

From Waste to Reuse: Manufacture of Ecological Composites Based on Biopolyethylene/Wood Powder with PE-g-MA and Macaíba oil

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Abstract

This work aimed to investigate the biopolyethylene (BioPE)/wood powder (WP) composites compatibilized with polyethylene-grafted maleic anhydride (PE-g-MA), using macaíba oil (OM) as a processing aid. The composites were prepared, initially, in an internal mixer and, later, the crushed flakes were molded by injection. Mechanical properties (impact, tensile, flexural and Shore D hardness), heat deflection temperature (HDT), Vicat softening temperature, differential scanning calorimetry (DSC), thermogravimetry (TG), water absorption, torque rheometry and scanning electron microscopy (SEM) were evaluated. The addition of 30% wood powder to the BioPE matrix increased the elastic modulus (tensile and flexural), Shore D hardness and heat deflection temperature (HDT), compared to neat BioPE. These properties were improved when 10% of the PE-g-MA compatibilizer was added, compared to neat BioPE and the non-compatibilized composite. There was a significant reduction in the torque of the composites with the addition of macaíba oil, indicating that it improved the processability. In addition, the incorporation of macaíba oil into the composites helped to reduce water absorption, as well as to increase impact strength. SEM micrographs illustrated a greater degree of interfacial adhesion when PE-g-MA and macaíba oil were added.

1. Introduction

The plastic industry is one of the sectors of the economy that has developed the most in recent years, generating a large number of products derived from conventional thermoplastics [1]. One of the major problems in the use of conventional plastic materials is the environmental impact caused by their disposal [2]. In recent years, there has been an increasing mobilization for sustainable development, with materials that are less aggressive to natural ecosystems [3–5]. The development of polymeric composites reinforced with natural fibers has received great prominence as an alternative technology for the processing of new materials that provide a lower environmental impact, associated with low density and interesting mechanical properties [6–8].

Natural fibers are commonly used in the production of polymeric composites, as they have advantages such as good availability, biodegradability, low cost, good thermal and mechanical properties, renewable, non-toxic and can be easily modified by chemical agents [9, 10]. Currently, there is great academic and industrial interest in the preparation of composites reinforced with jute fibers, sisal, cotton, macaíba, linen, palm fiber, coconut fiber, curauá and wood powder, using biodegradable polymers and green polymers [11–13]. These fibers are widely used to reinforce polyethylene (PE) and polypropylene (PP), aiming at the manufacture of materials for applications in the automotive and aerospace industries [14, 15].

Biopolyethylene (BioPE) produced from ethanol derived from sugar cane, although not biodegradable, maintains the balance of carbon dioxide (CO₂) in nature [16]. In addition to being an ecological material, BioPE has the same mechanical and processing properties as the resin made from fossil sources [17]. For this reason, the development of composites with BioPE polymeric matrix is of great importance, due to the potential for technological application and the reduction of environmental impacts [18]. There are

works in the literature on polyethylene composites reinforced with jute [19, 20], curauá [21], sisal [22, 23], coconut fiber [24, 25], cotton [26], rice husk [27] and wood powder [28, 29], suggesting good mechanical properties.

Castro et al. [30] developed biopolyethylene composites with curauá fibers, using castor and canola oils, as processing aids. When oil was added to the biopolyethylene/curauá fiber composite, it increased the mobility of the polymer chains, decreasing the viscosity of the system and improving processability. In general, the results indicated that the incorporation of oils, particularly castor oil, favored better mechanical properties compared to the composite without oil, indicating the potential of the oil as a compatibilizer.

The use of natural fibers as a reinforcement in composites with biopolyethylene has a great potential to replace synthetic fibers, which will have an impact both in reducing the dependence on materials from non-renewable sources, as well as with regard to environmental and economic aspects [31, 32]. At the same time, there is the possibility of reusing natural fillers, such as wood powder, which is widely discarded by the wood industry. Wood powder residues are sources of environmental pollution and waste of raw material with good properties, with ample potential for reuse in mixtures with polymers [33, 34]. In view of this, both the economic aspects and the environmental pollution are justifications for the research directed with the reuse of wood powder, aiming at the development of composites with good mechanical and thermomechanical properties. The problem associated with the development of composites with high natural filler content is the difficulty during processing, since it increases viscosity. The high viscosity makes it difficult to process the composites, because it is necessary to submit the material between narrow channels in short times, such as extruder and injector. Therefore, aiming at higher productivity with minimum disengage of the equipment, as well as to reduce energy consumption, it is necessary to reduce the composite viscosity. Consequently, it is relevant to investigate processing aid from natural sources, considering that it can improve the composite processing.

The objective of this work was to develop composites of biopolyethylene (BioPE) with wood powder residues, using polyethylene-grafted maleic anhydride (PE-g-MA) as a compatibilizing agent and macaíba oil (OM) as a processing aid.

2. Materials And Methods

2.1 Materials

- Green high density polyethylene, biopolyethylene (BioPE), sold under the code SHC7260® and melt flow index (MFI) of 7.2 g/10 min (190°C/2.16 kg), produced from sugar cane and supplied in the form of pellets by Braskem.
- Angelim wood powder (420 µm), with scientific name *Dinizia excelsa* Ducke, from the wood industry, based in the city of Campina Grande/PB, Brazil.

- Polyethylene-grafted maleic anhydride (PE-g-MA) was used as a compatibilizing agent, marketed under code 3029®, with 1.5 to 1.7% maleic anhydride (MA), supplied in the form of pellets by Addivant. Density of 0.95 g/cm³ and MFI = 4.0 g/10 min (190°C/2.16 kg).

2.2 Methods

2.2.1 Obtaining Macaíba Oil

The procedure for extracting macaíba oil was adopted in the literature [35]. For the assembly of the extractor, a heating blanket with a balloon support (1L) was used, with the Soxhlet (700mL) and the condenser, respectively. Coupled to the system, a coil was installed to cool the water, helping the condensation process. Before starting the extraction process, the filter paper cartridge was weighed on an analytical balance, the mass being determined. After assembling the cartridge with 70g of sample, a new weighing was performed to obtain the mass of the cartridge plus the sample. 1L of hexane was added to the flask and after coupling to the system, the time-controlled extraction process began. During the extraction, an attempt was made to maintain the flow of hexane in the extractor and with the aid of a thermometer, the internal temperature of the soxhlet was monitored, which varied uniformly between 65°C and 70°C.

2.2.2. Composites Processing and Injection Molding

Table 1 illustrates the mass proportions (%) of the compositions that were used in the development of the composites.

Table 1. Compositions of composites with mass proportions (%).

Samples	BioPE (% by weight)	WP (% by weight)	PE-g-MA (% by weight)	Macaíba Oil (% phr)
BioPE	100	-	-	-
BioPE/WP	70	30	-	-
BioPE/WP/oil	70	30	-	10
BioPE/WP/PE-g-MA	60	30	10	-
BioPE/WP/PE-g-MA/oil	60	30	10	10

The compatibilizer PE-g-MA and the wood powder were dried in a vacuum oven for 24 hours, at a temperature of 60°C. The composites were dry mixed and subsequently processed in an internal Haake Rheomix 3000 mixer from Thermo Scientific (310 cm³ capacity), with roller type rotors, at 180 ° C, rotor rotation speed of 60 rpm, under air atmosphere for 10 min. Biopolethylene (BioPE) was processed under the same conditions as composites. Subsequently, the processed material was ground in a knife mill.

The BioPE and the composites were injection molded, in an Arburg injection molding machine, Model Allrounder 207C Golden Edition, to obtain impact, tensile, flexion and HDT specimens, according to the ASTM D256, ASTM D638, ASTM D790 and ASTM standards D648, respectively. The molding conditions of the specimens are shown in Table 2.

Table 2. Injection molding conditions of specimens.

Parameters	Conditions
Injection pressure (bar)	1000
Temperature profile (°C)	170,170,170,170,170
Mold Temperature (°C)	20
Cooling time inside the mold (s)	25
Discharge pressure (bar)	500

2.2.3 Materials Characterization

Rheological curves were obtained by a Haake PolyLab QC da Thermo Scientific mixer with roller-type rotors at 170°C and 60 rpm rotor speed inside an air atmosphere for 10 min. The total mass (Mt) processed in the mixer was measured according to Equation (1):

$$Mt = 0.7 \times D \times V_n \quad (1)$$

where, D = material density to be processed; 0.7 = factor corresponding to 70% of the occupied volume inside the camera; Vn = free volume of mixing chamber (69 cm³).

Izod impact strength test was performed in notched specimens, according to ASTM D256, in a Ceast model Resil 5.5J, operating with a hammer of 2.75J, at room temperature (~ 23°C). Results were analyzed with an average of seven specimens.

Tensile test was performed with injected specimens, according to ASTM D638, using an universal test machine EMIC DL 2000, with a rate of 50 mm/min and load cell of 20 kN, at room temperature (~ 23°C). Results were analyzed with an average of seven specimens.

The flexural test was performed on a universal testing machine of the EMIC DL 2000 brand, according to the ASTM D790 standard, operating in flexion mode at three points, with a speed of 1.6 mm/min, separation between the supports of 40 mm and 20 kN load cell. The results were analyzed with an average of seven specimens.

The penetration resistance was determined according to the ASTM D2240 standard, on a Shore "D" durometer, with a 50 N load controlled by calibrated springs by means of standardized indenters. The results were analyzed with an average of seven penetrations.

Heat deflection temperature (HDT) experiment was measured according to ASTM D648 in a Ceast model HDT 6 VICAT, with a voltage of 455kPa and a heating rate of 120°C/h (method A). The temperature was determined after the sample deflecting 0.25 mm. Results were analyzed with an average of three specimens.

Differential Scanning Calorimetry (DSC) analysis was performed in a DSC-Q20 from TA Instruments. The scans were computed from 30 to 200°C, under a heating rate of 10°C/min, gas flow rate of 50 mL/min, in nitrogen atmosphere, samples with approximately 6 mg were tested.

The thermogravimetry (TG) analyzes were obtained in a simultaneous TG/DSC equipment from TA Instruments SDT Q600, employing about 5 mg of sample, with heating rate of 10 °C/min and gas flow rate of 100 mL/min, starting at a temperature of 30 to 510 ° C, under a nitrogen atmosphere.

The water absorption test was based on the procedure suggested in ASTM D570. The specimens were conditioned in the oven under vacuum, for 24 h at 60 ° C. After this period, they were immediately weighed on a precision scale, and the mass was obtained before immersion (M₀). Then, the specimens were submerged in water at room temperature (~ 23 ° C), so that the samples were completely immersed, obtaining the mass after immersion (M_f). At predetermined time intervals, the specimens were removed from the water (dried with a cloth), weighed on a precision scale and, again, placed in the bath. The percentage of water absorption was calculated from Equation 2.

$$\text{Water absorption (\%)} = \frac{M_f - M_0}{M_0} \times 100\% \quad (2)$$

Scanning Electronic Microscope (SEM) analysis was obtained with a VEGAN 3 TESCAN device with a 30 kV voltage and under high vacuum conditions. The fracture surfaces of impact samples were analyzed. The surfaces were coated with a gold layer.

