EFFECT OF ELECTRON BEAM IRRADIATION ON THE TENSILE PROPERTIES OF OIL PALM MESOCARP FIBRE/POLY(BUTYLENE SUCCINATE) BIOCOMPOSITES

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ABSTRACT

The present work deals with the utilization of oil palm mesocarp fibre (OPMF) and poly(butylene succinate) (PBS) to produce cost-effective biodegradable materials. The biocomposite was prepared from OPMF and PBS at a weight ratio of 70:30 via a melt blending technique without addition of any additive. This biocomposite showed relatively low tensile and water resistance properties. As a consequence, it was subjected to electron beam irradiation (EBI) treatment under various irradiation dosages ranging from 5–25 kGy, aiming at improving both its tensile properties and its water resistance. The results indicated that the biocomposite irradiated with 25 kGy of applied dosage showed a considerable improvement of 50% in tensile strength, 780% in tensile modulus and 38% in elongation at break. The water absorption and thickness swelling of the biocomposites were reduced after EBI treatment. The scanning electron microscopy showed improvement of the interfacial adhesion between OPMF and PBS after EBI treatment.

Keywords: Biocomposites; electron beam irradiation; poly(butylene succinate); oil palm mesocarp fibre

INTRODUCTION

Recently, natural fibre-filled polymer biocomposites have been increasingly utilized as an alternative to glass fibre-filled polymer composites in the automotive and construction industries due to their eco-friendliness, light weight and energy-saving characteristics [1]. Malaysia is one of the major oil palm plantation countries in the world, hence a huge amount of biomass is generated after the production of palm oil. Statistical analysis showed that approximately 4 kg of dry biomass is generated with every kilogram of palm oil produced. Oil palm mesocarp fibre (OPMF) is one of the
types of biomass generated in the palm oil mill after the extraction of oil from the oil palm fruit [2]. In 2010, the available amount of OPMF in Malaysia was reported to be 10.80 Mt [3]. Currently, it is either used in mulching or incinerated to generate electricity for the mill, but both these methods are detrimental to the environment, so a more sustainable method is required. Therefore, utilization of OPMF as a filler for biocomposites fabrication may play a key role in reducing the problem and at the same time produce a value-added product. Poly(butylene succinate) (PBS) is a commercially available biodegradable polymer produced from monomers of 1,4-butanediol and succinic acid via the polycondensation process [2]. Traditionally, those monomers are derived from a petroleum-based system, but nowadays they can be produced from renewable resources via a bacterial fermentation route [3]. This means that PBS has received equal attention to that given to other biodegradable polymers such as poly(lactic acid) (PLA) and poly(hydroxyl butyrate) (PHB), which are also derived from renewable resources. In comparison to PLA and PHB, PBS is relatively ductile and flexible in nature, with low rigidity, which has made it more suitable for various applications [4-8].

The biodegradability, renewability, availability and low cost of natural fibre have made it an attractive filler for biocomposites fabrication [9, 10]. However, its high moisture adsorption, relatively poor wettability and compatibility with some polymeric matrices have restricted many of its applications [6, 11-15]. Therefore, various treatment methods such as silane, alkali, peroxide, and isocyanate are employed to modify the surface of natural fibres [11]. These treatments are less environmentally friendly as they always involve the use of solvents and wastes are generated at the end of the process. Additionally, the solvents and chemicals used are expensive. Therefore, it would be desirable if an efficient and more environmentally friendly treatment process could be made available for improving the fibre/polymer interfacial adhesion. Recently, the electron beam irradiation (EBI) technique has been increasingly utilized for surface-modification and properties enhancement of various polymer materials like fibres, films and composites, as it is a dry, clean and cold process that is energy-saving, high speed and eco-friendly [16]. According to Han et al. [16], EBI treatment of natural fibre surfaces at certain dosages can significantly contribute to the improvement of the interfacial and mechanical properties as well as the thermal stability of natural fibre/polymer biocomposites. Therefore, there is no doubt that EBI treatment can be used in this work to enhance the properties of OPMF/PBS biocomposite. This paper highlights the effect of EBI treatment on the tensile and water absorption behaviours of OPMF/PBS biocomposite without the presence of any additive. The adhesion between OPMF and PBS was studied using scanning electron microscopy.

