanning wave photopolymerization (SWaP) that achieves an arbitrary alignment pattern by spatiotemporal light-triggered molecular diffusion. Here, we investigate the effect of crosslinker on molecular alignment by SWaP. We found that the controlled concentration gradient of polymer added with the optimal amount of crosslinker improved the degree of alignment of the liquid-crystalline polymers. SWaP could be employed as a simple and versatile fabrication process for preparing highly functional materials and devices that require alignment control.

Keywords: Liquid crystal, Molecular alignment, Polymer, Photopolymerization

1. Introduction
Controlling the alignment of various functional molecules such as liquid crystals (LCs) is key for generating a new function and enhancing their performances [1-3]. Therefore, molecular alignment methods play an important role in the development and the production of electronic, photonic, mechanical and biomedical organic devices [2-5]. Although mechanical methods are simple and practical processes for one-dimensional alignment over large areas, alignment patterning is difficult. Furthermore, the mechanical methods cause alignment defects due to dust contamination and static electricity [6].

As an alternative method for overcoming these difficulties, photoalignment method using axis-selective photoresponsive dye molecules has been developed [7-10]. A representative photoalignment method uses the photoisomerization of azobenzene, which was first reported by Ichimura and Seki et al [11]. Such significant success of a photoreactive alignment layer led to development of photochemical alignment control, referred to as a “command surfaces”. The photoalignment method also was applied to photoresponsive polymer systems such as side-chain LC polymers, supramolecules and block copolymers. However, there still remain some challenges, namely the requirement of polarized light and photoresponsive molecules to control the alignment. In addition to such necessity, the alignment processes become complicated in the induction of complex alignment patterns.

Recently, we reported a dye-free photoalignment process based on a new concept of scanning wave photopolymerization (SWaP) [12-18]. In this method, localized photopolymerization generates a concentration gradient of polymer between irradiated and unirradiated regions, resulting in molecular diffusion [19-21]. The diffusion induces the mass flow, and then the flow aligns LCs at the boundary of both regions. Therefore, this method has far fewer limitations on applicable molecular structures and can provide a complex alignment pattern in a single step. For further development of SWaP as a versatile alignment method, to
investigate the relation between the concentration gradient of polymer and the induction of molecular alignment is important. In this study, we fabricated LC films with different composition ratios of a monomer and a crosslinker, and then examined their molecular alignment behavior. As a result, we revealed that the SWaP fabricated the highly oriented LC film under the optimal condition.

2. Experimental

The chemical structures of materials used in this study are shown in Fig. 1. The photopolymerizable anisotropic monomer, 4-(4-propylecyclohexyl)phenyl-4-{4-[(3-ethyloxetan-3-yl)methoxy]butoxy}benzoate (OXT-M) was employed as previously reported [13]. The isotropic crosslinker, 3-ethyl-3\{[(3-ethyloxetan-3-yl)methoxy]methyl\}oxetane (OXT-221) was provided by TOAGOSEI Co., Ltd., Tokyo, Japan. The cationic photoinitiator, CPI-210S was provided by San-Apro Ltd., Tokyo, Japan. The surfactant, MEGAFAC R-40 were provided by DIC Corporation Tokyo, Japan. Propyleneglycol monomethyl ether acetate (PGMEA) was purchased from Tokyo Chemical Industry Co., Ltd., Tokyo, Japan.

Fig. 1. Chemical structures used in this study.

The mixtures with 99:1, 97:3, and 95:5 molar ratio of OXT-M:OXT-221 (OXT\textsubscript{99:1}, OXT\textsubscript{97:3}, OXT\textsubscript{95:5}), were prepared by adding CPI-210S to the mixtures at 5 wt%. We then prepared the solutions of 40 wt\% by adding PGMEA, and doped R-40 at 1000 ppm. The solutions were stirred at 70 °C for 10 min, and were spin-coated (MS-A100, MIKASA, Co., Ltd., Tokyo, Japan; spin rate, 800 rpm) for 30 s on an ozone-treated glass substrate. The solvent in the coated mixtures was removed at 120 °C for 2 min. Before photopolymerization, the substrate was cooled to 100 °C for each mixture, which was their optimized photopolymerization temperature. To induce 1D molecular alignment by photopolymerization, the mixture on the substrate was subsequently irradiated with ultraviolet (UV) light at wavelength of 365 nm through a 500-µm-slitted mask from a high-pressure mercury lamp (OPM2-502HQ, USHIO INC., Tokyo, Japan) with glass filters (IRA-25S, UV-D33S, AGC Techno Glass, Co., Ltd., Shizuoka, Japan). The light intensity and the scanning rate were optimized to be 3.2 mW/cm\(^2\) and 20 µm/s, respectively (Fig. 2). Finally, the samples were irradiated with UV light (wavelength: 365 nm, light intensity: 10 mW/cm\(^2\)) for 10 min to complete photopolymerization without slit to fix the molecular alignment.