2.2.4 Statistical Hypothesis Testing

The statistical hypothesis testing is a procedure that allows a decision to be made between two or more hypotheses (null hypothesis H₀ or alternative hypothesis H), using the observed data from a given experiment. Student's t test was applied to compare impact strength, tensile strength, flexural modulus, Shore D hardness and heat deflection temperature (HDT). The criterion adopted for comparative effect was the impact of adding macaiba oil to the properties. Therefore, similar composites BioPE/WP; BioPE/WP/oil and BioPE/WP/PE-g-MA; BioPE/WP/PE-g-MA/oil were evaluated. Table 3 shows the conditions adopted for a 95% confidence level.

Table 3. Conditions tested for Student's t test.

Property	Hypothesis	t_{table}^*
Impact strength (α_0)	H_0 : There is no significant difference.	2.446
	H: There is a significant difference.	
Tensile strength (α_0)	H_0 : There is no significant difference.	- 2.446
	H: There is a significant difference.	
Elastic modulus (α_0)	H_0 : There is no significant difference.	- 2.446
	H: There is a significant difference.	
Flexural modulus (α_0)	H_0 : There is no significant difference.	- 2.446
	H: There is a significant difference.	
Shore D hardness (α_0)	H_0 : There is no significant difference.	- 2.446
	H: There is a significant difference.	
HDT (α_0)	H_0 : There is no significant difference.	-3.182
	H: There is a significant difference.	

* From the literature [82].

3. Results And Discussion

3.1 Torque Rheometry

Torque rheometry is a technique used to evaluate the reactivity and processability of composites [36]. Figure 1 shows the torque versus time curves for neat BioPE and composites with and without macaíba oil. A significant increase in torque is observed in 1 min of processing, due to the solid material in the mixing chamber and, therefore, greater mechanical work to rotate the rotor. Composites have a higher intense peak torque compared to neat BioPE. Such behavior is due to the process of distribution and dispersion of wood flour particles, since they are harder. Apparently, the presence of macaiba oil did not significantly influence processing start of processing, possibly due to the high content of wood flour. After melting and mixing, the torque for all materials decreases and stabilizes, suggesting a homogeneous mixture. It appears that after 3 min of the process, the torque of the BioPE is constant, with a value around 1.8 N.m. This behavior indicates viscosity stability for the process conditions used, i.e., a speed of 60 rpm and a temperature of 180 °C. There is no evidence of degradation for BioPE, since there has been no continuous reduction in torque over time. The oil was more effective in reducing the torque after homogenizing the composites, as verified after 3 min.

The composites BioPE/WP and BioPE/WP/PE-g-MA presented in 10 min torques of approximately 2.5 N.m, indicating a 38.8% increase in torque, in comparison to BioPE. The addition of 30% wood powder hindered the melt flowing, since it acts as a filler and reduces molecular mobility, causing an increase in the viscosity of the composites, as also verified in the literature [37]. In this case, there is a higher energy consumption for processing the composites BioPE/WP and BioPE/WP/PE-g-MA, in relation to BioPE.

In the interval of 2 to 4 min there was a slight increase in the torque of the BioPE/WP/PE-g-MA composite compared to BioPE/WP, indicating a possible interaction between the compatibilizer PE-g-MA with the wood powder. It is believed that the maleic anhydride groups of PE-g-MA and the hydroxyl groups of wood powder have interacted [38], as shown in Figure 2. After 5.5 min the torque curves of the BioPE/WP and BioPE/WP/PE-g-MA composites are superimposed.

Despite containing 30% wood powder, the BioPE/WP/oil composite showed a reduction in torque (1.55 N.m), compared to BioPE. The same trend of torque reduction occurred for the BioPE/WP/PE-g-MA/Oil (1.98 N.m) composite, compared to the BioPE/WP and BioPE/WP/PE-g-MA systems. Macaiba oil is acting as a plasticizer in composites, increasing molecular mobility, reducing viscosity and equilibrium torque (10 min). In this case, it is suggested macaiba oil is acting as a processing aid for composites, improving the melt flowing. In this way, macaiba oil is able to improve the flow properties and decrease the adherence of the melt with the machine components. Thus, energy consumption is saved and productivity can be increased.

The BioPE/WP/PE-g-MA/Oil composite showed a torque 27.7% higher than the BioPE/WP/oil system, indicating that despite having the same proportion of wood powder (30%), the plasticization mechanism was different. The different behavior of these composites is associated with the compatibilizer PE-g-MA, suggesting there was interaction between the functional groups maleic anhydride, the hydroxyls of wood powder and macaiba oil. As a consequence, there was a balance between the plasticizer effect and the restriction of filler mobility in the BioPE/WP/PE-g-MA/Oil composite, generating an intermediate torque curve between BioPE and the BioPE/WP and BioPE/WP/PE-g-MA systems. The literature [40,41] reports that natural oils have technological potential to interact with wood powder, including to act as a compatibilizer.

3.2 Scanning Electron Microscopy (SEM)

Through scanning electron microscopy it was possible to evaluate the efficiency of the wood filler dispersion process in the BioPE matrix and the surface wetting capacity of the composites, as a way of evaluating the efficiency of the compatibilizing agent and the macaiba oil. Figure 3 (a, b, c, d) illustrates the micrographs obtained by SEM from composites, with and without macaiba oil. In Figure 1 (a), the BioPE/WP composite showed a large amount of agglomerated wood fibers, indicating that there was poor dispersion during processing. In this case, due to the presence of agglomerates, a localized stress concentration can occur during the mechanical test, with the nucleation of cracks and generating a premature failure in the composite [42]. In addition, there is a weak adhesion between the wood fibers and BioPE. The poor adhesion occurs due to the high interfacial tension existing between the components,

due to the hydrophilic nature of BioPE's wood and hydrophobic powder, which makes the interface with low resistance, corroborating the lower performance under the impact of this system [43].

In Figure 3 (b), it can be seen the BioPE/WP/oil composite did not present agglomerations, indicating that macaíba oil helped to disperse the wood fibers during processing. In addition, it presents well-adhered wood fiber in the BioPE matrix (see square) and, at the same time, poorly adhered particles (marked with a circle), with an interface without adhesion. However, clearly macaíba oil promoted a more stable morphology in the composite, improving the interfacial adhesion and the degree of dispersion of the fillers, confirming the better performance under the impact of the BioPE/WP/oil system, in relation to the BioPE/WP composite (Figure 4).

Figure 3 (c, d) shows that the addition of PE-g-MA and macaíba oil increased the interaction of the wood fiber and the BioPE matrix, improving the degree of interfacial adhesion. In this case, the applied stress is better distributed from BioPE to wood fiber, contributing to a higher performance under impact [44], as further on presented. At the same time, there is no evidence of delamination or agglomeration of fillers, suggesting that better distribution and dispersion occurred during processing. Apparently, the compatibilizer PE-g-MA favored higher efficiency of the level of wetting of the wood fiber by BioPE, suggesting interactions between the functional groups, as discussed in torque rheometry.

The composites BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil presented strong interfaces, Figure 3 (c, d), reinforcing the idea that PE-g-MA acts as a proper compatibilizing agent in these systems. The interfacial adhesion between the fiber and the matrix is essential to obtain high mechanical properties in composites [45]. As the stress applied on the composite material is transferred from the matrix to the fibers through the interface, a good interfacial adhesion is necessary to optimize the mechanical properties [46].

3.4 Impact Strength

Figure 4 shows the impact strength of BioPE and composites with and without macaíba oil, respectively. BioPE presented the highest impact strength (91.2 J/m), a typical ductile polymer value. The BioPE/WP composite reduced the impact strength by around 60%, compared to neat BioPE. This occurred due to the addition of 30% wood powder (WP), since it is a stiff filler and, consequently, tended to weaken the ductile matrix [47]. The fillers act as stress concentrators, restricting the mobility of the matrix and reducing the plastic deformation [48]. The BioPE/WP/PE-g-MA composite increased the impact strength by 32.5%, compared to BioPE/WP. The increase in impact strength is taken as an indication that the addition of PE-g-MA promoted the interaction between the present phases, as seen in the torque rheometry and SEM. As a consequence, it strengthened the system interface, which is essential for the tension transfer between the phases [49,50].

The BioPE/WP/oil composite showed a gain in impact strength of approximately 31% compared to the BioPE/WP, indicating macaíba oil increased flexibility, generating higher level of energy dissipation. Apparently, macaíba oil also minimizes the effect of the polar difference between the hydrophobic

matrix of BioPE and the wood powder with hydrophilic nature, improving the interaction at the interface, as verified in the SEM. In fact, the composites BioPE/WP/oil and BioPE/WP/PE-g-MA are resistant to impact within the experimental error range and, in this case, are comparable in this property. It is reasonable to suggest that macaíba oil is acting as a plasticizer, increasing the free volume and molecular mobility of the BioPE/WP/oil composite, contributing to increase the compatibility of impact properties.

The greatest impact strength was achieved with BioPE/WP/PE-g-MA/oil composite, with a value in the order of 56.8 J/m. The performance of this composite is the result of the combined behavior of the reinforcement element, compatibilizer and the macaíba oil. Figure 5 presents a schematic representation of the BioPE/WP/PE-g-MA/oil interaction. The compatibilizer PE-g-MA reacts with the hydroxyls of the wood powder and, at the same time, there is a physical entanglement with the BioPE polymer matrix chain. The macaíba oil has oleic acid in its constitution [51], which promotes interactions of hydrogen bonds with the hydroxyls present on the surface of the wood powder and plasticizes the composite, generating compatibility and increasing impact strength [52, 53].

The statistical evaluation was carried out to analyze the macaíba oil effect on impact strength. Therefore, similar composites were evaluated, that is, BioPE/WP; BioPE/WP/oil and BioPE/WP/PE-g-MA; BioPE/WP/PE-g-MA/oil. A significance level of 95% was adopted and the $t_{table} = 2,446$. BioPE/WP and BioPE/WP/oil composites presented a $t_{calculated} = 2.574$, indicating that macaíba oil provided significant increase in impact strength. On the other hand, the BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil mixtures indicated a $t_{calculated} = 2.132$ and, in this case, there is no significant difference in the impact strength of these composites ($t_{calculated} < t_{table}$). Statistical results indicate that macaíba oil has the potential to make the BioPE/WP system compatible, generating a gain in impact strength. However, the BioPE/WP/PE-g-MA/oil hybrid composite, despite having a gain in impact strength with the addition of oil, was not as effective compared to the BioPE/WP/oil system. Such trend indicates that it is more feasible to use PE-g-MA or macaíba oil individually aiming at compatible BioPE/WP composite, in order to optimize the impact strength.

3.5 Tensile Strength

Figure 6 shows the results of the elastic modulus of BioPE and composites, with and without macaíba oil, respectively.

