INVESTIGATION OF BIOPLASTIC PROPERTIES DEVELOPED FROM ACRYLATE EPOXIDIZED SOYBEAN OIL THROUGH RING OPENING POLYMERIZATION PROCESS

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ABSTRACT

Soybean oil is one of the major vegetable oils containing more than 99\% of triglycerides of saturated and unsaturated fatty acids and has become an interesting source to produce bioplastic. This study investigates the synthesis and characterization of bioplastic developed by the acrylate epoxidized soybean oil through ring opening polymerization process. The mechanical properties of the samples were characterized using Fourier Transform Infrared Spectroscopy, Thermogravimetric analysis and the physio-chemical properties of the bioplastics were studied. In this study, the bioplastic made up from soybean oil with the lowest glycerol concentration showed the best mechanical properties.

Keywords: Bioplastic, Acrylate epoxidized soybean oil, Biodegradable

1.0 INTRODUCTION

Plastics is one of the most highly valued materials mainly because they play an important role in human life such as food packaging and biomedical applications Piringer & Baner (2008) and Raheem(2012). The development of plastics is believed to have started around 1860 with an intention of developing (Browne, 1993) a new substitute for natural ivory. In addition, plastics have been one of the most highly valued materials mainly because of their extraordinary versatility and low cost (Murray, 2007). Plastics are everywhere in human’s life such as packaging, constructions and biomedical applications. Plastics are cheap with its wide ranges of usage yet it caused a huge environmental problem, because they do not degrade in landfills and cannot be composted (Hopewell et al, 2009). Conventional plastics made of petroleum and its aliened components have caused the depletion of natural resources and environmental issues as it does not degrade in landfill (Hopewell, et al 2009). Most of the conventional plastics are made from poly-olefins such as poly (propylene) (PP), poly (carbonate) (PC), poly (vinyl chloride) (PVC), poly (ethylene) (PE), poly (styrene) (PS) etc. These conventional plastics are made of petroleum and its allied components, these natural resources take millions of years to form and are finite in quantity. In addition, plastics derived from fossil resources are largely non-biodegradable. Thus, the depletion of petroleum resources and increasing environmental awareness and regulations have triggered for the development of next generation materials that are environment-friendly and available resourcefully to meet the increasing demand for plastics (Muniyappa et al, 1996). In addition, as a result of
advancing technology a lot of problems and damage has been done to the environment over the years, which has now become a huge concern. This threat to the ecosystem is extremely big due to which manufacturing industries, especially the packaging, construction, and automobile industries, are searching for new environmentally friendly biodegradable substitutes that can successfully replace their conventional synthetic non-biodegradable counterparts. One way to reduce the use of fully petroleum based plastic products, is to substitute the synthetic fibres to natural wood and agriculture fibres. Researchers have been conducting research on composites made from wood fibers, plastics and recycled plastics as well as hybrid composites. (Clemons 2002,. Ashori, 2008,. Rivai et al 2014). These kinds of composites have been found to exhibit improved properties such as mechanical, thermal, and morphological properties yet they are not fully bio based (Islam et al 2014,. Islam et al 2016). Hence, a lot of attention is now being paid on preparation and application of bio-based polymers because of environmental concerns (Mohanty & Misra (2002), Gandini(2008) and Raquez et al (2010).

Development of bioplastics which are made from renewable resources and biodegradable materials can reduce the depleting fossil fuels and environment impact. Bioplastics made from renewable resources are biobased, biodegradable or both. Biobased means the material or the product is partly derived from biomass (plants) while biodegradable means the materials can be converted into natural substances such as water and carbon dioxide by the chemical process of the micro-organisms in the environment. Today, bioplastics play an important role in various industry applications such as food packaging and composting bags. Vegetable oils are highly suitable starting materials for polymers due to their abundance, low cost, biodegradability, environmental benefits and variety of functional groups. They are triglycerides with different composition of fatty acids depending on the plant, crop, season, growing conditions and purification methods (Kim & Sharma, 2012). Some of the most commonly used vegetable oils to synthesis bioplastics included castor oil, palm oil, and sunflower oil. Triglyceride-based vegetable oils, such as soybean, jatropha, linseed, sunflower, palm, castor, nahar seed, and canola oil, are being considered as precursors in the production of polymers . Soybean oil is one of the major vegetable oils containing more than 99% of triglycerides of saturated and unsaturated fatty acids.