3. Results and discussion

The molecular alignment of the obtained LC polymer films was evaluated with a polarized optical microscope (POM, BX50, Olympus, Tokyo, Japan). The molecular alignment direction was confirmed by polarized UV-visible (vis) absorption spectroscopy. The order parameter (\(S\)), which quantitatively showed the degree of the in-plane alignment of mesogens, was calculated from the absorption bands corresponding to phenyl benzoate moieties (302–307 nm) by using the following equation:

\[
S = \frac{A_\parallel - A_\perp}{A_\parallel + 2A_\perp}
\]  

where \(A_\parallel\) and \(A_\perp\) are defined as the absorbance in which the direction of polarized incident light is parallel and perpendicular to the light scanning direction, respectively.
perpendicular. We measured birefringences of OXT99:1 and OXT97:3 films and then found to be 0.01 and 0.05, which were determined by POM observation with a Berek compensator. On the other hand, under crossed polarizers, POM image of OXT95:5 film always showed darkfield even though the stage of POM was rotated by 360° (Fig. 3f). To further investigate the molecular alignment behavior, we measured the polarized UV-vis absorption spectra and evaluated polar plots of these films. In OXT99:1 and OXT97:3 films, the absorption spectra showed a clear difference; the absorbance parallel to the light scanning direction was larger compared to the perpendicular direction (Figs. 4a-d). Furthermore, the $S$ values were found to be 0.19 and 0.26, respectively. This result indicated that the phenyl benzoate moieties in the OXT99:1 and OXT97:3 films were in-plane aligned along the light scanning direction. On the contrary, we observed the coincident spectra parallel and perpendicular to the light scanning direction in the OXT95:5 film, indicating that mesogens in the OXT95:5 was not aligned (Figs. 4e, f). These results revealed that addition of the appropriate amount of crosslinker is necessary to induce unidirectional molecular alignment by SWaP.

The alignment behavior in OXT99:1, OXT97:3 and OXT95:5 films is rationalized by the molecular alignment mechanism of SWaP. In SWaP, the spatiotemporal UV light irradiation generates local photopolymerization, which causes the spatial polymer concentration gradient between irradiated and unirradiated regions. The gradient induces the molecular diffusion. Such diffusion propagates along the direction of scanning light, triggering mass flow. The flow could impose shear stress on polymers, and subsequently align their main chains along the flow direction. Therefore, the concentration gradient is key to the induction of molecular alignment. The number of polymerizable groups in monomer or crosslinker in the systems determines the polymerization rate [20]. Therefore, the increase of the number of polymerizable groups enhances the polymerization rate, that is, the spatial polymer concentration gradient is steep. The OXT95:5 mixture with a large amount of crosslinker has faster polymerization, and thus finish the polymerization before the molecular alignment is induced. As a result, the alignment in OXT95:5 film became random. On the other hand, the OXT99:1 mixture with a small amount of crosslinker decreases the reactivity, and generates the spatial polymer concentration gradient with a gentle slope, leading to the small degree of alignment in OXT99:1 film. As a result, the alignment behavior depends on the amount of crosslinker due to the created polymer concentration gradient.
4. Conclusion
In summary, we fabricated LC films with various composition ratios of anisotropic monomer and isotropic crosslinker by SWaP, and investigated the effect of the polymer concentration gradient on molecular alignment. SWaP achieved the unidirectional molecular alignment in the LC films with 1 and 3 mol% of crosslinker, and the film containing 3 mol% of crosslinker showed the highest degree of alignment. On the other hand, the LC film with 5 mol% of crosslinker was not aligned. SWaP improves the degree of alignment of LC polymers with the controlled polymer concentration gradient, which is determined by the reactivity of polymerization. We believe that SWaP provides a powerful new method for fabricating highly oriented functional materials and devices.

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