BioPE presented the lower elastic modulus with a value around 400 MPa, indicating that it is the most flexible material, as verified in the impact strength. The elastic modulus increased with the addition of 30% wood powder in the BioPE/WP composite, reaching a gain of 53%, compared to neat BioPE. This increase is due to the fact that wood powder is more stiff than BioPE matrix and prevents its free mobility [54]. A more expressive increase in the elastic modulus was achieved for the BioPE/WP/PE-g-MA composite, with gains of 65.6% and 8.2%, compared to BioPE and BioPE/WP, respectively. Thus, there is evidence that the compatibilizer PE-g-MA showed interfacial action, promoting a better adhesion between

the BioPE matrix and the wood powder, contributing to increase the elastic modulus [55]. The higher performance in the stiffness of the BioPE/WP/PE-g-MA composite may be related to the greater wettability of BioPE with wood powder, as seen in the SEM (Figure 3c).

The composites BioPE/WP/oil and BioPE/WP/PE-g-MA/oil did not show significant differences between them, as they are within the margin of experimental error. In comparison to BioPE, the composites BioPE/WP/oil and BioPE/WP/PE-g-MA/oil increased 32.5% and 39%, respectively. However, composites with macaíba oil became more flexible and reduced stiffness, compared to BioPE/WP and BioPE/WP/PE-g-MA. This behavior can be explained by the plasticizing action of macaíba oil, which probably acts in the amorphous phase of BioPE, inserting itself between its macromolecules and favoring sliding and molecular mobility [56], as seen with the reduction of viscosity in torque rheometry. The BioPE/WP; BioPE/WP/oil composites and BioPE/WP/PE-g-MA; BioPE/WP/PE-g-MA/oil composites were evaluated by statistical test, with a 95% significance level and the $t_{table} = -2.446$. The BioPE/WP and BioPE/WP/oil composites presented a $t_{calculated} = -1.964$, while the BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil systems indicated a value of $t_{calculated} = -2.183$. These results show that although macaíba oil reduced the elastic modulus, there was no drastic reduction in this property. In this case, even the macaíba oil increasing the flexibility of the composites kept the tensile elastic modulus high, even surpassing the neat BioPE.

The tensile strength results of BioPE and composites with and without macaíba oil are shown in Figure

The tensile strength of the BioPE/WP composite reduced approximately 10% compared to BioPE, indicating that the wood powder is acting as filler. At the same time, the BioPE/WP composite showed agglomerates of wood particles, as seen in Figure 3 (a), generating a concentration and intensification of the stress [57], contributing to the decrease in tensile strength. When the BioPE/WP composite was made compatible with PE-g-MA, the tensile strength reached the highest performance with 29.6 MPa, corresponding to gains of 39.6% and 54.9%, compared to BioPE and the composite BioPE/WP. This behavior can be associated with good adhesion and homogeneous dispersion among the constituents of the composite, as verified in the SEM. The fracture surface morphology of the BioPE/WP/PE-g-MA composite showed a good wettability of the wood powder with the BioPE matrix, providing stress transfer from the matrix to the filler, resulting in an increase in tensile strength.

The BioPE/WP/oil composite showed a marked reduction in tensile strength, with decreases of 29.7% and 21.9%, in relation to BioPE and the BioPE/WP composite, respectively. This behavior indicates that the macaíba oil plasticized, reducing the maximum resistance supported by the BioPE/WP/oil composite. The BioPE/WP/PE-g-MA/oil system did not significantly modify the tensile strength (22.6 MPa), only maintaining the value compared to BioPE. However, the BioPE/WP/PE-g-MA/oil composite showed a gain of 18.3%, compared to BioPE/WP. Clearly, the addition of macaíba oil impaired the effect of wood powder reinforcement on the BioPE/WP/PE-g-MA composite, leading to a loss of mechanical strength due to the plasticizer effect. The statistical evaluation was carried out to analyze the effect of macaíba oil on the tensile strength, with a significance level of 95% and the $t_{table} = -2.446$. The comparative test was applied

to similar composites, that is, BioPE/WP; BioPE/WP/oil and BioPE/WP/PE-g-MA; BioPE/WP/PE-g-MA/oil. The BioPE/WP and BioPE/WP/oil composites showed $t_{\text{calculated}} = -2.578$, while the BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil systems indicated a value of $t_{\text{calculated}} = -2.613$. The tests indicated that $t_{\text{calculated}} < t_{\text{table}}$, therefore, there is a statistical difference in tensile strength. Such behavior suggests there was significant reduction in tensile strength, that is, macaíba oil contributed significantly to deteriorate this property, due to the plasticizer effect. As consequence, composites deformed at lower stresses.

3.6 Flexural Modulus

Figure 8 shows the results of flexural modulus of BioPE and composites, with and without macaíba oil, respectively. The BioPE/WP composite showed a flexural modulus value higher than the BioPE, with a gain of 89%. A more expressive increase in the flexural modulus was verified with the BioPE/WP/PE-g-MA composite, reaching gains in the order of 140% and 26.9%, in relation to BioPE and the BioPE/WP composite. The higher performance of the BioPE/WP/PE-g-MA system can be explained by the interactions triggered by PE-g-MA at the BioPE/wood powder interface, when hydroxyl groups (OH) react with maleic anhydride (MA) providing greater anchoring to the composite [58,59].

The addition of macaíba oil caused a significant decrease flexural modulus of the composites BioPE/WP/oil and BioPE/WP/PE-g-MA/oil, making them more flexible. The addition of macaíba oil to the composites, regardless of whether they are compatible or not, compromised the stiffness, corroborating the same trend of the elastic modulus. Such behavior is due to the diffusion of macaíba oil to the amorphous structure of BioPE, with a plasticizing effect and increasing molecular mobility [60]. The statistical test was performed to verify if there was significant reduction in the flexural modulus, with significance level of 95% and the $t_{\text{table}} = -2.446$. BioPE/WP and BioPE/WP/oil composites showed a $t_{\text{calculated}} = -3.764$, while the BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil systems indicated a value of $t_{\text{calculated}} = -4.183$. The statistical results demonstrated macaíba oil addition severely impairs the flexural modulus ($t_{\text{table}} > t_{\text{calculated}}$). Therefore, if required rigid composite which will be submitted to flexural effort, macaíba oil should be avoided, as it impairs this property. The result is opposite to that presented by the elastic modulus, however it is important to mention that the mechanical effort is different for the two tensile tests.

3.7 Shore D Hardness

Figure 9 shows the results of Shore D hardness of BioPE and composites, with and without macaíba oil, respectively. It is observed that BioPE has the lowest Shore D hardness, due to its more flexible behavior, as seen in the impact strength and elastic modulus. The addition of 30% wood powder increased the Shore D hardness of all composites, in relation to BioPE. The composites BioPE/WP and BioPE/WP/PE-g-MA increased 10.7% and 11.2% when compared to BioPE. This is due the fillers are more stiff than the matrix, reducing mobility and increasing stiffness [61], as seen in the elastic modulus and flexural

modulus. As a consequence, there is an increase in the penetration resistance, contributing to the increase in Shore D hardness of the composites.

The penetration resistance is a quick test to evaluate the influence of macaíba oil in the composites, as seen in Figure 9. The trend is the reduction in Shore D hardness of the composites, BioPE/WP and BioPE/WP/PE-g-MA, with the addition of macaíba oil, due to the increase in molecular spacing [62]. The composites BioPE/WP/oil and BioPE/WP/PE-g-MA/oil showed slight gains of 6% and 7.2%, in relation to BioPE. These composites do not present significant differences between them, considering that they are within the margin of experimental error and, in this case, have penetration resistance comparable.

Shore D hardness was assessed by the statistical test, with a 95% significance level and the $t_{table} = -2.446$. Again, BioPE/WP; BioPE/WP/oil and BioPE/WP/PE-g-MA; BioPE/WP/PE-g-MA/oil composites were evaluated in order to analyze the effect of macaíba oil. The tests indicated a $t_{calculated} = -1.841$ for the BioPE/WP and BioPE/WP/oil composites, while the BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil systems presented $t_{calculated} = -1.893$. The results show that there is no significant difference in Shore D hardness between similar composites ($t_{calculated} > t_{table}$), that is, the addition of macaíba oil did not severely impact this property.

Heat Deflection Temperature (HDT)

Heat deflection temperature (HDT) is an extremely important property in the area of polymers, since it indicates the dimensional stability of the materials, at high temperatures [63,64]. Figure 10 shows the results of the heat deflection temperature of BioPE and composites, with and without macaíba oil, respectively. It is observed that the HDT value for BioPE was 70.3°C, value close of the literature [65].

The heat deflection temperature is strongly influenced by the stiffness of the system, with a tendency to increase the deflection temperature, with the addition of the elastic modulus [66]. The composites BioPE/WP and BioPE/WP/PE-g-MA showed very significant increases in HDT, with gains of 44.1% and 48.1%, compared to BioPE. These results indicate that the addition of 30% wood powder in BioPE improved the thermomechanical resistance, impacting on a greater structural stability at higher temperatures. In comparative terms, the heat deflection temperature of the composites BioPE/WP and BioPE/WP/PE-g-MA was, on average, 34°C higher than that of BioPE, suggesting that the incorporation of a stiff filler contributed to the corresponding increase this property, corroborating the results of elastic modulus and flexural modulus. Apparently, the influence of the compatibilizer PE-g-MA was not significant in this HDT property, since the composites BioPE/WP and BioPE/WP/PE-g-MA are within the experimental error margin.

The composites BioPE/WP and BioPE/WP/PE-g-MA additivated with macaíba oil showed gains of 26.3% and 30.3%, in relation to BioPE. However, the performance of these composites with macaíba oil decreased when compared to similar BioPE/WP and BioPE/WP/PE-g-MA without oil, indicating that the plasticizer effect impairs this property, due to reduced stiffness, as verified in the tensile and flexion tests.

The statistical test with a 95% significance level and the $t_{table} = -3.182$ was applied to analyze the impact of macaíba oil. BioPE/WP and BioPE/WP/oil composites showed a $t_{calculated} = -2.324$, while the BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil composites indicated a $t_{calculated} = -2.187$. As $t_{calculated} > t_{table}$ there was no significant reduction in heat deflection temperature (HDT), with the addition of macaíba oil. The results indicate that although macaíba oil reduces the HDT of the composites, there was no severe loss in this property, contributing to keep good thermomechanical stability.

The increase in thermomechanical resistance caused by the presence of 30% wood powder provides an important technological contribution to the BioPE/wood powder composites, since it significantly increased HDT. Although macaíba oil reduced the HDT value, its presence still keeps high HDT and, at the same time, improves processability, as seen in torque rheometry.

3.8 Differential Scanning Calorimetry (DSC)

The melting, crystallization parameters and the degree of crystallinity of BioPE and composites are summarized in Table 4. The DSC scans obtained during the second heating cycle can be seen in Figure 11 (a, b).

Table 4. Melting and crystallization parameters of BioPE and composites.

