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Synthesis of Hydroxylated Polyisoprene-Graft-Polylactide Copolymer
(Sintesis Kopolimer Poliisoprena Terhidroksil-Cangkuk-Polilaktida)

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ABSTRACT
Polyisoprene (PI) has been widely used in many industries for decades. Many researches have reported that most significant weaknesses of polyisoprene are caused by unsaturated double bond C=C. The aim of this research was to synthesis and characterize a new copolymer utilizing the unsaturated double bond C=C of polyisoprene. PI is first modified to form hydroxylated polyisoprene (PIOH). The absence of alkene proton peak in NMR spectrum of PIOH is a strong evidence that the unsaturation of PI has been reduced. After that, PIOH is subjected as an initiator for the ring-opening polymerization of D,L-lactide in bulk condition to form hydroxylated polyisoprene-graft-polylactide copolymer (PIOH-g-PLA). The NMR spectrum of the new copolymer structure showed an unique peak at 4.09 ppm corresponding to methine proton of polyisoprene backbone adjacent to the PLA chains, indicating the grafting of D,L-lactide is successful to form PIOH-g-PLA. The average molecular weight, M_n of PIOH-g-PLA was significantly increased compared to PIOH, from 38260 to 36870 according to GPC. The surface of PIOH-g-PLA displayed significantly higher wettability and hydrophilicity than polyisoprene with water contact angle of below 30°. This owes to the terminal hydroxyl groups of PLA chains that lead to the formation of hydrogen bonds. Thermal stability studies by TGA and DTG of PIOH-g-PLA indicated two thermal degradations at T_max 260 and 392 ℃ corresponding to PLA side chains and PIOH backbone, respectively, with PIOH exhibiting highest thermal stability compared to PI and the graft copolymer.

Keywords: Graft copolymer; hydroxylated polyisoprene; polylactide; ring opening polymerization

INTRODUCTION
Natural rubber (NR) is an irreplaceable resource in modern day production industry. NR is used in manufacturing of more than 40,000 products which more than 400 are medical devices (Kind & Hull 2012; Nampoothiri et al. 2010). The usage of NR has been steadily increasing throughout the years, from 7.32 million tons in 2010 to 14.01 million tons in 2018 worldwide (MRB 2018).

The need of NR steadily increases due to the good properties of the resulting material, such as strong elasticity, high molecular weight, high tensile strength, renewable, impact resistant, and abrasion resistant (Anon.
NR is purified to obtain polyisoprene (PI) which is the particle of natural rubber that provides all these properties. PI is hard to be mimicked by synthetic materials.

Modification of PI has been conducted for more than 50 years, and the common modifications are hydroxylation (Azhar et al. 2017), chlorination (Paoprasert & Chanroj 2017), dehydrogenation, microwave technique (Idris et al. 2019), grafting (Wongthong et al. 2013), oxidation (Zahari et al. 2018a), vulcanization, self-assembly polymerization (Caldas et al. 2020), blending (Ahmad et al. 2015; Hisham et al. 2011), epoxidation (Azhar et al. 2017), mastication, hydrogenation (Azhar et al. 2016; Yakubchik et al. 1962) and cyclization (Swartling et al. 2018). Most modifications focus on the unsaturation of PI. One of the significant weakness of PI is the unsaturated double bond C=C that causes PI to be weak against weathering and heat. PI is also readily decomposed upon heating due to random chain scission. In some cases, researchers utilize the unsaturation of PI. In recent research, polyisoprene was blended with carbon black and low-density polyethylene to form shape memory composite by utilizing the unsaturated double bond C=C of polyisoprene and polyethylene to form specific crosslink region and crystallized region (Xia et al. 2019).