**EXPERIMENTAL SET-UP**

**Materials**

Poly(butylene succinate) (PBS), under the commercial name of BIONOLLE 1903MD, was purchased from Showa Denko, Japan. It has a density of 1.26 g/cm³ and a melting point of ~115 °C. Its molecular structure is shown in Figure 1. OPMF was collected from FELDA Serting Hilir Oil Palm Mill, Malaysia. It was first soaked in distilled water for 24 hours, then rinsed with hot water (60 °C) and acetone prior to drying at 60 °C in an oven. This process was carried out to remove waxes and impurities from the fibre.
surface. The dried fibre was then ground, sieved into sizes of 150–300 µm and stored in a sealed polyethylene bag for use in biocomposite fabrication.

![Molecular structure of PBS](image)

Figure 1. Molecular structure of PBS.

**Fabrication of OPMF/PBS Biocomposite**

Before processing, both PBS and OPMF were oven dried at 60 °C. The OPMF/PBS biocomposite was fabricated by melt blending of PBS and OPMF in a Brabender internal mixer at 120 °C with 50 rpm rotor speed for 15 minutes. The weight ratio of OPMF/PBS was fixed at 70:30. PBS was first loaded into the heating chamber for 2 min to melt. After that, OPMF was added slowly into the heating chamber and mixing continued for another 13 min. This compounded material was then compressed into a 1 mm sheet by a hydraulic hot press also at 120 °C under a pressure of 150 kg/cm² for 5 min, followed by cold pressing at 30 °C for 5 min.

**Electron Beam Irradiation**

Electron beam irradiation treatment was carried out using an electron beam machine, EPS Model-3000. Before treatment, the previously compressed sample sheets were put in a polyethylene bag and vacuum-sealed. The samples were then placed on an aluminium tray and irradiated at room temperature within a dosage range of 5–25 kGy at 5 kGy/pass with accelerator energy of 1 MeV, beam current of 2 mA and conveyor speed of 0.94 m/min. One-sided irradiation was employed for all the samples.

**Tensile Properties Measurement**

Tensile testing was carried out using a Universal Testing Machine, Instron 4302. The specimens for the tensile test were cut according to ASTM D638-5. The tensile test was performed at 25 °C under a load cell of 1 kN and crosshead speed of 5 mm/min. The results were expressed in terms of tensile strength, tensile modulus and elongation at break. Five specimens were tested for each group and their average values were reported.

**Scanning Electron Microscopy**

The scanning electron micrographs of the tensile fracture surfaces of biocomposites were captured using a JEOL JSM-6400 scanning electron microscope operated at 15 kV. The scanning electron micrographs were captured at a magnification of 200x. All the samples were coated with gold by a Bio-rad coating system before viewing to avoid charging of samples.
Water Absorption Test

A water absorption test was conducted according to ASTM D570. The samples were cut to dimensions of 10 x 10 x 1 mm³ to perform the test. Prior to the water absorption test, the samples were oven dried at 60 °C until constant weight was obtained and denoted as $W_0$. The samples were then immersed in distilled water at 25 °C. The samples were removed from the distilled water periodically, wiped with tissue paper to remove excess water on their surface and weighed immediately to obtain the wet weight ($W_t$). After the weighing process, the samples were returned into the distilled water. This process was repeated up to 168 hours. Three specimens were tested for each set of experiments and their average values were reported. The water uptake was calculated based on Eq. (1):

$$\text{Water uptake (\%)} = \frac{W_t - W_0}{W_0} \times 100\%$$

Thickness Swelling Test

The thickness swelling test was conducted according to the European Standard EN 317 (1993). Samples with dimensions of 10 x 10 x 1 mm³ were cut and used to carry out the test. The samples were oven dried at 60 °C prior to analysis. The thickness of the samples was measured before immersion in distilled water for 24 hour at 25 °C. This initial thickness was denoted as $T_0$. After that, the samples were removed from the distilled water and wiped with tissue paper to remove excess water on their surface. The thickness of the wet samples ($T_{24h}$) was measured immediately. Three specimens were tested for each set of experiments and their average values were reported. The thickness swelling was calculated based on Eq. (2):

