. Alkali Treatment On Sugarcane Bagasse To Improve Properties Of Green Composites Of Sugarcane Bagasse Fibers-Polypropylene

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Submission date: 05-Dec-2020 10:51PM (UTC+0700)

Submission ID: 1465721153

File name: f Green Composites Of Sugarcane Bagasse Fibers-Polypropylene.pdf (1.05M)

Word count: 3886

Character count: 20247

ALKALI TREATMENT ON SUGARCANE BAGASSE TO IMPROVE PROPERTIES OF GREEN COMPOSITES OF SUGARCANE BAGASSE FIBERS-POLYPROPYLENE

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ABSTRACT

This work evaluated the improved performance of sugarcane bagasse fibers-polypropylene (PP) composites as a result of alkali treatment done on sugarcane bagasse as reinforcing fibers. Alkali solution of 10% NaOH with a temperature of 60-70°C was used to soak the bagasse for 2 to 6 hours. Hot pressed samples were prepared from a mixture of sugarcane bagasse fibers and PP with fibers/PP weight ratios of 20/80, 25/75, and 30/70. Hot pressed samples using untreated bagasse fibers were also produced for a comgrison. Tensile test was performed to all samples and the fracture surface from tensile test samples were observed using scanning electron microscopy (SEM) to study how the presence of reinforcement in PP matrix and its plali treatment influenced the strengthening and failure characteristics of the composites. The alkali treatment carried out resulted in a relevant increase of tensile strength of the composite to 229,29 MPa at 20/80 fibers/PP weight ratio or an increas 43% comparing to the composite samples which their fibers were untreated with alkali. The tensile strength of the composite with a 25/75 fibers/PP weight ratio increased to 290,30 MPa and decreased to 214,17 MPa as the fibers/PP weight ratio increased to 30/70. Microstructural study using SEM showed a different outlook on the fiber's surface after alkali treatment which the treatment caused the elimination of the impurities, waxy materials, lignin as well as hemicellulose, which was responsible for an easy deformation of the cellular networks. Alkali treated-sugarcane bagasse gave the best reinforcement for composites after 2 hours of treatment which resulted in the excellent fiberresin bonding.

Keywords: sugarcane bagasse, alkali treatment, fiber-resin bonding

INTRODUCTION

The use of green composites for engineering applications has gained high attention from industries in recent years as their potential to reduce waste from non-degradable synthetic materials and their carbon footprint. These composites using agro-fibers as reinforcement of thermoplastic materials are attractive to the automotive industry for light weighting of components (such as for automobile interiors) while much taining strength and cost-competitiveness, leading to improved fuel efficiency as well as to satisfy the public demand for 'green cars'. [1,2] Natural fiber composites are also suitable materials to make panels, ceilings, blocks, and partition boards to substitute wood, flooring tiles, etc. in building and construction industry. In comparison with synthetic fibers like glass and carbon, these natural files are gaining importance due to its many advantages, such as environment-friendly, reduced greenhouse gas emissions, low energy consumption, low cost, low density, and acceptable

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specific strength properties. [3-6] Indonesia has abundant availability of natural fibres such as pineapple, ramie, coir, bamboo, banana, and sugarcane. Their use can provide value added novel applications of natural fibers based composites and at the end it gives social benefits and major income to communities who cultivate them. This work focused on the potential of wester fibers from sugarcane named bagasse when used as reinforcement in polypropylene matrix. Bagasse is a fibrous residue and about 30% that remains in a sugarcane mill after crushing the sugarcane stalk and extracting of its juice. [7] It contains short fibers (Figure 1) and consists of water, and small amounts of soluble solids. [8,9] Contribution percentage of each of these components varies according to the variety, maturity, method of harvesting and the efficiency of the crashing plant. The bagasse, thus, produced after juice extraction poses a disposal problem and it is one of the largest agriculture residues in the world. Bagasse is mainly used as a burning raw material (85%) in sugarcane mill furnaces. [9] However the efficiency of the burning process is low due to low caloric power of bagasse. The excess of bagasse is deposited on empty field altering the landscape. Therefore its existence presents a serious environmental problem which many researchers have performed research work to develop technology, procedure or method that may help to exploit their efficient use. Studies on the impacts, performances and applications of Sugar Cane Bagasse in its original condition; transformed forms; treated with various chemicals and/or processes [10-14]; in combination with materials of distinct properties [15-19], and manipulation of manufacturing methodologies [20] have been explored and reviewed. This paper focused on the investigation to develop bagasse fiber reinforced composite material with 20 wt%, 25 wt%, and 30 wt% of bagasse fiber mixed in polypropylene matrix. On the basis of previous works which alkali treatment using NaOH has been used to modify other natural fibers, surface, [21] the same treatment was employed to bagasse fibers in this work. Their surface adhesive bonding to polypropylene resin or their reinforcement capacity to obtain the maximum strength was investigated.