As many modifications have been done over the years, most modifications focus on reducing the unsaturation of the double bond C=C in PI, which is one of the factors of low heat resistance of PI. In many previous studies, synthesized copolymers show the properties of both polymers, which further improve the material in being more competent in the industry. Thus, the aim of this research was to synthesize a copolymer using polyisoprene and other eco-friendly material, which is D,L-lactide to form a grafted copolymer. The graft polymerization will take place at the unsaturated double bond C=C, thus reducing the unsaturation of PI and producing a competent grafted copolymer.

Polyisoprene is used as the backbone chain of the copolymer and D,L-lactide is the grafting monomer. During graft polymerization, D,L-lactide is polymerized and forms poly-D,L-lactide (PDLLA). PDLLA is a widely researched material due to the properties of biodegradable, biocompatible, and renewable. This research is envisaged to achieve grafted copolymer of hydroxylated polyisoprene-graft-poly(lactide) (PIOH-g-PLA). Referring to some past research, it is expected that the synthesis of PIOH-g-PLA is successful (Azhar et al. 2017, 2016) and it is expected to inherit both PI and PLA’s properties. The graft copolymerization is carried out by melt ring-opening polymerization of cyclic D,L-lactide using hydroxylated polyisoprene (PIOH) initiator with the presence of Sn(Oct)2, catalyst. Scheme 1 shows the reaction scheme of PIOH-g-PLA synthesis. PIOH-g-PLA is characterized in terms of structure, thermal properties, surface properties and molecular weight.

**SCHEME 1. Reaction scheme of PIOH-g-PLA synthesis**

**MATERIALS AND METHODS**

Hydrogen peroxide, H2O2, AR grade and methanol, CH3OH, AR grade was purchased from R&M Chemicals. Sodium tungstate dihydrate, H2Na2O6W (≥99%) was purchased from Acros Organics (New Jersey, USA). Acetic acid, CH3COOH (≥ 99 %) was purchased from J.T Baker. Polyisoprene, C5H8, sodium carbonate, Na2CO3 (≥ 99 %), AR grade, D,L-lactide, C6H8O4 (99 %) and tin(II) 2-ethylhexanoate, C16H30O4Sn (92.5 - 100 %) was purchased from Sigma Alrich (Germany). Toluene, C7H8, AR grade was purchased from Chemiz.
PREPARATION OF PIOH
PIOH was synthesized following a previously reported method (Azhar et al. 2017), using epoxidation and oxidation via sodium tungstate complex. The polyisoprene samples were subjected to epoxidation and hydroxylation according to the previous outlined procedure with some modifications. Modifications are the use of polyisoprene instead of liquid natural rubber (LNR) solution and of the amount used. The amount used was modified from 17.0 g of LNR solution to 3.3 g of PI.

PREPARATION OF PIOH-GRAFT-PLA
The graft copolymer of PIOH-g-PLA was synthesized via ring opening polymerization of D,L-lactide in bulk using PI-OH as initiator and Sn(Oct)2 as catalyst under nitrogen gas flow. PIOH (1 g) and D,L-lactide (1 g) were added into a Schlenk flask and the flask was heated at 120 °C in a silicone oil bath, followed by addition of Sn(Oct)2 (41 µL) into the flask. The reaction was kept at 120 °C and constantly stirred for 6 h. The flask was then left to cool down. The graft copolymer was transferred from the flask into a vial for characterization.

CHARACTERIZATION
PI, PIOH, and PIOH-g-PLA were each diluted in CDCl3 (15 mg in 460 μL) and analyzed by using Nuclear Magnetic Resonance Spectroscopy (1H and 13C NMR) (JEOL, JNM-ECP 400, Japan) to identify the proton and carbon present after polymerization. Samples of 5 mg in 5 mL THF were used for Gel Permeable Chromatography (GPC) (Waters/1515 Isocratic HPLC Pump equipped with a Waters 2414 Refractive Index detector, Waters Corporation, USA) with styragel type column, THF as solvent and polystyrene as calibrant with flow rate of 1 mL min⁻¹ at 40 °C to determine number average molecular weight (Mn), weight average molecular weight, (Mw) and polydispersity index (PDI). Thermogravimetric Analysis (TGA) (Mettler Toledo, TGA/SDTA 851e) was used to determine the thermal stability of the prepolymers and graft copolymer using aluminum crucible, heated from 30 to 600 °C at the heating rate of 10 °C min⁻¹ with constant flow of nitrogen gas. Static contact angle test was used to test the wettability of the surface of PIOH-g-PLA via film formation. A thin film of 10 wt. % PIOH-g-PLA diluted in CHCl3 (1 mL) were casted on glass and let dry for 3 days before static contact angle test (Contact Angle LSE-B100W).