Samples	Tc (°C)	ΔH_c (J/g)	Tm (°C)	ΔH_m (J/g)	Xc (%)
BioPE	115.6	187.3	133.9	176.6	60.2
BioPE/WP	119.1	126.0	132.8	115.2	56.2
BioPE/WP/Oil	118.3	94.7	131.6	99.1	48.3
BioPE/WP/PE-g-MA	117.9	117.0	132.8	107.7	61.2
BioPE/WP/Pe-g-MA/Oil	118.4	109.8	131.0	104.4	59.4

* T_m = Melting peak temperature; T_c = Crystallization peak temperature; ΔH_c = Crystallization enthalpy; ΔH_f = Melting enthalpy; X_c = Degree of crystallinity, $X_c = \Delta H_f / w * \Delta H_0$, where: w = mass fraction of BioPE; ΔH_0 = Melting enthalpy BioPE at 100% of crystallinity, 293 J/g [21].

In Table 4, it is possible to see a single melting peak around 133.9 °C, commonly found in a polymeric polyethylene matrix [67]. The melting temperature (T_m) of the composites did not change significantly in relation to BioPE, only subtle changes. Composites additivated with macaíba oil, although not a significant variation, tended to have the lowest T_m . As the reduction in the melting temperature is often associated with the degree of crystals perfection [68], possibly macaíba oil may have helped to diffuse small particles of wood powder to the amorphous region of BioPE, reducing its degree of crystals perfection. There is a marked reduction in the melting enthalpy (ΔH_f) of the composites compared to BioPE, due to the high content of wood powder (30%) and the reduction of the BioPE content in the

formulation. When the composites BioPE/WP and BioPE/WP/PE-g-MA are additivated with macaíba oil, there is a more evident reduction in the melting enthalpy, suggesting that the presence of macaíba oil reduces the energy consumption to melt the crystals, corroborating with torque rheometry.

The crystallization temperature (T_c) of the composites increased in relation to neat BioPE, indicating that the wood powder accelerates the crystallization process. From a practical point of view, injection molded composites can be removed hotter from the molds, as they solidify at higher temperatures. The enthalpy of crystallization of composites requires less energy to promote crystallization, compared to BioPE. The composites BioPE/WP and BioPE/WP/PE-g-MA with macaíba oil show the lowest enthalpy of crystallization, suggesting that the oil contributed to the structural organization during cooling.

Biopolyethylene showed a degree of crystallinity in the order of 60.2%, a value close to reported in literature [69]. The BioPE/WP composite showed 56.2% crystallinity, a reduction compared to BioPE, due to poor wettability and the low interfacial interaction between BioPE and wood powder, as seen in the SEM (Figure 3a). When the BioPE/WP composite contains macaiba oil, a significant reduction occurred in the degree of crystallinity, with a value around 48.3%. As macaíba oil is acting as a plasticizer in the BioPE/WP composite, increasing molecular mobility, the degree of crystallinity would have increased or at least maintained the crystallinity of the BioPE/WP composite without oil. Apparently, the addition of macaiba oil directly to the BioPE/WP system hindered the formation of crystals, reinforcing the hypothesis that the oil helped diffuse small particles of wood powder into the amorphous chains of BioPE, creating difficulties for molecular packaging and reducing crystallinity [70].

The composites BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil showed a degree of crystallinity similar to neat BioPE, indicating that PE-g-MA acts in order to improve the structural organization, increasing the degree of crystallinity. This finding is related to a better dispersion of wood powder in BioPE matrix, as well as a greater wettability between the phases, as verified in the SEM.

3.9 Thermogravimetry (TG)

Figure 12 illustrates the TG's curves of BioPE and composites, with and without macaíba oil, respectively. It is observed that BioPE presented only one stage of decomposition at temperatures above 400°C, which guarantees good thermal stability for the process conditions used. A reduction in thermal stability is seen for all composites, compared to BioPE, due to the presence of wood powder [71]. The composites presented three evident decomposition steps, the first in the range of room temperature up to 100°C, attributed to the loss of water by desorption, considered a thermal event of physical origin, as there is no loss of mass of the material [72]. The second loss after 250°C is attributed to the decomposition of wood powder components, which are cellulose, hemicelluloses and lignin, as well as macaiba oil [73]. The third mass loss event is attributed to the decomposition of the BioPE chain and remaining lignin.

In order to evaluate the relative thermal stability of the composites, the temperatures for 10% and 50% of mass loss ($T_{0.1}$ and $T_{0.5}$) were calculated and are shown in Table 5. BioPE is the most thermally stable material, as verified with $T_{0.1}$ and $T_{0.5}$, i.e., there is a loss of mass at higher temperatures. The BioPE/WP

composite showed the lowest thermal stability at 10% of mass loss, since it started to decompose at a lower temperature, around 312°C. This behavior is related to the weak interactions between BioPE and wood powder, as seen in the mechanical results presented previously, as well as in the morphological analysis (Figure 3a). The addition of the BioPE/WP composite with macaíba oil moved the weight loss ($T_{0.1}$) to a higher temperature, around 325°C. The use of macaíba oil clearly improved the interfacial adhesion between BioPE and wood powder (Figure 3b), generating an increase in the thermal stability of the BioPE/WP/oil composite. The literature [74] has shown that the use of natural oil in composites can improve thermal stability, with an effect similar to a compatibilizing agent.

The composites BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil showed a higher stability temperature for the loss of mass in $T_{0.1}$, which is an indication of interaction between the components of the mixture, as seen in torque rheometry and SEM. As a consequence, from the greater synergism there was a stabilizing effect. The literature [75] indicates that the improvement in thermal properties reflects the stabilization of morphology, in which the components present interfacial interactions, with better compatibility. The 50% mass loss temperature practically does not change between composites, only similar losses.

Table 5. Thermal stability of neat BioPE and composites, with and without macaíba oil.

Samples	$T_{0.1}$ (°C)	$T_{0.5}$ (°C)	Residue at 520°C (%)
BioPE	433.8	467.2	0.0
BioPE/WP	312.9	461.5	1.1
BioPE/WP/Oil	324.9	458.1	1.4
BioPE/WP/PE-g-MA	341.0	461.3	2.9
BioPE/WP/PE-g-MA/Oil	333.4	468.9	3.2

In Table 5, the composites at 510°C showed residual carbonaceous material, resulting from decarboxylation, decomposition and depolymerization of cellulose and hemicellulose fragments, as reported in literature [76, 77].

3.10 Water Absorption

The degree of water absorption is an important characteristic to be evaluated in composites with natural fibers, as it assesses the potential of these materials for use in certain applications, such as in external applications in which there is contact with moisture [78]. Figure 13 shows the water absorption curve for BioPE and composites, with and without macaíba oil, respectively.

BioPE practically did not absorb water during 1728 h, suggesting a stability against humidity, due to the hydrophobic character of this polymer. The composites showed a fast water absorption in the first 200 hours, due to wood powder. Natural fibers have a hydrophilic behavior, due to the presence of hydroxyl

groups in their structure, thus absorbing water [79]. Over time, the curve changes its slope smoothly until it reaches a linear level, which represents the level of moisture saturation.

The macaíba oil and the compatibilizer, impact on the water absorption kinetics, as shown in Figure 13. The BioPE/WP composite showed the highest level of water absorption in 1728 h, due to the ease of diffusion of water molecules among micro-voids polymer chains, as well as defects resulting from the low adhesion between BioPE and wood powder, as seen in the SEM. When the BioPE/WP composite was additivated with macaíba oil, a reduction in water absorption is verified. This finding indicates that macaíba oil acts as a barrier effect, reducing the diffusion of water into the composite.

The composites BioPE/WP/PE-g-MA and BioPE/WP/PE-g-MA/oil significantly reduced water absorption in 1728 h, compared to the BioPE/WP and BioPE/WP/oil systems. This finding indicates that there was an increase in the interaction between the components of the composites, reinforcing the reactions proposed in Figures 2 and 5. The literature [80, 81] attributed the reduction in water absorption to the reaction of the hydrophilic groups –OH present in the fibers with the groups anhydrides present in the compatibilizer. As a consequence, there is a decrease in the availability of hydrophilic sites in wood powder, impacting on the reduction of water absorption.

4. Conclusions

The impact of macaíba oil on bio-polyethylene/wood powder composites compatibilized with PE-g-MA was investigated. The addition of macaíba oil improved the processability of the composites, with a plasticizer effect. However, the presence of macaíba oil tended to reduce the elastic modulus, Shore D hardness and heat deflection temperature, in comparison to composites without macaíba oil. In contrast, the use of macaíba oil improved the interfacial adhesion between BioPE and wood powder, generating gains in impact strength and reduction in water absorption. This last result is important, since the decrease in water absorption can increase dimensional stability. When the composite formulation is BioPE/wood powder/PE-g-MA/macaíba oil, good results are achieved, generating a synergism of properties and processing. The results indicate a good perspective of application of these composites, with a reduced price due to the possibility of reusing up to 30% of a material that would be discarded. In addition, the harmful effect of this residue on the environment can be minimized and added value to a rejected material.

Declarations

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Figures

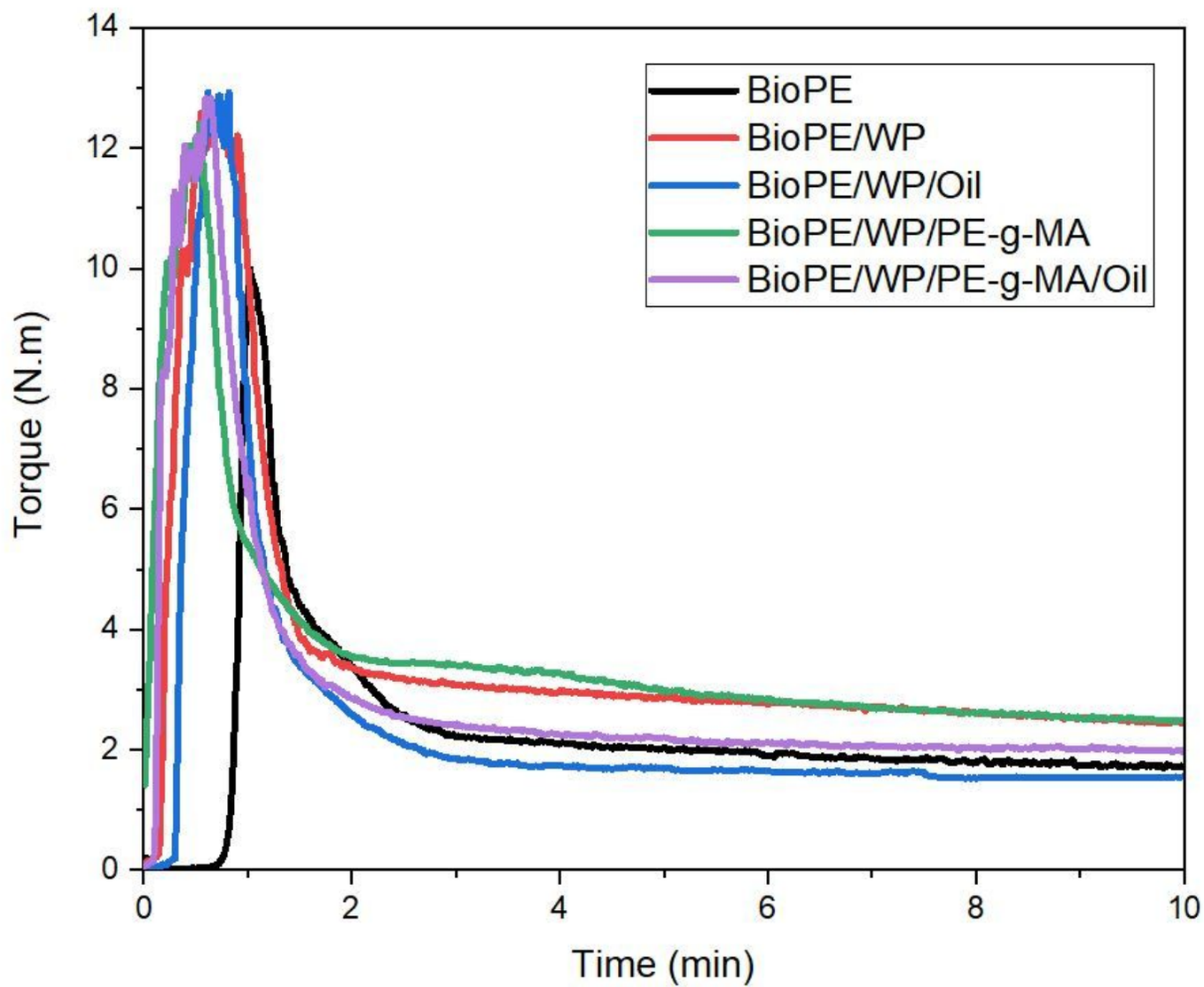


Figure 1

Torque versus time curves for neat BioPE and composites, with and without macaíba oil.