$$\text{Thickness swelling (\%)} = \frac{T_{24h} - T_0}{T_0} \times 100\%$$

RESULTS AND DISCUSSION

In previous work [17], we have reported the maximum amount of OPMF that can be incorporated into the biocomposite of OPMF/PBS to be 70 wt%. Therefore, this ratio was chosen in this study to evaluate the effect of EBI treatment on the tensile and water resistance properties of this biocomposite without applying any additive.

Tensile Properties

The tensile test is carried out to study the effectiveness of EBI treatment in tuning the tensile properties of OPMF/PBS biocomposite. The tensile strength, tensile modulus and elongation at break of OPMF/PBS biocomposite as a function of EBI dosages are illustrated in Figures 2, 3 and 4, respectively. The tensile strength of non-irradiated OPMF/PBS biocomposite shows a value of 13.86 MPa (Figure 2). This value increased gradually with increasing EBI dosages. The maximum increment is observed at an EBI dosage of 25 kGy, which shows a value of 21.50 MPa or an improvement of 50% relative to non-irradiated OPMF/PBS biocomposite. During EBI treatment, several reactions may occur in the polymer microstructure, such as chain-scission, cross-linking,
and recombination of broken chains. Additionally, dislocating or abstracting of hydrogen atoms from the polymer chain may also happen [18]. The sites where hydrogen has been abstracted can instantly be joined with adjacent molecular chains to form strong bonds. According to Suhartini et al. [19], PBS does not cross-link within the EBI dosage range of this study. Therefore, the enhancement in tensile strength of OPMF/PBS biocomposite is mainly due to the formation of chemical bonds between the polymer matrix and fibre [20].

![Figure 2. Tensile strength of OPMF/PBS biocomposites at different irradiation dosages.](image)

![Figure 3. Tensile modulus of OPMF/PBS biocomposites at different irradiation dosages.](image)

An increasing trend is also observed for the tensile modulus upon EBI treatment, as shown in Figure 3. The tensile modulus of OPMF/PBS biocomposite increases gradually with increasing EBI dosages of up to 15 kGy. A drastic increment in tensile modulus is observed at an EBI dosage of 20 kGy. Beyond that (25 kGy), the increment is rather small. At low EBI dosages (5–15 kGy), fewer radicals are generated due to insufficient energy, hence a limited reaction occurs. However, once a suitable dosage (20 kGy) is applied, a reaction may occur rapidly owing to the presence of a large
amount of free radicals. After that, it reaches a plateau region (25 kGy) before degradation of chains starts to occur at higher EBI dosages. Figure 3 shows that the tensile modulus of non-irradiated OPMF/PBS biocomposite is 94.80 MPa and it has increased to a value of 843.10 MPa after EBI treatment at 25 kGy, showing an improvement of 780%. Previous studies by Ahmad et al. [21] also reported an increase in tensile modulus with increasing irradiation dosages and a decrease at higher irradiation dosages. It is interesting to note that the elongation at break of the irradiated biocomposite is enhanced by 38% upon EBI treatment at 25 kGy, as shown in Figure 4. This observation can probably be attributed to the better bridging formed at the interfacial region, by either a van der Waals interaction or a covalent bond between the PBS and OPMF after irradiation. The improved interfacial adhesion between OPMF and PBS (clearly shown in Figure 5(b)) will then prolong the deformation of the biocomposite, resulting in a higher value of elongation at break.

Figure 4. Elongation at break of OPMF/PBS biocomposites at different irradiation dosages.