RESULTS AND DISCUSSION
HYDROXYLATED POLYISOPRENE-GRAFT-POLYLACTIDE
Ring opening polymerization in bulk was employed in the preparation of PIOH-g-PLA. The catalyst, Sn(Oct)2 was added into the flask after both PIOH polymer D,L-lactide monomer melted at 120 °C. The addition of the catalyst initiates the ring opening polymerization of D,L-lactide to form poly-D,L-lactide (PLA). During theROP of lactide, the first ring opened lactide is inserted between the oxygen and hydrogen atoms of the hydroxyl groups in PIOH repeating unit as shown in Scheme 2. This process repeats as subsequent lactides are inserted at the terminal hydroxyl to form PLA as the side chain of PIOH. This process is a grafting-from method and is a living polymerization.

SCHEME 2. Grafting “from” living polymerization of PIOH-g-PLA graft copolymer
MICROSTRUCTURE OF PIOH-GRAFT-PLA

Nuclear magnetic resonance spectroscopy (NMR) was used to determine the proton and carbon present in the microstructure of the synthesized graft copolymer. Figure 1 shows the NMR spectrum (1H NMR) of the synthesized PIOH-g-PLA, whereas Table 1 shows the results extracted from Figure 1. As can be seen in Table 1, there are some important peaks that signifies the success of PIOH-g-PLA synthesis. The signal at chemical shift of 4.09 ppm belongs to the methine proton at the polyisoprene backbone, and is adjacent to PLA side chain. This signal is important as this proton was originally an alkene proton and first appeared at chemical shift of 5.19 ppm in polyisoprene, which was determined by running 1H NMR on the prepolymer polyisoprene and compared with other references (Azhar et al. 2016, 2017; Gemmer & Golub 1979; Sato & Tanaka 1979; Tanaka & Sato 1976; Zell et al. 2002). After modifying polyisoprene to hydroxylated polyisoprene (PIOH), the proton shares its carbon with hydroxyl group and resonated at 3.42 ppm, which still can be seen in Figure 1. The changes of chemical shifts are due to the shared carbon’s deshielding effect towards the hydrogen in which this sudden change of environment affects the chemical shift (Azhar et al. 2017; Zell et al. 2002). The chemical shift of the proton in polyisoprene backbone becomes more deshielded when D,L-lactide inserted in between the hydrogen and oxygen atom of the hydroxyl group at PIOH, causing a change in distortion of electron cloud. The change of chemical shift of this proton shows that the hydroxylated polyisoprene has successfully grafted with polylactide. The signal at 5.18 ppm shows the methine proton of the polylactide repeating units (Pretula et al. 2016; Rasal et al. 2010; Thakur et al. 1997a, 1997b; Zhong et al. 2002). The proton at the unsaturated double bond C=C of polyisoprene starting material that initially observed at 5.05 ppm was not observed in the PIOH-g-PLA copolymer product spectrum (Figure 1), indicating that after hydroxylation of polyisoprene, the double bond conversion was nearly 100% as the alkene peak cannot be detected in the 1H NMR spectrum.