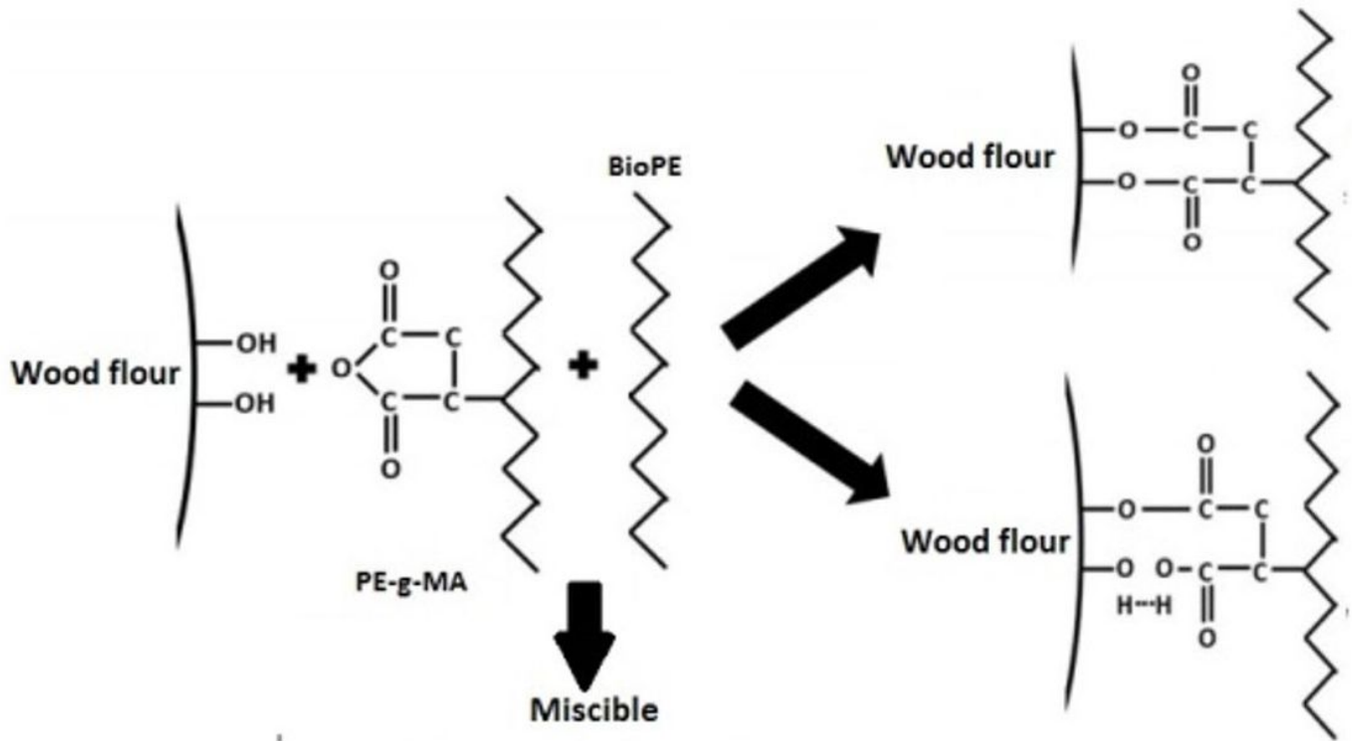


Figure 2

Proposed reaction for the compound BioPE/WP/PE-g-MA, with adaptation of [39].

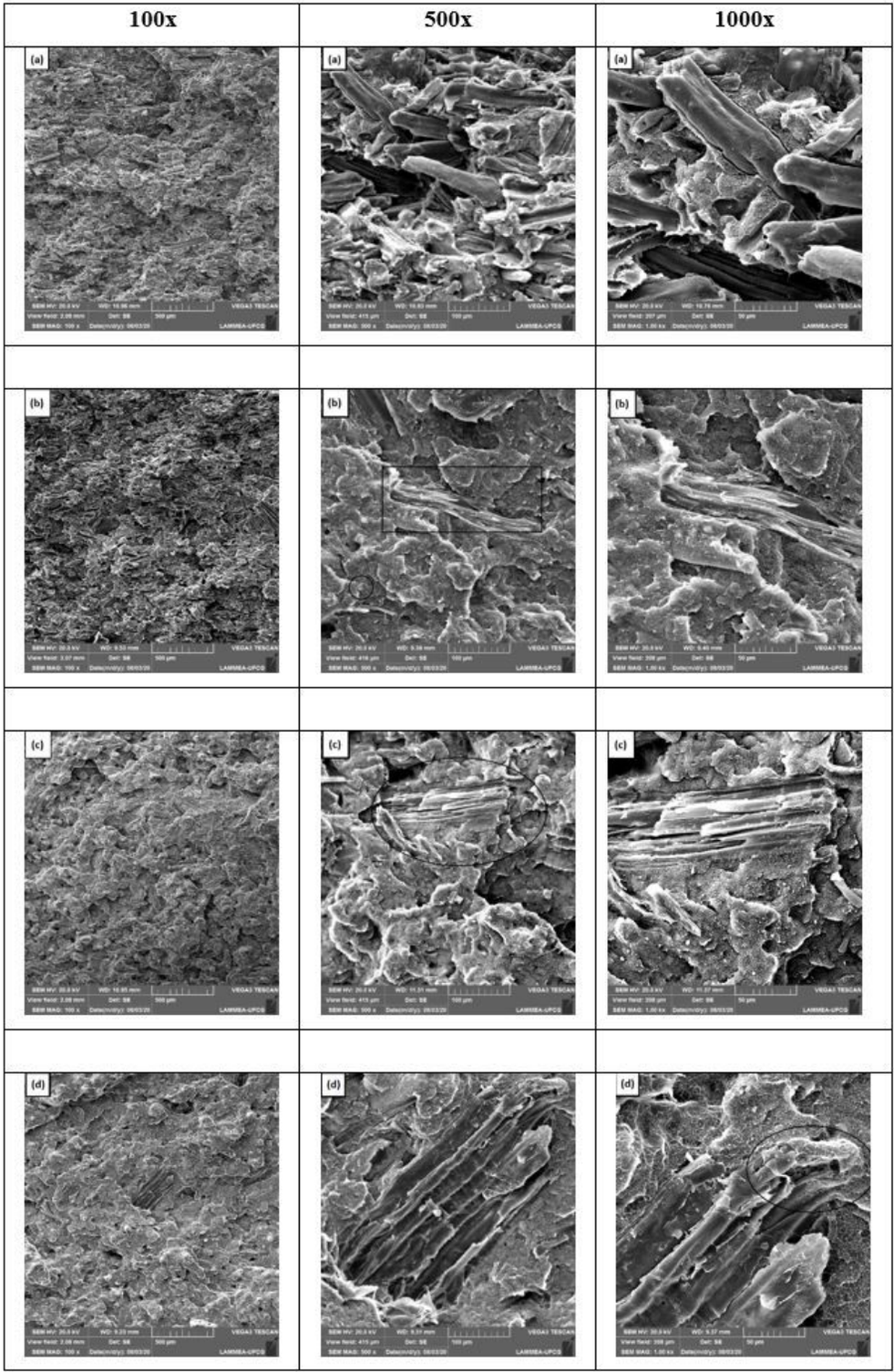


Figure 3

SEM images of the fractured surfaces, with 100x; 500x and 1000x magnification: (a) BioPE/WP; (b) BioPE/WP/oil; (c) BioPE/WP/PE-g-MA; (d) BioPE/WP/PE-g-MA/oil.

