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http://dx.doi.org/10.5772/36784

1. Introduction

Originally there are three states of matter; solid, liquid and gas. The emergence of an exotic and extraordinary form of matter, which is known as liquid crystal has been considered as one of the major breakthrough in polymer science. Liquid crystal can be defined as an intermediate of solid (crystal) and liquid (Knight & Vollrath, 2002) where the molecules have the capabilities to flow like a liquid (mobility) as well as possessing the common property associated to solid, which is the degree of order (Doldeny & Alder, 1998). In addition, liquid crystal materials are self assembling by nature and can offer a very elegant and effective way of controlling and tuning the physical properties that ultimately define the self-organizing and self assembly process (Zhang et.al., 2008). One of the exciting developments involving this unique material is the introduction of liquid crystalline behavior in polyurethane elastomers (PUE) where the first of this kind was synthesized by Iimura in 1981 (Lin et.al., 2001).

Polyurethane [PU] is one of the most versatile class of polymeric materials known today. Wide variety of structural changes can be produced with the different hydroxyl compounds and isocyanates leading to a wide spectrum of properties and applications (Yeganeh et.al., 2007). It contains a high concentration of polar groups, in particular the urethane group, resulting from isocyanate-hydroxyl reactions. The interactions between these polar entities are of great importance in determining the properties of PU of all types (Lee et.al., 1999) besides the composition and characteristic of the polyol, diisocyanates and the additives utilized during the synthesis (Pachecho et.al., 2009).

High toughness, excellent wear and tear properties and good oil resistance are among the advantages displayed by PUE (Wright & Cumming, 1969). Moreover, not only they have good mechanical and physical properties, PUE are also benefited with biocompatibility
characteristics for biomedical applications (Barikani et.al., 2009). Despite all the great aforementioned properties, modifications and improvements are done to conventional PUE in order to meet the qualities in more advanced applications.

Diisocyanates, polyol and low molecular weight diamine or diol (chain extender) are the basic building blocks of conventional PUE (Yeganeh & Mehdizadeh, 2009). In order to synthesize liquid crystal polyurethane elastomers (LCPUE), the low molecular weight diamine or diol used in conventional PUE was substituted with the mesogenic unit. Incorporation of geometrically anisotropic moieties (mesogenic unit) within polymer architecture can drive the formation of liquid crystalline phase from strictly steric repulsion considerations (Abe & Ballauf, 1991; Rowan & Mather, 2007). Furthermore, the insertion of mesogenic unit in the backbone of PUE will impart unique physical properties to the polymer and also improve its mechanical, optical and electrical characteristics (Jia et.al., 1996).

Various mesomorphic behaviors are exhibited with different types of mesogenic units in preparation of LCPUE. In this research work, mesogens consumed were thiazolo [5,4d] thiazoles based and it is known as an important class of bicyclic aromatic molecule comprising two fused thiazole rings (Knighton et.al., 2010). Thiazolo-thiazole rigid fused ring can enhance the rigidity of the polymer and the conjugation (Osaka et.al, 2007) which makes it a best candidate to be part of the hard segment in the LCPUE network. The hard segments consisted of either 2,5-bis(4-hydroxyphenyl) thiazolo-[5,4d] thiazole or 2,5-bis(4-hydroxy-3-methoxyphenyl) thiazolo [5,4d] thiazole and 4,4'- methylene diphenyl diisocyanate (MDI). As for the soft segments, polyethylene glycol (PEG) 1000, 2000 and 3000 were involved.

The ultimate aim of this work is to synthesize new LCPUE with the presence of thiazolo-[5,4d] thiazole as a chain extender. Study and analysis were carried out to determine the effects and consequences of the introduction of thiazolo-[5,4d] thiazole moiety and the influence of various lengths of polyols on the properties of LCPUE.

2. Experimental

2.1. Materials

Vanillin and 4, 4'-methylene diphenyl diisocyanate (MDI) were purchased from Aldrich Co. (United States). Rubeanic Acid (dithiooxiamide) and 4-hydroxybenzaldehyde were obtained from MERCK Co. (Germany). Polyethylene glycol with molecular weight of 3000, 2000 and 1000 (PEG: Mn= 3000, 2000 and 1000) were purchased from Fluka Chemica (Switzerland). All the chemicals were utilized as received without any further purification. N,N-Dimethylformamide purchased from Aldrich (United States) was distilled over Calcium Hydride (CaH₂) through vacuum distillation before being used.