![Figure 1](image_url)

**Figure 1.** 1H NMR spectrum of PIOH-g-PLA (27 °C, 400 MHz, CDCl3 solvent)

**Table 1.** 1H NMR data for PIOH-g-PLA

| Labelling | Chemical shift (ppm) | Type of proton | Assignment |
|-----------|----------------------|----------------|------------|
| (a)       | 5.19                 | Methine proton of PLA repeating unit | -CH-       |
| (b)       | 4.09                 | Methine proton of PI backbone    | -CH-       |
| (c)       | 4.39                 | Methine proton of PLA terminating unit | -CH-       |
| (d)       | 3.91                 | Hydroxyl proton of OH           | -OH-       |
| (e)       | 1.85                 | Methylene proton of PI backbone | -CH2-      |
| (f)       | 1.58                 | Methyl group of PLA repeating unit | -CH3-      |
| (g)       | 1.26                 | Methyl group of PI backbone      | -CH3-      |
| (h)       | 1.49                 | Methyl group of PLA terminating unit | -CH3-      |
Figure 2 shows the $^{13}$C NMR spectrum of PIOH-g-PLA, whereas Table 2 shows the results extracted from Figure 2. The methine carbon at PI backbone appears at 129 ppm. Based on previous reports (Anancharoenwong 2011; Gemmer & Golub 1978; Tanaka & Sato 1976), all the peaks that appear on the $^{13}$C NMR spectrum of PIOH-g-PLA is accurate in comparison. The change in adjacent atom after the graft polymerization causes a change of the appearance of the methine carbon at PI backbone in the $^{13}$C NMR spectrum.

**FIGURE 2.** $^{13}$C NMR spectrum of carbon atoms in PIOH-g-PLA

**TABLE 2.** $^{13}$C NMR data for PIOH-g-PLA

| Labelling | Chemical shift (ppm) | Functional group | Type of carbon |
|-----------|----------------------|------------------|----------------|
|           | 76-78                | CDCl$_3$         |                |
| a         | 20                   | Methyl carbon of PLA repeating unit | -CH$_3$       |
| b         | 16                   | Methyl carbon of the PLA terminating unit | -CH$_3$       |
| c         | 26                   | Methyl carbon of PI backbone | -CH$_3$       |
| d         | 29                   | Methylene carbon of PI backbone | -CH$_2$       |
| e         | 31                   | Methylene carbon of PI backbone | -CH$_2$       |
| f         | 69                   | Chiral carbon of PLA repeating unit | -C=          |
| g         | 84                   | Chiral carbon of PLA terminating unit | -C=          |
| h         | 129                  | Methine carbon of PI backbone | -CH=         |
| i         | 138                  | Chiral carbon of PI backbone | -C=          |
| j         | 169                  | Ester carbon of PLA repeating unit | -C=O         |
| k         | 175                  | Ester carbon of PLA terminating unit | -C=O         |
THERMAL STABILITY OF PIOH-GRAFT-PLA