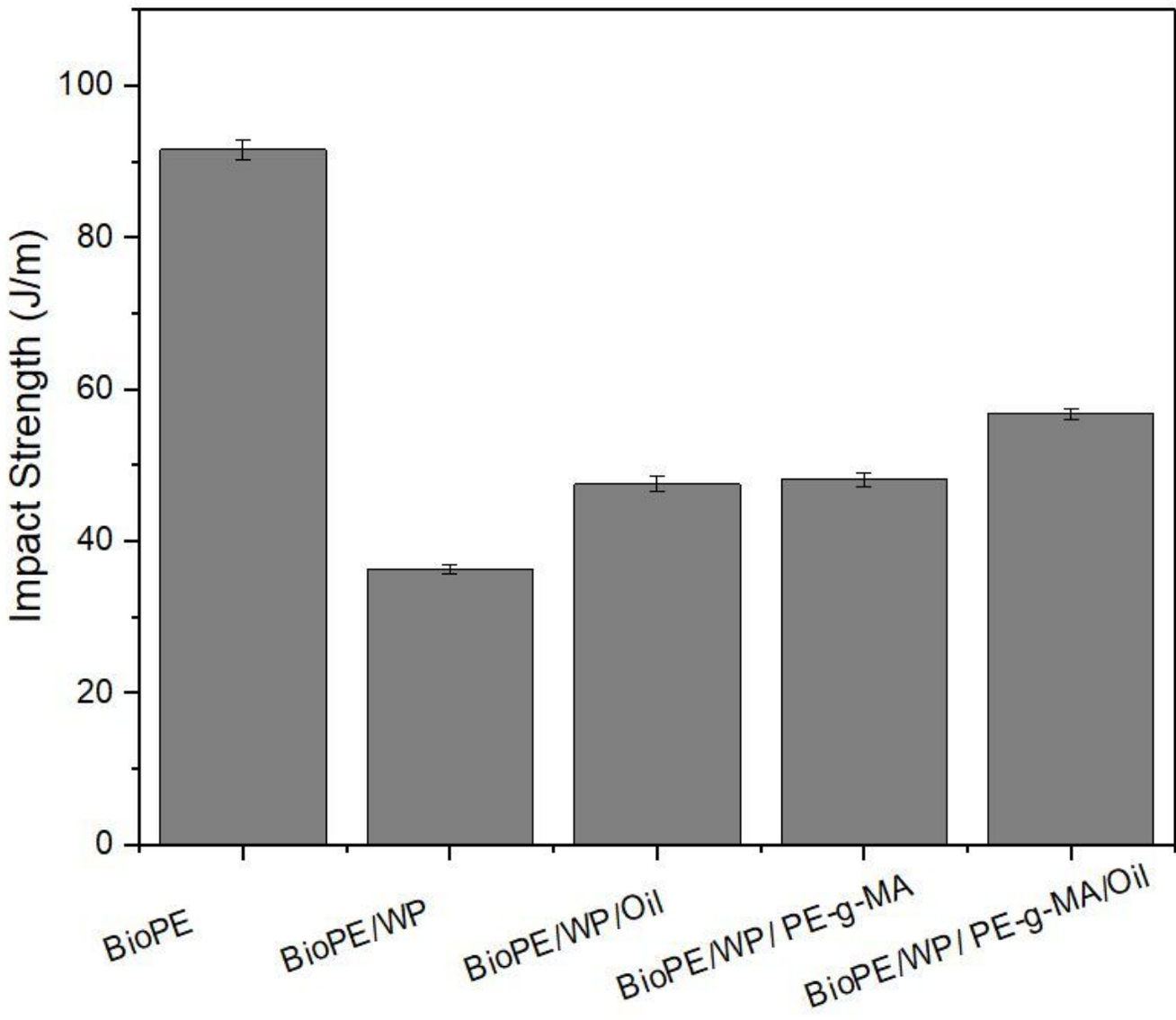


Figure 4

Izod impact strength of BioPE and composites, with and without macaiba oil.

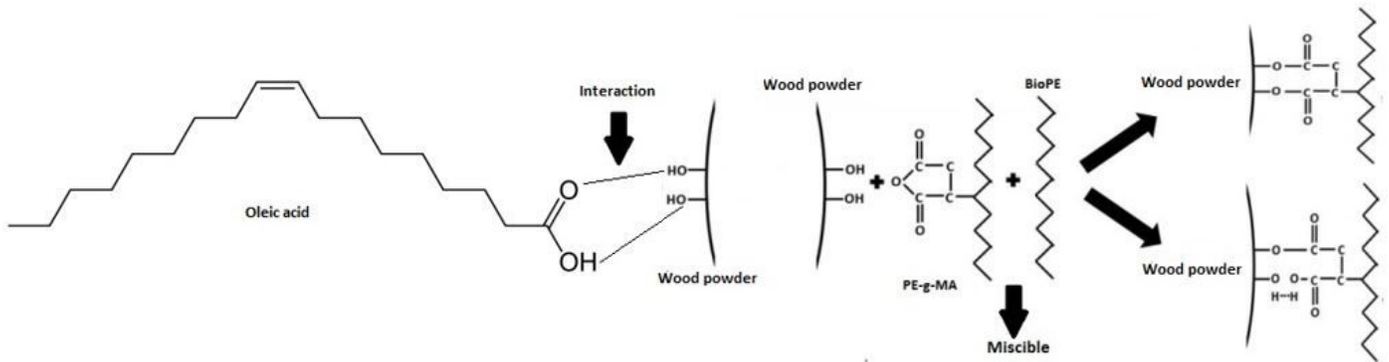


Figure 5

Proposed reaction for the compound BioPE/WP/PE-g-MA/oil, with adaptation of [39].

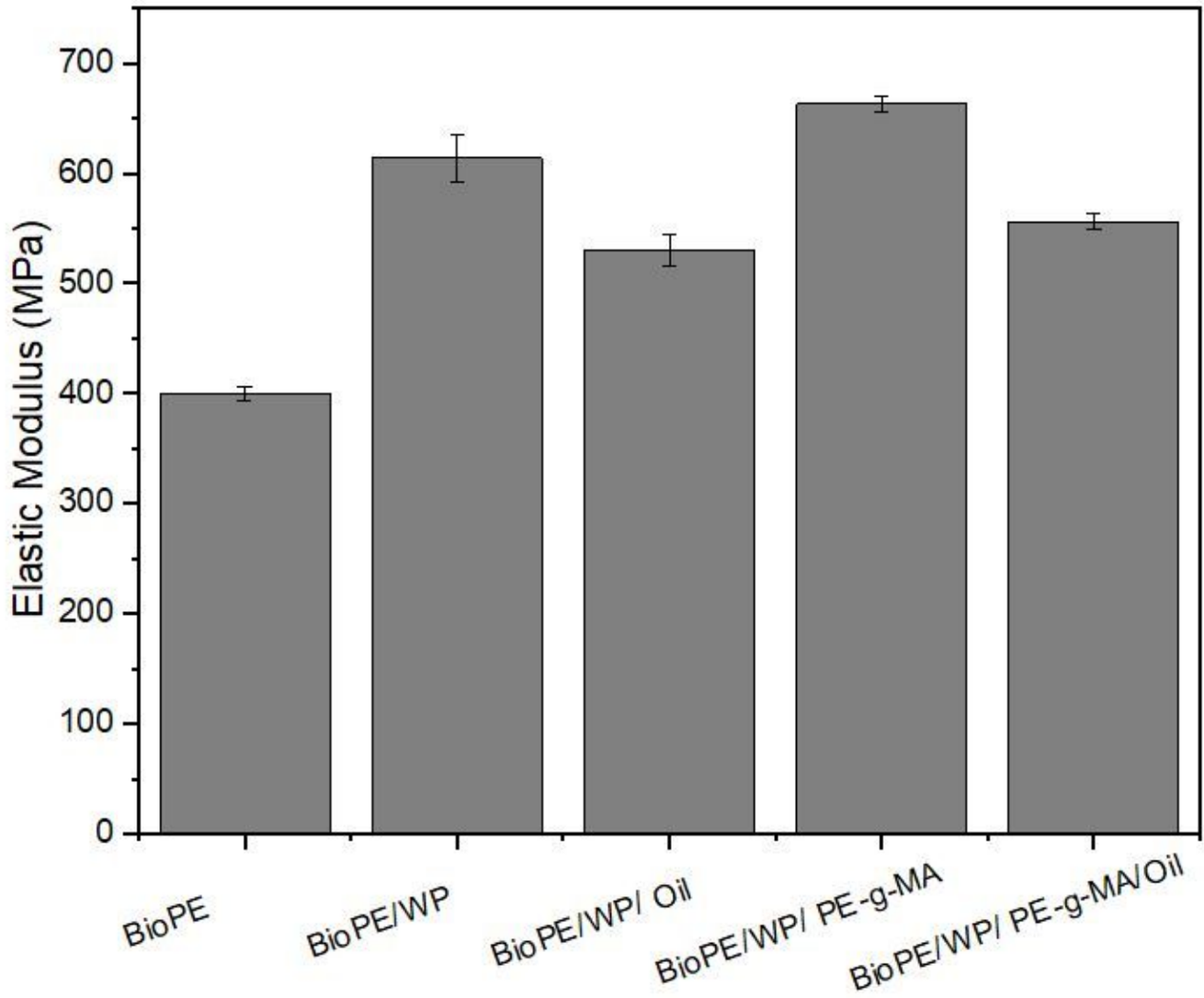


Figure 6

Elastic modulus of BioPE and composites, with and without macaiba oil.

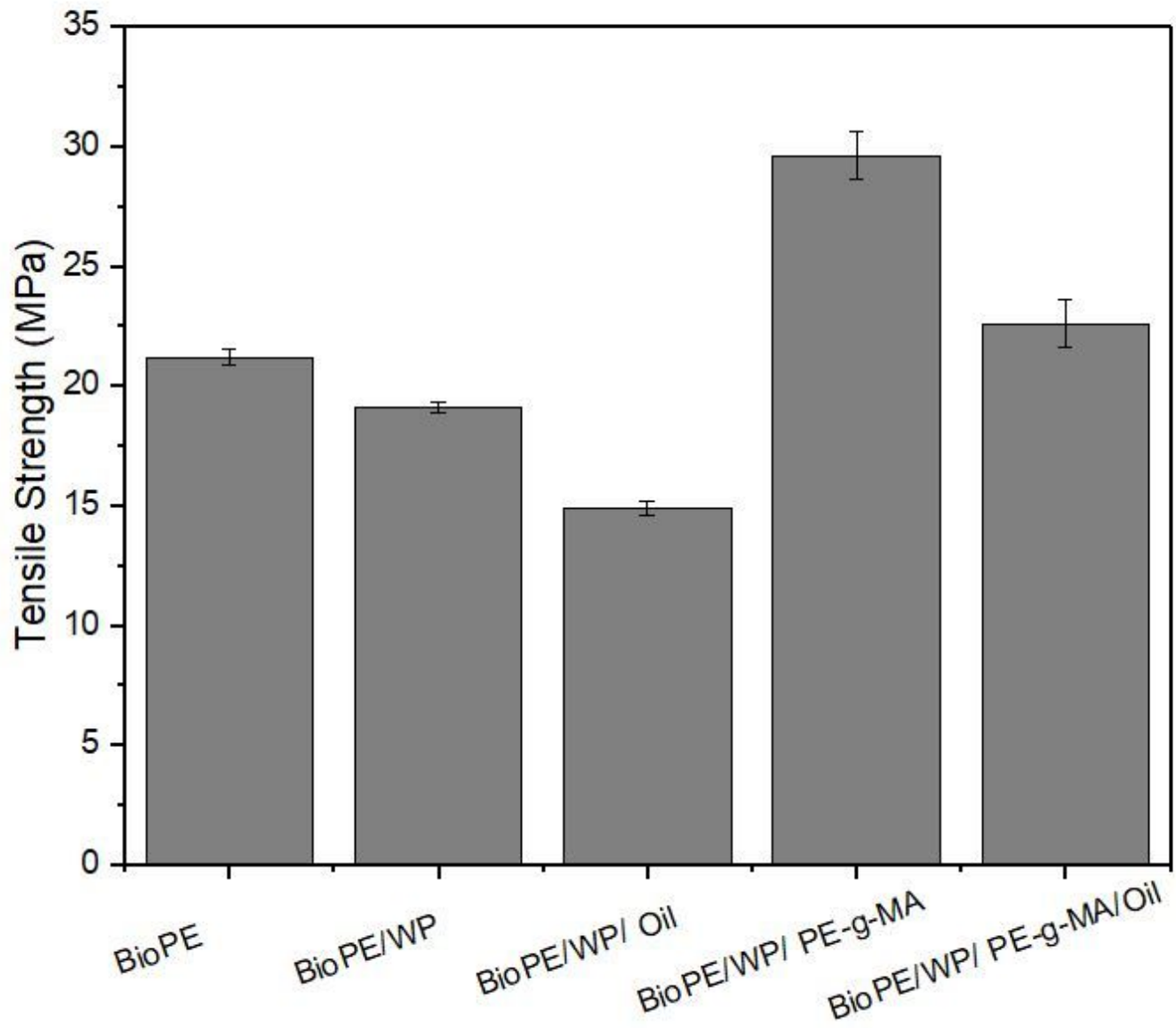


Figure 7

Tensile strength of BioPE and composites, with and without macaíba oil.