Soybean oil is available abundantly across the world and is the second largest crop in the United States of America. In addition, the unsaturated sites in soybean oil can be used to introduce new functional groups like epoxides. Epoxidised soybean oil can be used as a stabilizer and plasticizer in the polymers or as an additive in lubricants and can also be converted into polyls by ring opening of the oxirane groups, resulting in intermediates that are used in polyurethane (Kaitsimkul, 2008). Its sustainability, low price and ease of manufacture make this bio resin an attractive alternative to petroleum-based plastics. One of the problems of bioplastics made from triglyceride-based materials is that they exhibit low rigidity and strength, which limit their use for practical applications (Khot et al, 2001). To improve the properties of biodegradable plastics, various types of fillers have been studied which included inorganic fillers (eg: calcium carbonate, nanoclay), natural fibers (both wood and plant fibers) and other types of filler such as carbon nanotubes (Mohanty et al (2000), Andrew and Weisenberge (2015)).
New applications of AESO based polymers are also desirable because of their availability from renewable resources and biodegradability. In this study, ring-opening polymerization of AESO using glycerol as plasticizer (Luo et al., 2011) is reported. There are two (2) objectives of this research which are to synthesis bioplastics by ring opening polymerization of acrylated epoxidised soybean oil (AESO) with plasticizer and to characterize the properties of AESO bioplastics. The scope of this research was to study the performance of bioplastics developed by mixing of 10wt% to 20wt% of glycerol in AESO. Characterization of bioplastics was performed using FT-IR analysis, TGA, DSC analysis, XRD analysis and SEM analysis, and to run mechanical testing (tensile strength and Young’s modulus) on the synthesized bioplastics.

2.0 METHODS AND MATERIALS
Acrylate epoxidised soybean oil (AESO) with 4,000 ppm monomethyl ether hydroquinone as inhibitor was used as the base materials for the synthesis of bioplastics and it was supplied by Sigma-Aldrich Company. Glacial acetic acid (99.85%) is supplied from Chemmart Asia Sdn. Bhd. The catalyst used in this research is 4-dimethylaminopyridine, a derivative of pyridine with the chemical formula (CH3)2NC5H4N and it was supplied by Chemmart Asia Sdn. Glycerol ACS reagent, anhydrous, (99.5%) is also supplied from Chemmart Asia Sdn. Bhd.

Setup of Synthesis Ring Opening Polymerization
Different types of equipment were used in the lab for this research work. The ring opening polymerization system consisted of a reactor assembly designed to achieve better mixing and dispersion control of lactic acid and nano fillers into AESO. It includes three neck reaction pot made up of quartz glass, equipped with a dropping funnel, condenser, high stirring motor, thermometer, nitrogen gas supplies and an electro-thermal heater. An additional, opening has been provided on the base of the reaction vessel in order to collect pre-polymer and polymer solutions in due process of experiment. Few other apparatus and equipment are vacuum oven, vacuum pump, oven, temperature and speed controlled mixer, glass wares (beakers, measuring cylinders, and etc), disposable syringes and a fume hood with water. LA and 4-dimethylaminopyridine (DMAP) catalyst are dried under vacuum at 70˚C for 24 hours, AESO is dehydrated in a vacuum oven at 60˚C, MMT is dried and dehydrated in vacuum oven at 100˚C before use.