2.2. Synthesis of monomers and polymers

2.2.1. Synthesis of 2,5-bis(4-hydroxyphenyl)thiazolo-[5,4d] thiazole (I)

Briefly, 3 g (25 mmol) of dithiooxiamide (Rubeanic acid) and 15 g (123 mmol) of 4-hydroxybenzaldehyde with the presence of 9 g (97 mmol) of phenol were charged all at once
in a 500 ml round bottom flask fitted with condenser and left to be refluxed for 2h. Precipitates were obtained by pouring the hot mixtures to the cold water. Subsequently, the yield was filtered off and washed with ethanol followed by ether. The product obtained was dried at 70°C in a vacuum oven for 24 hours. Recrystallization from cyclohexanone was performed giving an orange-yellowish powder. Yield: 35% with melting point 364°C. Fourier transform infrared (FTIR; KBr, cm⁻¹): 3492 (-OH), 1606 (C=N), 1596 (C=C), 855 (p-substituted benzene), 1H-NMR (400 MHz, DMSO-d₆ ppm): δH 7.12 (m, aromatic protons), 9.8 (s, -OH). Elemental analysis: Found: C, 59.16; H, 3.28; N, 8.84, C₁₆H₁₀N₂O₂S₂ Calc.: C, 58.89; H, 3.09, N, 8.59.

2.2.2. Synthesis of 2,5-bis(4-hydroxy-3-methoxyphenyl) thiazolo [5,4d] thiazole (II)

The same procedure was applied to the synthesis of 2,5-bis(4-hydroxy-3-methoxyphenyl) thiazolo [5,4d] thiazole except that 4-hydroxybenzaldehyde was substituted with vanillin. Orange-yellowish powder was obtained as the end product. Yield: 26% with the melting point of 259°C. Fourier transform infrared (FT-IR; KBr disc): 3534 cm⁻¹ (OH), 1608 cm⁻¹ (C=N), 1510 cm⁻¹ (C=C), 842 cm⁻¹ (-CH out of plane), 1H-NMR (400 MHz, DMSO-d₆ ppm): δH 7.09 (m, aromatic protons), 9.5 (s, -OH) and 3.87 (s, OCH₃). Elemental analysis: Found: C, 55.60; H, 4.03; N, 6.89, C₁₈H₁₄N₂O₄S₂ Calc.: C, 55.95; H, 3.62, N, 7.25.

2.2.3. Synthesis of liquid crystalline polyurethane elastomers (LCPUE)

Preparation of LCPUE was achieved by two steps solution polymerization reaction, where isocyanate terminated pre-polymer was synthesized initially in the first stage. To produce pre-polymer, 0.01 mol of PEG (Mw = 1000, 2000, and 3000) and 0.02 mol of MDI were mixed in 500ml of reactor flask equipped with condenser, thermometer, nitrogen inlet and mechanical stirrer. The mixture was allowed to be stirred and heated for 4h at 70°C in the presence of 15 ml of DMF as solvent and nitrogen gas was kept flowing to provide inert atmosphere. The reaction was followed by chain extension process, using either compound (I) or (II), where the chain extender was added dropwise within 1h to complete the formation of LCPUE. Subsequently, the temperature was increased to 100°C and the reaction continued for another 9 hr. The hot viscous solution was then poured into 200ml of cold water for precipitation, before subjected to filtration. Later, the filtered product was washed with ethanol several times and finally with ether, before being dried overnight in a vacuum oven at 60°C.

2.4. Measurements

100mg mixture of samples and KBr (grounded) were pressed into translucent disc before being subjected to Nicolar Avatar Model 360 Fourier Transform infrared spectrometer devices to obtain FT-IR spectra. Data was collected in the range of 4000-400cm⁻¹. 1H-NMR and 13C-NMR spectra were obtained using Bruker 400 MHz NMR spectrometer consuming DMSO-d₆ as solvent and TMS as internal standard. Thermal stability of LCPUE was determined by thermogravimetric analyzer (Perkin Elmer Pyris series 6) under nitrogen purge and with 10°C/min of heating rate and the heating was done up to 800°C. Liquid crystalline behavior was verified by means of differential scanning calorimetry (DSC) to
observe the behavior of polymers such as glass transition point (T_g), melting point (T_m) and isotropic temperature (T_i). It was conducted utilizing Perkin Elmer Pyris Series 7 thermal analyzer under Nitrogen flux at 10°C/min rate of heating. Textures of mesomorphic phases were displayed by Nikon Eclipse E600 polarized microscope equipped with MS600 Linkam Hot stage and SONY CCD-IRIS Color Video Camera. The heating rate was 5°C/min and 10°C/min for the cooling rate. Sample was placed between two thin round glasses and it was then transferred onto microscope fitted with the hot stage to be analyzed. Siemens X-ray Diffractometer model D5000 equipped with a CuKα target at 40KV and 40mA was used in obtaining X-ray scattering curve. Tensile strain properties of LCPUE films were measured by Instron Testing instrument at a constant speed of 500mm/min (speed) where the measurements were performed at room temperature. Brookfield viscometer was used to measure the fluid viscosity where suitable spindle and speed were chosen and it was also performed at room temperature.