Thermogravimetric analysis (TGA) was used to determine the thermal stability of PIOH-g-PLA, PI and PIOH. Figure 3 shows the thermogram of PIOH-g-PLA, PI and PIOH whereas Figure 4 shows the DTG curve of PI, PIOH and PIOH-g-PLA. Table 3 shows the tabulated results of Figures 3 and 4. Based on Table 3, PI started to decompose at 352 °C (T_{onset}) and exhibited maximum decomposition temperature (T_{max}) at 382 °C, whereas PIOH showed degradation at T_{max} of 345 °C and T_{max} of 408 °C. PIOH-g-PLA exhibited two degradation stages, with T_{max} values of 260 and 392 °C, respectively. The prepolymer PI underwent complete thermal degradation while PIOH and PIOH-g-PLA have residues of 4.9 and 10.0%, due to the presence of tungsten and tin based catalysts used in the preparation of PIOH and PIOH-g-PLA, respectively. The thermal stability increased after modification from PI to PIOH due to the removal of the unsaturated double bond C=C via hydroxylation to form PIOH. For the graft copolymer PIOH-g-PLA, the overall thermal stability decreased compared to the intermediate prepolymer, PIOH. The first decomposition stage which belongs to polylactide graft segment whereas the second decomposition stage is the main polymer backbone. PLA has weaker thermal stability compared to PI, thus degrades earlier than PIOH backbone. PIOH-g-PLA which has 50:50 weight ratio of PIOH and PLA during synthesis, decreases in weight by 48.0% after 330 °C. The molecular weight and D,L-isomer ratio in PLA affects the heat resistance of PLA. Research has shown that L-lactide have a stronger thermal stability than D-lactide when heated between 300 and 400 °C (Feng et al. 2018; Pluta 2004; Schmidt & Hillmyer 2001). PLA can degrade via intra and intermolecular ester exchange resulting in the formation of CO_{2}, CO, acetaldehyde and methylketene (Kopinke et al. 1996). The polylactide in this research uses D,L-lactide as the monomer, thus having a lower thermal stability which result in a lower degradation temperature. Many researches have shown that the degradation temperature of PLA occurs at different temperature.

FIGURE 3. TGA thermograms of PI, PIOH and PIOH-g-PLA

FIGURE 4. DTG curves of PI, PIOH, and PIOH-g-PLA
TABLE 3. $T_{\text{max}}$, $T_{\text{onset}}$, $T_{\text{end}}$ and residue of PI, PIOH, and PIOH-g-PLA

| Sample            | $T_{\text{max}}$ (°C) | $T_{\text{onset}}$ (°C) | $T_{\text{end}}$ (°C) | Residue (%) |
|-------------------|------------------------|--------------------------|------------------------|-------------|
| PI                | 382                    | 352                      | 450                    | 2.6         |
| PIOH              | 408                    | 345                      | 451                    | 4.9         |
| PIOH-g-PLA        | 260                    | 392                      | 230                    | N/A         |

N/A = cannot be observed due to overlapping of first and second thermal degradation

MOLECULAR WEIGHT OF PIOH-GRAFT-PLA

Gel permeation chromatography (GPC) was used to determine the molecular weight of PIOH-g-PLA and the prepolymer. In terms of number average of molecular weight ($M_n$), weight average of molecular weight, ($M_w$) and polydispersity index (PDI), Table 4 shows the results of analysis. Based on the table, it can be concluded that the overall molecular weight of PIOH-g-PLA is higher than the prepolymer PIOH. Because of the growing of polylactide chains on PIOH, the $M_n$ of PIOH-g-PLA shows significant increase, from $M_n$ of 38260 to 56870. The increase of PDI is due to the limited control of the grafting mechanism. During the ring opening of D,L-lactide, two hydroxyl groups of PIOH readily to form bond with D,L-lactide. The carbon grafting sites, which one contains methyl group and the other contains hydrogen atom, causes steric hindrance towards D,L-lactide to graft onto PIOH. As the steric hindrance of the methyl group at PI backbone affects the grafting site of D,L-lactide, it also affects the grafting reactivity of D,L-lactide. This causes weight distribution along the chain to be broad and affects the PDI of PIOH-g-PLA copolymer. Besides that, as polysoprene is readily decompose upon heating, this causes a phenomenon known as random chain scission (Kind & Hull 2012) during copolymerization. Random chain scission starts to occur, targeting the ‘weaker link’. This weaker links is formed due to remainders in PI and PIOH such as initiator, or some head-head repeating units during the formation of long polysoprene chain via radical polymerization in natural latex (Gemmer & Golub 1978; Kind & Hull 2012; Zahari et al. 2018b). This further affects the $M_w$ of PIOH-g-PLA as during the synthesis of the graft copolymer, long duration of heat was applied. This is also one of the factors that causes the decrease in molecular weight from PI to PIOH.