Sample Preparation
AESO was dehydrated in a vacuum oven at 60˚C before use. AESO was mixed with acetic acid (20 wt%) and glycerol (10 wt%) at around 120˚C for five (5) hours using mechanical stirrer. The mixture was degassed for five (5) minutes to remove gas bubbles produced in the process. Five (5) ml of mixture was poured into aluminium weight boat and cured in an oven at 110˚C for a week to obtain the AESO plastic film. An analogous procedure was used to prepare AESO plastic film containing 15.0 wt% and 20.0 wt% glycerol.

Fourier Transform Infrared Spectroscopy (FTIR)
FTIR analysis was performed using Fourier transform infrared Spectrometer (Perkin-Elmer: Model 1000 Series) instrument equipped with a universal attenuated total reflectance (UATR) accessory was used to analyze biocomposites and raw materials. The data suggested that spectra were recorded between 4000 cm⁻¹ and 500 cm⁻¹ frequency ranges.
**X-Ray Diffraction (XRD)**

XRD measurement was carried out by using a Shimadzu XRD 600 X-ray diffractometer with CuKα radiation (\(\lambda = 1.542 \text{ Å}\)) operated at 30 kV and 15 mA. Data were collected within the range of scattering angles (2\(\theta\)) of 5° to 50° at the rate of 0.02°/min. The basal spacing was derived from the peak position in the XRD diffractogram according to the Bragg equation (\(\lambda = 2d \sin \theta\)).

**Scanning Electron Microscopy (SEM)**

The surfaces of synthesized bioplastics film were studied under a Hitachi’s Tabletop Electron Microscope TM3030Plus at an accelerating voltage of 15 kV. SEM was used to identify the molecular structure of the sample to check if the mixtures are well-mixed during the polymerization process.

**Thermo gravimetric analysis (TGA)**

Experiments were carried out for performing TGA using a thermal analyzer Mettler Toledo Instruments under a nitrogen atmosphere. Each sample was heated from room temperature to 900°C at a rate of 10°C/minute in air with a flowrate of in general 50ml/minute. The weight change of each sample was recorded as a function of temperature. Then mass loss and differential thermal analysis profiles were recorded.

**Differential Scanning Calorimetry (DSC)**

DSC analysis was also performed by TA-Instrument DSC/Q 1000 to study the nonisothermal crystallization kinetics. The specimen weight was in general between the ranges of 3mg – 5mg and the heating rate employed 10°C per minute from 40°C to 350°C.

**Tensile Test**

Tensile properties were tested using Shimadzu AG-X series universal testing machine. The samples were cut into dumbbell shapes following ASTM D638 (type V) standard. A load of 5.0 kN was applied at constant crosshead speed of 1 mm/min at room temperature. The tensile strength and tensile modulus were evaluated from the force-displacement data. Each sample included three tested replicates to obtain a reliable mean and standard deviation.

**Statistical Analysis**

Each batch of samples was prepared with different composition of glycerol. Analysis of variance (ANOVA) was performed by using the Data Analysis tools in Microsoft Excel 2013, and a least significant difference (LSD) test was used to compare the means with a confidence interval of 95%.

**3.0 RESULTS AND DISCUSSION**

The spectrum obtained from the six (6) samples was compared to the existing known materials and identified the material’s component and the highest possible matched in each sample is ricinoleic acid ethyl ester.

**FTIR Analysis**

The results of FTIR analysis are displayed in Figure 1. The spectrum of cluster peaks below 3000 cm\(^{-1}\) down to around 2850 cm\(^{-1}\) are the peaks from the stretching of sp3-hybridised C-H bonds.
The alcohol O-H group peak is around or about 3300 cm\(^{-1}\). A sharp peak right around 1700 cm\(^{-1}\) could possibly mean that there is a carboxylic acids group in the mixture (Key, 2015).

![Figure 1: FT-IR graph of AESO-based bioplastics](image)

The criteria of the peaks meet the criteria of the existence of an ester group, which are carbonyl peak around 1740 cm\(^{-1}\), alky peaks below 3000 cm\(^{-1}\) and a very strong peak around 1200 cm\(^{-1}\) for esters which representing the C-O single bond vibration (Wang et al, 2008).