EXPERIMENTAL PROCEDURES

(1) Materials Preparation

The experiment was initiated from the preparation of the sugarcane bagasse (Figure 1a). The bagasse contains of bacteria which can cause unpleasant smell. To remove that unpleasant smell, the bagasse was submerged in ethanol of 90% for 1 hour and then they were dried up in air in room temperature for about 6 hours before going to an oven drying at temperature of 200° C for 30 seconds. Figure 1b shows dried bagasse fibers after oven drying. A chemical treatment was performed on the bagasse by submerging the fibers 300 ml 10% NaOH for 2, 4, and 6 hours at temperature of $60 - 70^{\circ}$ C. Then they were rinsed with distilled water to cleanse them from NaOH excess on the surface then followed by drying up in an oven at temperature of 200° C for 30 seconds. Figure 15 shows bagasse fibers after alkali treatment. The fibers were then wrapped in aluminum foil to prevent it from moisture sorption from surrounding air and moist.



Figure 1 Bagasse fibers a) its original form, b) dried fibers after alcohol cleaning, and c) dried fibers after alkali treatment

Polypropylene (PP) used in this research was in fibers form as shown in Figure 2. The PP fibers were obtained from PT. Classic Prima Carpet Industries (Indonesia) which use the fibers as materials to make carpet. Those fibers were cut to obtain a size of 1 mm in length. That size was chosen to ensure that PP fibers were able to be well mixed with bagasse.

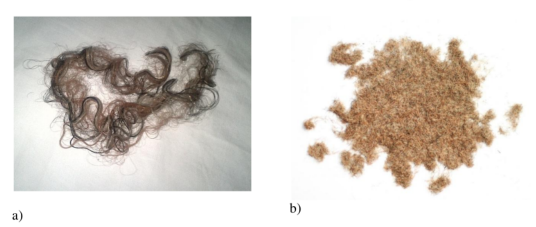


Figure 2 Polypropylene fibers a) in a continuous fiber and b) in 1 mm length before mixing with bagasse fibres

Bagasse and PP fibers in different weight ratio (Table 1) were mixed using a mixer of *Signora* brand, Jumbo Mix type with rotation speed of 17.500 rpm for 3 minutes. Flat type blade was used to prevent the fibers from being fragmented during mixing. A mixture of each different weight ratio of bagasse-PP fibers was hot pressed in a size of 100 mm x 200 mm using a pressure of 9800 Pa which was held for 3 minutes at temperature of 175°C. The hot-pressed samples were cooled in air for half an hour to let the curing of PP to take place.

(2) Materials Characterisation

Tensile test specimens (Figure 3) were prepared from those hot pressed samples in accordance with ASTM D638M-03 [22] and they were identified as shown in Table 1. Tensile test was performed using Gotech tensile test machine with a capacity of 30 tonnes. To obtain a better understanding of the surface adhesive bonding at the interface of the bagasse-PP fibers as well as to learn the sectiveness of the alkali treatment in modifying fibre surface properties, A Carl Zeiss SEM was used to observe the surface morphology and microstructure of untreated and alkali treated bagasse fibres. Fracture surface of the tensile test specimens were also studied using SEM to investigate the interfacial properties between bagasse fibers and PP matrix.