3. Results and discussion

3.1. Preparation of chain extender

The preparation of 2,5-bis(4-hydroxyphenyl)thiazolo-[5,4-d] thiazole and 2,5-bis(4-hydroxy-3-methoxyphenyl) thiazolo [5,4-d] thiazole were conducted according to the reaction shown in Scheme 1. The starting reagent involved for the synthesis of both the compounds were rubeanic acid and either 4-hydroxybenzaldehyde or vanillin with the presence of phenol. Subsequently, both chain extenders prepared were being used in the preparation of LCPUE. Identification of the chemical structures of the aforementioned products was monitored primarily with FT-IR spectroscopy and further confirmation was carried out by 1H-NMR spectrophotometer.

Scheme 1. Preparation of Compound I and II
3.2. Polymer synthesis

LCPUE based on thiazolo [5,4d] thiazoles moiety were synthesized from long chain of diol (PEG 3000, 2000 and 1000) with an excess of disiocyanate (MDI) via addition reaction to give the terminal reactive group which results in the formation of ‘extended diisocyanate’ or isocyanate pre-polymer. Then, 2,5-bis(4-hydroxyphenyl) thiazolo-[5,4d] thiazole [I] and 2,5-bis(4-hydroxy-3- methoxyphenyl) thiazolo-[5,4d] thiazole [II] were added acting as a chain extender in order to convert the pre-polymer into long chain LCPUE. The general route for the preparation of LCPUE was outlined in Scheme 2, yield and viscosity of LCPUE were listed in Table 1 and the data showed that the range of the viscosities and yields obtained were 10,744 to 40 692 cP and 76-87 %, respectively. Range of the viscosities obtained also provides the information of the molecular weight of each polymer synthesized where high value of viscosity indicates high molecular weight of the polymer produced and vice versa (Bagheri & Pourmoazzen, 2008). In this case, all LCPUE samples displayed fairly high molecular weight in accordance with the results demonstrated.

![Scheme 2. General route for the preparation of LCPUE VI (a-c) and VII (a-c)](image)

3.3. Structural elucidation

FT-IR was employed to verify functional groups of the pre-polymer, compound I and II, and LCPUE. Prior to the formation of LCPUE which is referring to the pre-polymer state, in the region of 2270 cm\(^{-1}\) a peak was observed which was assigned to \(-\text{N}=\text{C}=\text{O}-\)
Polyurethane (diisocyanate) whereas according to the IR spectra of compound I and II, a peak was found at 3492 cm$^{-1}$ and 3334 cm$^{-1}$ which corresponds to –OH functional group in the chemical structure. The disappearance of both the bands of -N=O- in pre-polymer and –OH of compound I and II, indicates the completion of the reaction of preparation of LCPUE and this fact was also supported with the appearance of new absorption bands at 3356.84 cm$^{-1}$ (N-H stretching) and 1782.5 cm$^{-1}$ (carbonyl group) which were attributed to the urethane linkage, –NHCOO- (Zhang et al., 2008; Issam, 2007). Furthermore, the peak at 2884.89 cm$^{-1}$ was ascribed to –CH stretching, whereas the band representing C=C aromatic can be found at 1598.59 cm$^{-1}$. Figure 1 displayed the FTI-R spectrum of LCPUE VIIa and based on the results obtained, the characteristic absorption bands of FT-IR spectra for the other LCPUE were almost identical to one another. The fact that differentiates LCPUE VI and LCPUE VII was the presence of the methoxy group and it was proven in the FTIR spectrum of LCPUE VIIa, where a peak displayed at the region of 1024.27 cm$^{-1}$ corresponded to the methoxy group.