**XRD Analysis**

Figure 2, shows the graph of comparison of XRD analysis with different concentrations of glycerol. The graph demonstrates peaks which indicate the strong crystalline characteristics. The shift to higher angle indicates a decrease in the corresponding interlayer spacing, which means that the blend component have an ordered structure. The increase of d-spacing shows that this blend has a less ordered structure, thus crystallization becomes more difficult (Chieng et al, 2014).
Figure 2: XRD Graph of AESO-based Bioplastics

SEM Analysis
Figure 3A shows the images of SEM analysis of 10 wt% of glycerol, Figure 3B shows the SEM images at 15 wt% of glycerol and Figure 3C shows the SEM images at 20 wt% of glycerol. Figure 3A shows very good compatible morphologies without edge, cavity and holes. This phenomenon shows a good adhesion between the components with a diffused polymer-plasticizer interface, which is attributed to the occurrence of chemical interactions between AESO, glycerol and acetic acid (Fenollar, 2009). Thus, the glycerol was well-dispersed to form a homogeneous matrix with evident signs of plasticization in the AESO matrix, without separation at the interface producing single phase morphology (Silverajah et al, 2012).
Figure 3: SEM graphs of (A) 10 wt% glycerol AESO bioplastics film; (B) 15 wt% glycerol AESO bioplastics film; (C) 20 wt% glycerol AESO bioplastics film.

On the other hand, SEM analysis on 15 wt% glycerol AESO plastics films (Figure 3B) and 20 wt% glycerol AESO plastics films (Figure 3C) show undissolved particles and presence of microvoid. As compared to 15 wt% glycerol AESO plastics films (Figure 3B), 20 wt% glycerol AESO plastics films (Figure 3C) shows more voids and undissolved particles. Presence of empty microvoids may due to the degree of dispersion of the plasticizer in the polymer matrix is better at lower plasticizer content, the tendency to form empty voids and phase separation increases when the glycerol content increases (Sanyang et al, 2015).

TGA Analysis
Figure 4, shows the graph of weigh percentage across room temperature to 900°C obtained through TGA analysis. From Figure 4, it can be seen that the AESO bioplastics with different glycerol concentrations have the same trend of weight loss. These Tonset shown in Figure 4 are identical to the research finding where the degradation occurred around 360°C. Besides, Derivative Thermograms (DTG) was also plotted and shown in Figure 5. From Figure 5, it is shown that the maximum rate of oxidation happens at around 400°C.
Figure 4: TGA curves of different concentrations of glycerol

Pure Poly-Lactic Acid (PLA) plastics has Tonset of 274.26°C, with 5 wt% of epoxidised palm oil is 313.54°C, and with 2 wt% of epoxidised Soybean oil is 330.40°C (Fenollar et al, 2009). A different study shows that starch-based films have three (3) thermal degradation events where the initial stage started at temperature less than 100°C, the second stage happened ~125°C to 290°C and the third staged happened at heating beyond 290°C with the Tonset occurred around 300°C (Mulla, 2011).

Figure 5: Derivative Thermograms (DTG) of different glycerol concentrations

Therefore, as compared to other biodegradable plastics, AESO bioplastics with glycerol as plasticizer have better thermal stabilities. The AESO bioplastics were observed thermally stable at temperature below 350°C. Theses higher degradation temperature as
compared to other biodegradable plastics may due to the increase molecular weight of interaction between AESO and glycerol or AESO matrix itself. Besides, the presence of glycerol which are small molecules that dispersed homogeneously in the PLA polymer acts as a barrier sheet to prevent oxidation, prevent the permeability of volatile degradation out from the blend materials and helps delay the thermal degradation process (Yang et al., 2015). It was reported in a study that the increase of glycerol concentration significantly increase the thermal degradation rate at constant temperature (Mulla, 2011). However, in the present study, increase glycerol concentration has an insignificant effect on the thermal stability. Thermal degradation rate as well as thermal stability which was found similar to the this study (Zhang et al., 2014). This study reported that TGA curves trend similar regardless of the composition of other components.