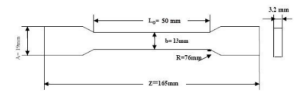


Figure 3 Tensile test specimen

Table 1. Samples Treatment, Weight Ratio, and Identification

Sample Code	Weight of Bagasse Fibers (g)	Weight of PP fibers (g)	Bagasse:PP Weight Ratio (%)	Alkali Treatment
A _i				None
A_j	4	16	20/80	2 hours
A_k				4 hours
A _l				6 hours
B _i	5	15	25/75	None
\mathbf{B}_{j}				2 hours
\mathbf{B}_{k}				4 hours
\mathbf{B}_{1}				6 hours
C _i			30/70	None
C _j	6	14		2 hours
C_k				4 hours
Cı				6 hours

RESULTS AND DISCUSSION

(1) Thickness Dimension

The thickness of hot pressed bagasse fibers-PP composites prepared from various bagasse:PP weight ratios and different soaking time in alkali was measured using a caliper. The thickness measurer tent was shown in Figure 4. The chart indicates that the thickness increment was in accordance with the increase weight of bagasse fiber in the composite and the decrease of soaking time of bagasse fiber in NaOH. Adding more bagasse fibers into PP matrix causing the stacking layer of fibers in the composite increased thus it increased the thickness of the composite. The lowest thickness of 2,17 mm was obtained in sample A_i and the highest thickness was obtained from sample C_i with 3,81 mm thick. Samples with "j", "k", and "l" codes (samples with their bagasse fibers were soaked in NaOH) had decrease thickness compared with the thickness of samples with "i" code (their bagasse fibers were not soaked in NaOH). Those thickness data informed that the longer time the fibres were soaked in NaOH, the thickness of the composites obtained got decrease. That happened because alkali treatment in NaOH causing the surface of the bagasse fibers was cleaned by removing hemicellulose and lignin which leads to a decrease in diameter and an increase in fibers crystallinity. [21]

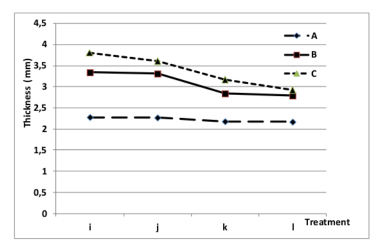


Figure 4 Composite's thickness against baggase:PP weight ratio and bagasse soaking time in NaOH.

(2) Tensile Properties

Tensile test were performed to three samples of each composition and treatments. The test results were compared to learn the effect of the increase of bagasse to PP weight ratio and soaking time on tensile strength. Most samples tested were failed in the gage section except one sample from C_i , C_k , and C_l respectively was failed outside the gage area. Figure 5 shows that in comparison with tensile strength of the composite samples A_i , B_i , and C_i (using untreated bagasse fibers), the strength of samples $A_{j,k,l}$, $B_{j,k,l}$, and $C_{j,k,l}$, was improved as increments of 15-43%; 31-80%; 43-61% respectively were identified if the bagasse fibers were treated with NaOH. The highest tensile strength was identified in B_j sample with bagasse:PP weight ratio of 25/75 and its bagasse experienced 2 hours alkali treatment in NaOH. A trend of decrease in tensile strength is shown in the graph in Figure 5 with respect to the soaking time in NaOH

which the lowest strength was identified in samples coded "I" which their bagasse fibers were soaked for 6 hours in NaOH. However this value of strength is still higher 15% to 43% compared to samples with untreated bagasse fibers (coded 'i'). Highest tensile strength of the composite was achieved when soaking time of the bagasse in NaOH was performed for 2 hours. Soaking time more than 2 hours, i.e. 4 and 6 hours caused the tensile strength to decrease. This is relevant with the microstructural observation on the fracture surface area of those samples which will be explained later. Other interesting findings from tensile test data were as the amount of bagasse fibers increased in the composite more than 25% resulted in the decrease of the composite's tensile strength. Composites samples C which had bagasse fibers of 30 wt.% experienced a decrease in strength as shown in Figure 5. Similar findings were also identified by Rout *et al* who studied composites of polyester reinforced with coir fibres. [23]