**Surface Morphology**

SEM analysis is performed on the tensile fracture surfaces for direct observation of the biocomposite structure, particularly to examine the degree of adhesion at the fibre/polymer interface region. The SEM micrographs of non-irradiated and irradiated OPMF/PBS biocomposites are shown in Figure 5. The SEM micrograph of non-irradiated OPMF/PBS biocomposite (Figure 5(a)) shows a two-phase system, with numerous voids present on the fracture surface due to the detachment of OPMF from the PBS matrix. Additionally, visible gaps can also be seen at the interface of the OPMF and PBS, resulting from poor interfacial adhesion between the hydrophilic OPMF and hydrophobic PBS. It is noted that the surface of the OPMF is relatively clean with no PBS adhering on it during fibre pull-out, indicating that their bonding is relatively poor. An obvious morphological change is observed upon EBI treatment at 25 kGy (Figure 5(b)). The interfacial adhesion between OPMF and PBS appears to be improved after EBI treatment. This is clearly manifested by the presence of fewer voids and the disappearance of visible gaps between the OPMF and PBS. In addition, the PBS also seems to be adhered on the surface of the OPMF during fibre pull-out (white circle in
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Figure 5(b)), indicating that their bonding strength is relatively stronger. This result coincides with the results for tensile strength, tensile modulus and elongation at break of biocomposites, which show improvement after EBI treatment.

![SEM micrographs of (a) non-irradiated OPMF/PBS and (b) irradiated OPMF/PBS at 25 kGy.](image)

**Figure 5.** SEM micrographs of (a) non-irradiated OPMF/PBS and (b) irradiated OPMF/PBS at 25 kGy.

**Water Absorption**

Water absorption is one of the major concerns in polymer/fibre biocomposite for industrial application, since natural fibre is hydrophilic in nature and very sensitive to moisture. For this reason, a water absorption test is carried out to study the effect of EBI treatment on the water absorption behaviour of the OPMF/PBS biocomposite. The water absorption of non-irradiated and irradiated OPMF/PBS biocomposites as a function of immersion time in water is graphically displayed in Figure 6. Both biocomposites show an increase in water absorption with prolonged immersion time in water. The increase in water absorption is probably due to the hydrophilic nature of OPMF, which contributes to water absorption [22, 23]. Non-irradiated OPMF/PBS biocomposite shows higher
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water absorption than that of irradiated OPMF/PBS biocomposites. This is because the gaps present at the interface of OPMF/PBS biocomposite may also act as a pathway to water absorption [24]. The water absorption of OPMF/PBS biocomposite at 168 hours is reduced from 25.41 to 14.61% after EBI treatment at 25 kGy, showing a 43% reduction. The improvement in interfacial adhesion between the OPMF and PBS may be the reason for the decrease in water absorption after EBI treatment. A similar observation has also been reported by other researchers [25, 26].

![Figure 6. Water absorption of non-irradiated and irradiated OPMF/PBS biocomposites at different immersion times.](image)

![Figure 7. Thickness swelling of non-irradiated and irradiated OPMF/PBS biocomposite after 24 hours of immersion in distilled water.](image)
Thickness Swelling

The thickness swelling test is conducted to study the dimensional stability of OPMF/PBS biocomposite in the presence of water. Figure 7 shows the thickness swelling of non-irradiated and irradiated OPMF/PBS biocomposites after immersion in water for 24 hours. It indicates that the thickness swelling of the OPMF/PBS biocomposite decreases with increasing EBI treatment dosages. The initial thickness swelling value (10.81%) of the OPMF/PBS biocomposite is reduced to 6.84% upon EBI treatment at 25 kGy, showing a 37% reduction. This is attributed to the decrease in water absorption of the OPMF/PBS biocomposite after EBI treatment, as thickness swelling is proportional to the amount of water absorbed.

CONCLUSIONS

Electron beam irradiation was successfully employed to modify the tensile properties, water absorption and thickness swelling of OPMF/PBS biocomposite. The tensile strength, tensile modulus and elongation at break of the OPMF/PBS biocomposite were improved considerably after electron beam irradiation treatment at an applied dosage of 25 kGy. The electron beam irradiation treatment successfully reduced the water absorption and thickness swelling of the OPMF/PBS biocomposites by 43 and 37%, respectively. The interfacial adhesion between the OPMF and PBS was improved after treatment with electron beam irradiation.

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REFERENCES


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