Further confirmation of chemical composition of LCPUE produced was carried out by means of Nuclear Magnetic Resonance spectroscopy (NMR). $^1$H-NMR spectrum of LCPUE VIIa was illustrated in Figure 2. A singlet peak centered at 8.76 ppm was assigned to –NHCOO- and this proved the formation of urethane linkage. The appearance of multiplet peaks at 7.53-6.99 ppm and singlet peak at 3.87 ppm was attributed to the aromatic protons and the protons in methoxy group, respectively. Aliphatic chain of polyol (PEG 1000) was detected in the region of 1.23-1.64 ppm.
Other than FT-IR and $^1$H-NMR analysis, $^{13}$C-NMR was performed in order to clarify the structure of LCPUE prepared. $^{13}$C-NMR spectra portrayed in Figure 3 which represents LCPUE VIIa shows that the formation of urethane linkage (NHC0O) was determined by the
appearance of the peak at 173.4 ppm. The methylene group presence in the soft segment of PEG can be seen as a sharp and intense peak at 25-29 ppm. More peaks can be observed at 117.8 to 158.7 ppm and 56.2 ppm where they were assigned to the aromatic carbons and the carbon in methoxy group respectively. Significant peaks in all characterization analysis (FT-IR, \textsuperscript{1}H-NMR and \textsuperscript{13}C-NMR) were consistent and adequately provide the evidences to support the fact that the reaction of all materials took place and LCPUE was successfully prepared.

3.4. Thermal and liquid crystalline behavior of polymers

The DSC analysis was conducted at a heating rate of 10°C to understand phase separation behavior of all synthesized LPCUE where the transition occurs, observed under polarizing optical microscope (POM) equipped with heating stage and the results obtained from both measurements were listed in Table 1. Based on the DSC thermograms, upon heating, one step transition and two endothermic peaks were detected where each of them indicates glass transition (T\textsubscript{g}), melting endotherm, (T\textsubscript{m}) and isotropic endotherm (T\textsubscript{i}) respectively, which is also the evidence of the existence of mesophase. LCPUE derived from 2,5-bis(4-hydroxy-3-methoxyphenyl) thiazolo [5,4d] thiazole have transition temperatures lower than those derived from 2,5-bis(4-hydroxyphenyl)thiazolo-[5,4d]thiazole. Methoxy group, which acts as a substituent attached to the phenyl ring has the capability to lower the melting and isotropization temperature and caused thermal suppression of the molecule to occur (Al-Dujaili et.al., 2001). The fact was supported by the results illustrated in Fig.4 where it depicts the DSC thermograms of LCPUE. LCPUE VIIa displayed melting point (T\textsubscript{m}) at 164°C and isotropization temperature (T\textsubscript{i}) at 187°C whereas for LCPUE VIa, T\textsubscript{m} was detected at 176°C and T\textsubscript{i} at 205°C. The substituent could also act to reduce the coplanarity of adjacent mesogenic groups and increase the diameter or decrease the axial ratio of the mesogens [Li and Chang, 1991]. Due to the higher range between T\textsubscript{m} and T\textsubscript{i} of LCPUE VIa, the thermal properties of this polymer are higher and more stable compared to LCPUE VIIa. The types of diisocyanates also contribute to the thermal behavior of LCPUE, where MDI based PU was known for having better order of the rigid chain that approaches the decomposition temperature, giving high melting point to the polymer produced (Jieh & Chou, 1996). As for glass transition, it involves mobility of the chain segments and the T\textsubscript{g} will be affected by the mobility restriction on the chain segments, (Suresh et.al., 2008) it therefore explains the varying pattern of the T\textsubscript{g} values displayed in Table 1. The decreasing values of T\textsubscript{g} can be observed as the length of soft segments increases, indicating that the long chain of polyol gave great flexibility characteristics towards the polymer chains where less mobility restrictions occurred and hence resulting in the lower T\textsubscript{g} values.

POM was utilized to investigate the type of mesophase by displaying the phase transition that occurred, subsequently providing the polarizing optical microphotographs of the target compounds. The morphology observed on heating and transition temperatures obtained were given in Figures 5 and 6 and the results were summarized in Table 1. It was revealed that all LCPUE showed mesophases upon melting temperature where the thread texture of the nematic phases can be seen. From the photographs taken by POM, the crystal to mesophase transition occurred at temperature ranging from 129 to 181°C. The samples were
further heated after the crystal-nematic transition temperature, and resulted in the disappearing of the texture when reaching the isotropization stage. There were no traces of mesophase transition during the cooling process from POM indicating all samples possessed thermotropic type of liquid crystal. Phase transition temperatures observed through POM were found to be consistent with the corresponding DSC thermograms.