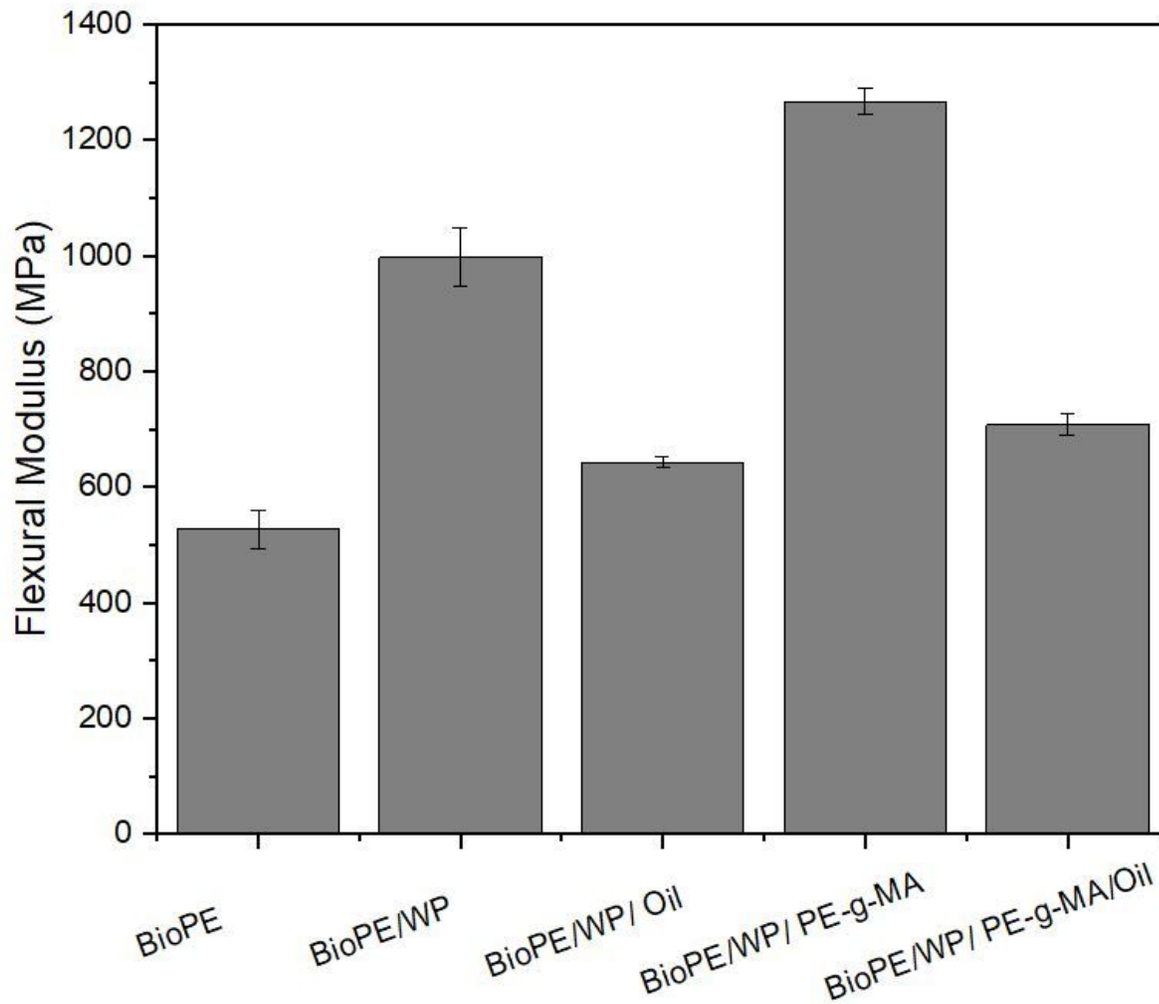


Figure 8

Flexural modulus of BioPE and composites, with and without macaíba oil.

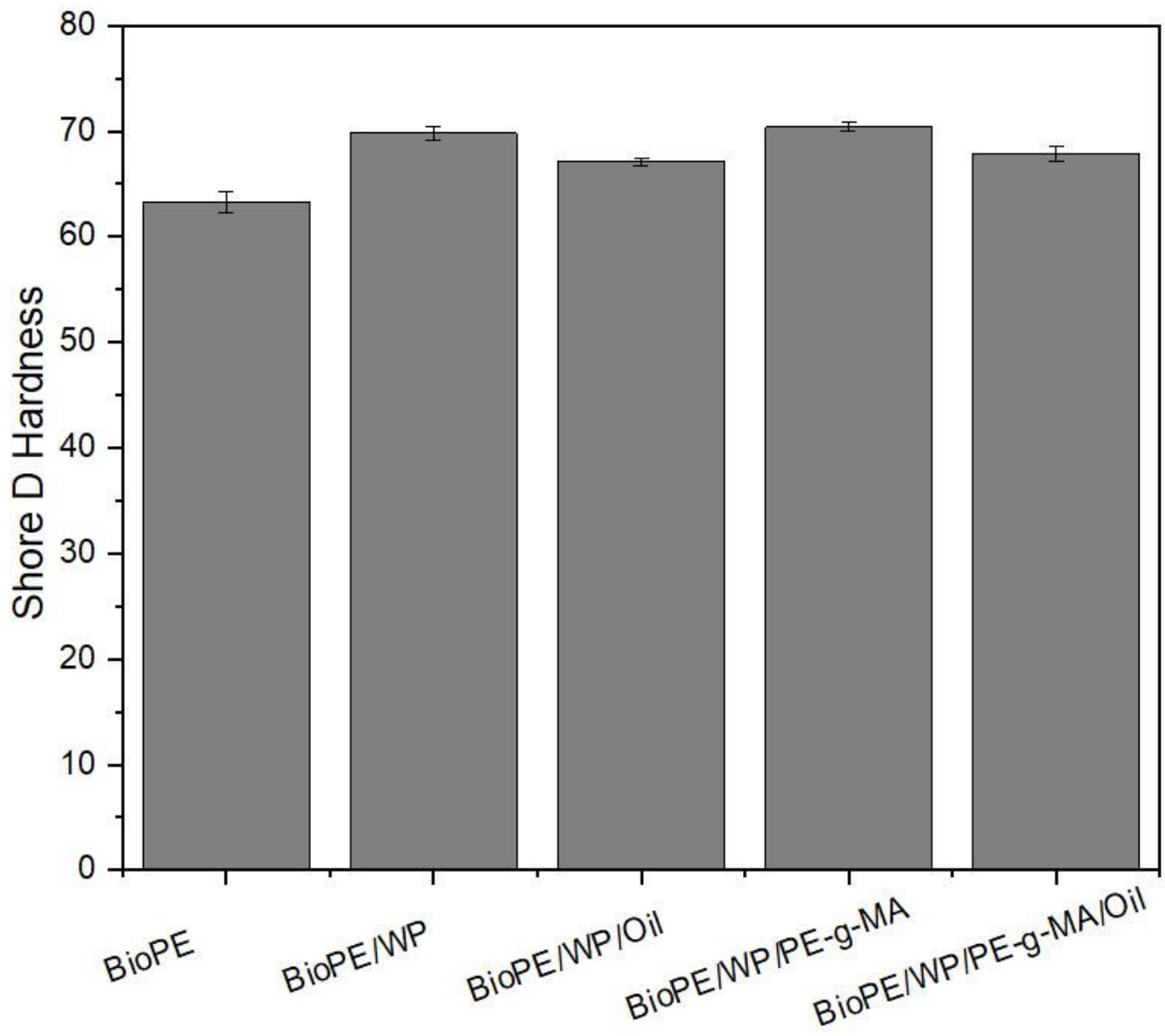


Figure 9

Shore D hardness of BioPE and composites, with and without macaiba oil.

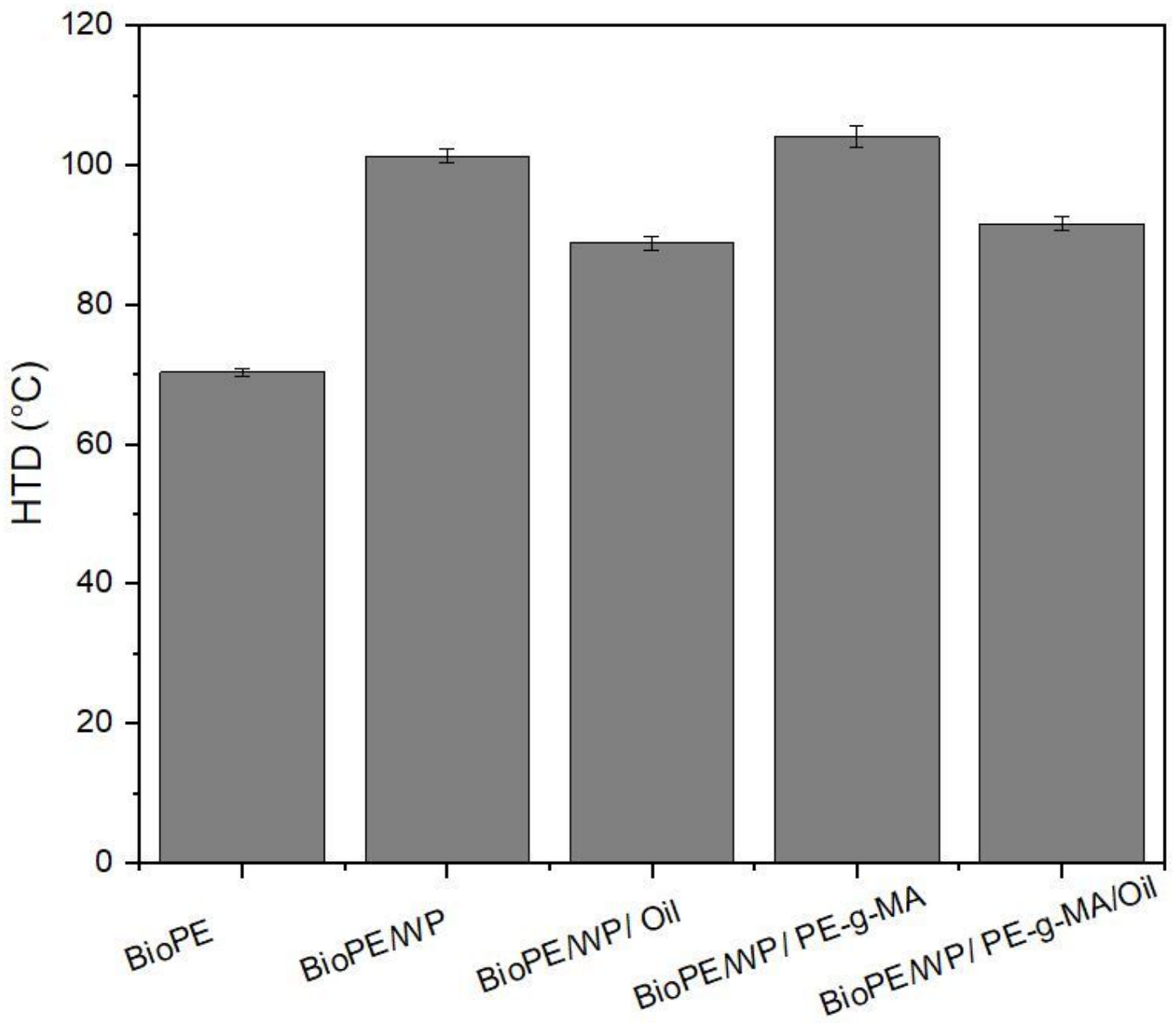


Figure 10

Heat deflection temperature of BioPE and composites, with and without macaíba oil.