**DSC Analysis**
The thermal properties of AESO bioplastics with different concentrations of glycerol were studies by DSC in range from room temperature to 350°C as shown in Figure 6.

![DSC analysis for different glycerol concentrations](image)

The DSC curves show that the melting temperature is around 41°C. Scala and Wool (2005), reported that glass transition temperature, Tg of triglycerides with acrylate functionality (AESO, for example) increases approximately linearly (~50 to 92°C) with cross-linking density induced thermally. The Tg of different glycerol concentration is around 40°C which is identical to the pure AESO composite at 40.50°C (Yang et al., 2015). By definition, Tg is the temperature at which the forces binding the molecules in the matrix are relaxed to allow large-scale molecular movement (Wittaya, 2013). The addition of plasticizer into the plastic film reduces the Tg, but in (Mulla, 2011) the decrease of Tg values are
insignificant when the plasticizer concentrations increases which is similar to the current study.

**Tensile Tests**

*Tensile strength*

Figure 7, shows the graph of tensile strength versus concentration of glycerol. Pure AESO has a tensile strength of 4.90 MPa it is shown that with the increases of concentration of plasticizer, the tensile strength decreases. Tensile strength of 15 wt% of glycerol and 20 wt% has insignificant difference (Suyatma, 2005). As compared to starch-based bioplastics, AESO bioplastics have relatively lower tensile strength which the tensile strength value of 15 wt% of glycerol-plasticized sugar palm starch film is 9.59 MPa while AESO film is just 0.898 MPa.

![Graph of tensile strength](image)

**Figure 7:** Graph of tensile strength of different concentrations of glycerol

When compared to epoxidized soybean oil-plasticized PLA films, PLA films have much higher tensile strength of 32 MPa. Tensile strengths of AESO-based plastics films generally have a lower tensile strength values as compared to other biodegradable plastics such as starch-based and PLA-based plastics film.

*Young’s Modulus*

Figure 8, shows the graph of Young’s modulus versus concentration of glycerol. It is shown that with the increases of glycerol concentration, the Young’s modulus decreases. Tensile strength of 15 wt% of glycerol and 20 wt% has insignificant difference. As compared to starch-based bioplastics, AESO bioplastics have relatively lower tensile strength which the tensile strength value of 15 wt% of glycerol-plasticized sugar palm starch film is 9.59 MPa while AESO film is just 0.898 MPa. When compared to epoxidized Soybean oil-plasticized PLA films, PLA films have much higher tensile strength of 841 MPa. The effect of plasticizer concentration on the Young’s modulus was
observed. Young’s modulus of the bioplastic decreased with 15% glycerol concentration compared to 10% glycerol concentration. However, it increased with the 20% glycerol concentration, this was found similar to the graph of elongation at break with glycerol-plasticized starch plastics. Young’s modulus is the function of stress and strain where strain is the function of elongation at break, therefore it is validated that the Young’s modulus has an inverse behavior. Young’s modulus of AESO-based plastics films generally has a lower Young’s modulus values as compared to other biodegradable plastics such as starch-based and PLA-based plastics film.

![Graph of Young’s modulus of different concentrations of glycerol](image)

**Figure 8:** Graph of Young’s modulus of different concentrations of glycerol

### 4.0 CONCLUSIONS

Bioplastics were synthesized successfully through ring opening polymerization using acetic acid to form polyols and glycerol as the plasticizer. AESO bioplastics synthesized were made up from ricinoleic acid ethyl ester. Glycerol was found to be compatible with AESO, bioplastics film surfaces were studied using SEM and showed no hole at 5 wt% glycerol AESO bioplastics. Thermal stability of AESO bioplastics was observed better than other biodegradable plastics. It can be concluded that they were thermally stable below 350°C but showed insignificant effect with various compositions of glycerol. This study suggests that tensile strength and Young’s modulus values decreased with the increase of glycerol.

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