TABLE 4. $M_n$, $M_w$, and PDI of PI, PIOH, and PIOH-g-PLA

| Polymer                              | Number average molecular weight ($M_n$) | Weight average molecular weight ($M_w$) | Polydispersity index (PDI) |
|--------------------------------------|----------------------------------------|----------------------------------------|---------------------------|
| Polysoprene (PI)                     | 4320                                   | 96220                                  | 22.3                      |
| Hydroxylated polysoprene (PIOH)      | 2580                                   | 38260                                  | 14.9                      |
| Hydroxylated polysoprene-graft-polylactide (PIOH-g-PLA) | 1410                                   | 56870                                  | 40.3                      |

SURFACE PROPERTIES OF PIOH-GRAFT-PLA

Static contact angle test was used to determine the surface properties of PIOH-g-PLA. The sample was prepared using 10 wt. % of polymer diluted in chloroform and was left to dry for 3 days before characterized. Figure 5 shows the static contact angle plot where y-axis is angle of the droplet (°) and x-axis is duration of the test (ms). According to Figure 5, there is a significant increase in wettablility of PIOH-g-PLA copolymer compared to PI. Although polysoprene and polylactide are hydrophobic
polymers, previous research showed that the properties of terminating functional group and the molecular weight of the polymer affects the hydrophilicity of the material (Liu et al. 2010). The addition of hydrophilic functional group can increase the hydrophilicity of the polymer and that the shorter the length of the polymer, the more it increases the hydrophilicity of the material (Hassan et al. 2016; Kasalkova et al. 2015; Liu et al. 2010; Tretinnikov & Ikada 1997). PI is hydrophobic due to the lack of functional groups to form hydrogen bonding with water. However, after modification into PIOH, the addition of hydroxyl groups increases the hydrophilicity of PIOH.

The PIOH-PLA copolymer is comprised of grafted PLA with hydroxyl group at one end, in addition to remaining hydroxylated polyisoprene backbone, further increases the amount of free carbonyl group and free hydroxyl group that can form hydrogen bonding with water. Therefore, the hydrophilicity of PIOH-g-PLA is high although both PI and PLA are hydrophobic material. The wettability of the surface of PIOH-g-PLA is high due to the addition of functional groups that can interact with water and form hydrogen bonding. Figure 6 shows the image obtained from static contact angle test at 0.5 and 25.5 s as a reference.

**FIGURE 5.** Static contact angle test plotted with angle (°) against time (ms)

**FIGURE 6.** Image obtained from static contact angle test at 0.5 and 25.5 s for PI, PIOH, and PIOH-PLA
CONCLUSION

The synthesis of PIOH-g-PLA was successful through the ring opening bulk polymerization of D,L-lactide with PIOH as initiator at reaction conditions of 120 °C and 6 h, yielding PIOH-g-PLA with interesting properties. The weight average molecular weight of PIOH-g-PLA was relatively high with values of 56870 due to the presence of PLA side chains, in comparison to 38260 for PIOH. The molecular weight distribution of PIOH-g-PLA was broad as the grafting mechanism was affected by the steric hindrance from the methyl group in PIOH and the occurrence of thermally induced random chain scission during copolymerization. The surface of PIOH-g-PLA shows hydrophilic properties, with contact angle of less than 30° which contradicts PLA hydrophobic characteristics. This was owed to the hydroxyl end group of PLA chains, allowing hydrogen bonding and improved surface wettability. The thermal stability of PIOH-g-PLA was relatively lower in comparison to PIOH and PI, with \( T_{\text{max}} \) of 260, 408, and 382 °C, respectively, due to the presence of PLA. Overall, the unsaturation in PI was successfully modified to obtain PIOH and PIOH-g-PLA with interesting properties. The unique advantage of PIOH-g-PLA synthesized using this method is that the material is solvent-free and using safe catalyst, making the copolymer potentially suitable in applications such as encapsulation material in drug delivery and as compatibilizer in biodegradable composites.

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