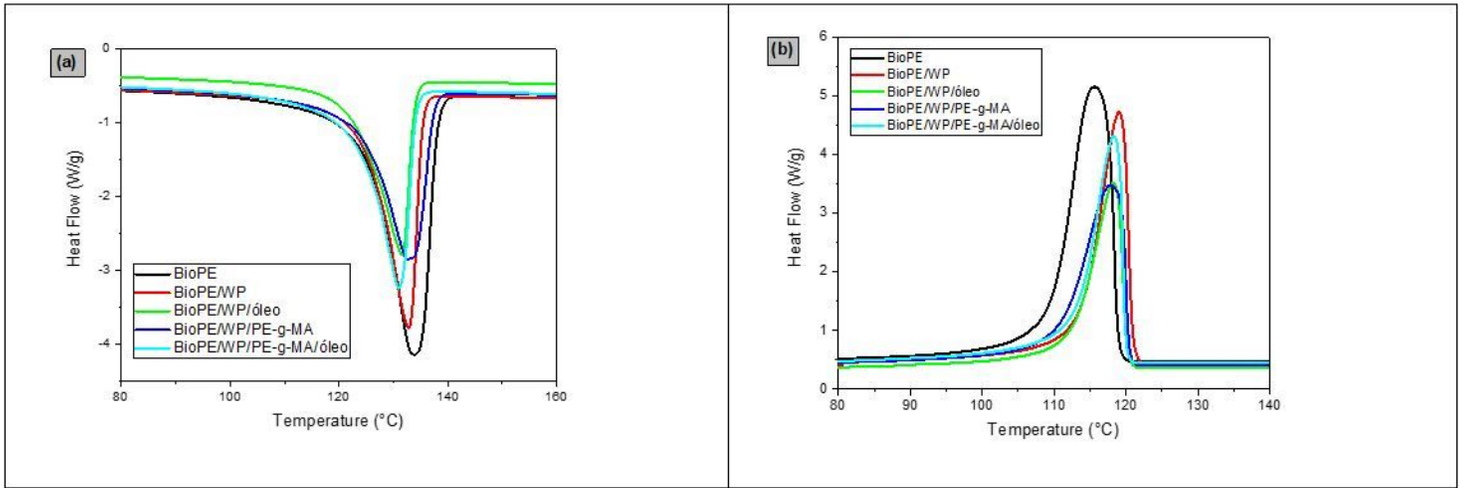


Figure 11

DSC curves for neat BioPE and composites: (a) melting temperature and (b) crystallization temperature.

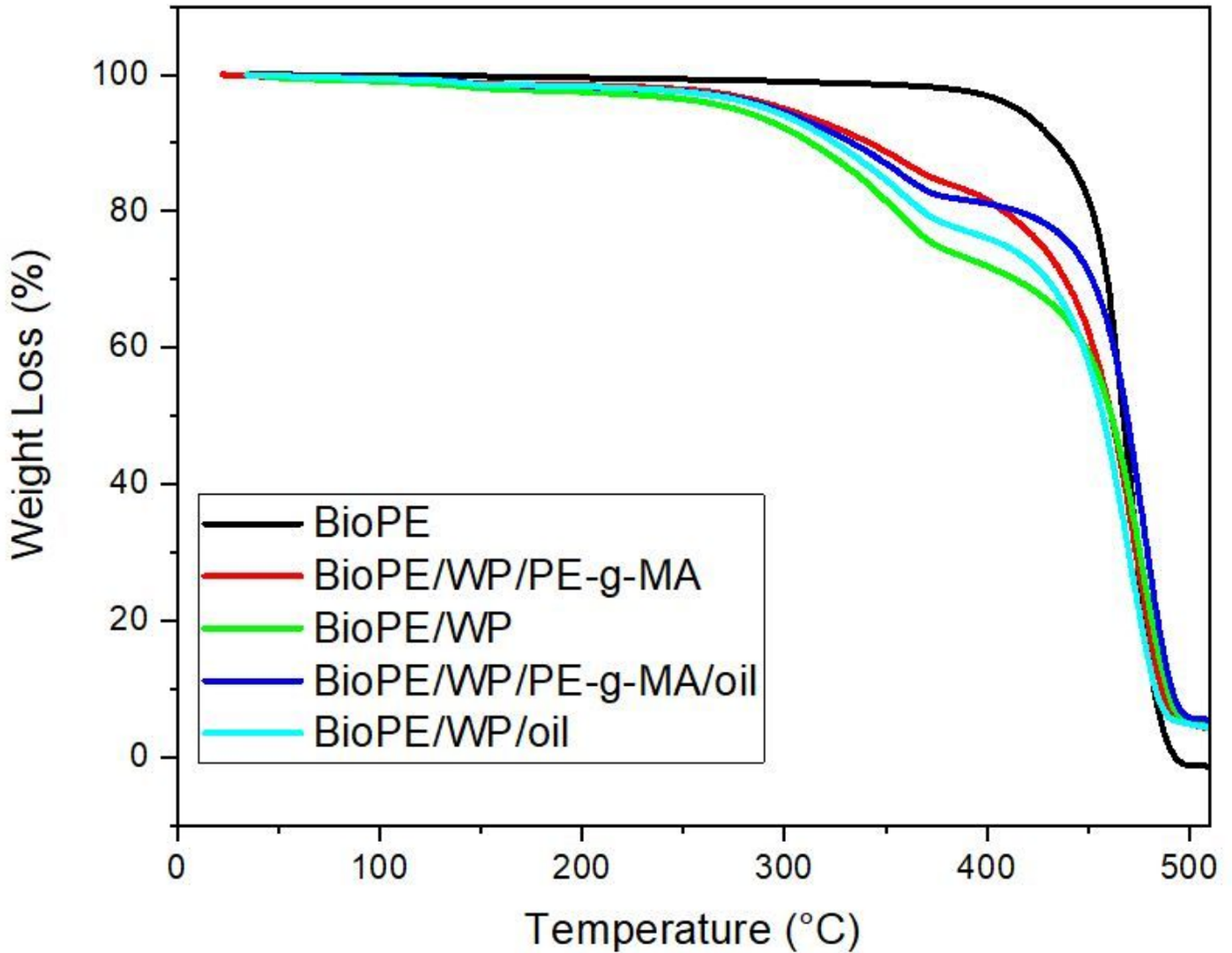


Figure 12

TG's curves of BioPE and composites, with and without macaiba oil, respectively.

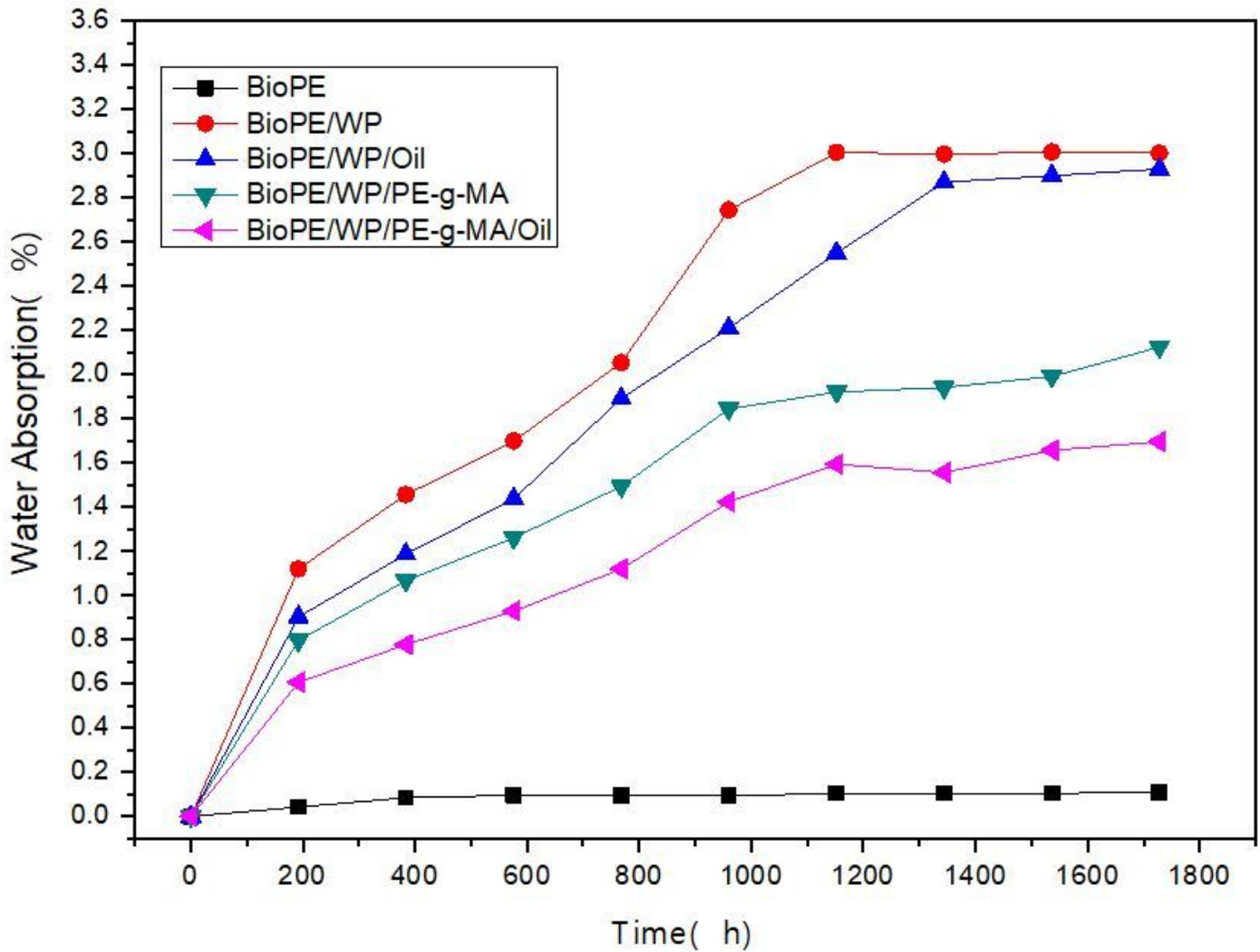


Figure 13

Water absorption of BioPE and composites, with and without macaiba oil, respectively.