Table 1. Thermal properties of LCPUE VI (a-c) and LCPUE VII (a-c) by DSC and POM
Figure 5. Polarized optical images of (a) LCPUE VIa (181°C), (b) LCPUE VIb (162°C) and (c) LCPUE VIc (133°C)

Figure 6. Polarized optical images of (a) LCPUE VIIa (170°C), (b) LCPUE VIIb (148°C) and (c) LCPUE VIIc (129°C)

X-ray diffraction analysis of LCPUE was conducted at room temperature to obtain information on both the mesophase structure and crystallinity of LCPUE. The measurements exhibited several peaks in the range of 2θ = 15 – 25° as observed in Figure 7.
and this indicated semi crystalline character possessed by LCPUE. The results obtained in above range also provide details related to the d-spacing of 3.56 and 4.92 Å, thus supporting the characteristic of nematic liquid crystalline phase (Jeh & The, 1994) as displayed through POM.

Figure 7. X-ray diffraction scales of LCPUE VI (a-c) and LCPUE VII (a-c)

Thermal stability of prepared LCPUE was investigated by thermogravimetric analysis (TGA). Incorporation of liquid crystalline properties into the polymer structure would enhance the thermal properties (Jahromi et al., 1994) and the theory has proved to be applicable from the results obtained. This may be partly due to favorable interactions between hard domain interface and the liquid crystalline phase. All synthesized LCPUE possessed good thermal stabilities, however, PU elastomers eventually undergo thermal degradation when exposed to high temperatures. Degradation process occurred in two step pattern where the initial degradation occurs in the hard segment involving the urethane linkages, while the second stage indicated the degradation of soft segments. TGA curves in Figure 8 demonstrated the thermal degradation of all LCPUE prepared where 10% weight loss of LCPUE occurred at about 315-341°C and the maximum degradation temperature was in the range of 430-470°C, signifying a high thermal stability property. Furthermore, it can be observed that LCPUE VIIc demonstrated the lowest degradation temperature among the others and this proved that the length of polyethylene glycol (soft segment) influenced the thermal stability of LCPUE where the order of LCPUE due to their thermal stability can be arranged as LCPUE VIa>VIIa>VIb>VIIb>VIIc>VIIc.
3.5. Tensile properties

Table 2 demonstrates tensile properties of the synthesized LCPUE. As seen, all of the polymers possessed good elastic properties with high elongation at break. Due to the data listed, the higher the molecular weight of the soft segments, the greater the elongation at break, but decrease of tensile strength and tensile modulus can be observed. When the molecular weight of polyol increased, the number of urethane groups in the polyol chain was reduced at the same time, and hence the number of rigid segments is lower, consequently, the possible number of intermolecular hydrogen bonds goes down in which –NH and C=O groups are active (Kro & Pitera, 2008). However, the presence of enhanced rigid and high aspect ratio mesogenic unit as part of hard segment in the synthesized LCPUE, is able to give both high strength and good elastic properties to LCPUE even with long soft segments, which

<table>
<thead>
<tr>
<th>Sample</th>
<th>Tensile modulus (Mpa)</th>
<th>Tensile strength (Mpa)</th>
<th>Elongation at break (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCPUE VIa</td>
<td>17.1</td>
<td>28.2</td>
<td>290</td>
</tr>
<tr>
<td>LCPUE VIb</td>
<td>13.4</td>
<td>24.1</td>
<td>450</td>
</tr>
<tr>
<td>LCPUE VIc</td>
<td>11.5</td>
<td>19.9</td>
<td>570</td>
</tr>
<tr>
<td>LCPUE VIIa</td>
<td>17.5</td>
<td>28.3</td>
<td>330</td>
</tr>
<tr>
<td>LCPUE VIIb</td>
<td>13.7</td>
<td>24.3</td>
<td>460</td>
</tr>
<tr>
<td>LCPUE VIIc</td>
<td>11.2</td>
<td>19.8</td>
<td>560</td>
</tr>
</tbody>
</table>

Table 2. Mechanical properties of LCPUE VI (a-c) and LCPUE VII (a-c)
is unusual in conventional PUE (Jeong et al., 2000). Better phase separation will lead to good mechanical properties; hence the introduction of the mesogens unit as chain extender into LCPUE can be said to easily induce the matter (phase separation) to occur.

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Acknowledgement
The author would like to thank University Sains Malaysia for short term grant no.304.PTEKIND.6311031 and the fellowship scheme for funding the research.

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