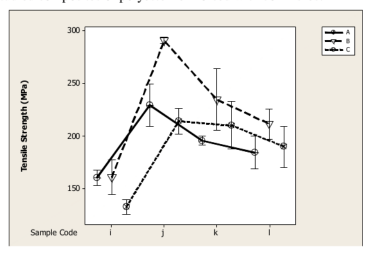


Figure 5 Tensile strength of Bagasse fibers – PP composites in various bagasse: PP weight ratio and soaking time

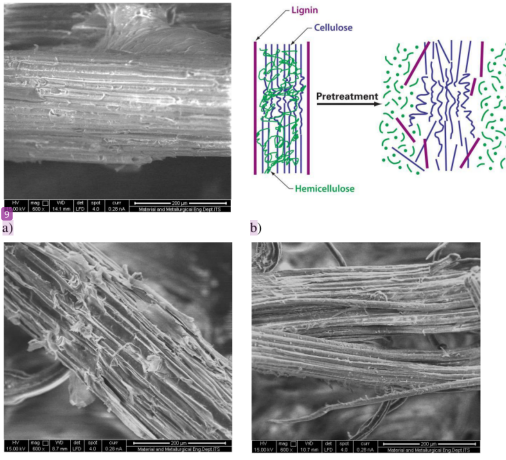
(3) Microstructural Study

Sugarcane Fiber Structure

To confirm the effect of NaOH treatment on morphological characteristics of bagasse fibers, the samples were studied using SEM. Figure 6 shows that differences are apparent on surface morphology between untreated and treated fibres.

The components of bagasse include cellulose 43.8%, hemicellulose 28.6%, lignin 23.5%, ash 1.3%, and other components 2.8%. [20] Cellulose is a semi crystalline polysaccharide made up of D-glucopyranose units linked together by β -(1-4)-glucosidic bonds [24] and the large amount of hydroxyl group in cellulose gives natural fiber hydrophilic properties. Therefore when the fibres are used to reinforce hydrophobic matrix, it results in a very poor interface and poor resistance to moisture absorption. [25] Hemicellulose is strongly bound to cellulose fibrils presumably by hydrogen bonds. Hemicellulosic polymers are branched, fully amorphous and have a significantly lower molecular weight than cellulose. Because of its open structure containing many hydroxyl and acetyl groups, hemicellulose is partly soluble in water and hygroscopic [26]. Lignins are amorphous, highly complex, mainly aromatic, polymers of phenyl propane units [13] but have the least water sorption of the natural fiber components. [26]

Figure 6a shows micrograph of untreated bagasse fibers which a superficial layer was evident. The cellulose microfibrils were still intact with a layer of fatty tissue and sugarcane lignin on the fiber surface. To optimize the interfacial properties between fiber and polymer matrix due to hydrophilic nature of natural fibers, a chemical modifications using 10% NaOH solution was used to eliminate lignin and hemicellulose on fibers surface. A significant change in morphological structure of the bagasse fiber occurred after treatment with NaOH for 2 hours (Figure 6c). Figure 6c, d, e show a highly fibrillated surface with a decrease of fibres diameter or an increase of the surface area due to the cleaning of the bagasse surface from impurities, wax cuticle layer, and hemicellulose structures which begin to dissolve in the solvent due to the nature of the alkali-soluble hemicellulose in alkali. The fibrils were starting to separate from each other as shown in Figures 6d and 6e because the lignin, the cementing component, had been removed by the action of the alkaline solution leading to an increase of the surface area and potentially improving the per-matrix adhesion in the composite. [21] Alkali treatment also affects the fibers' structure by solubilizing hemicellulose, reducing crystallinity, and increasing the available surface area and its pore volume. Figure 6b shows an illustration of how that treatment acts on the matrix of cellulose, hemicelluloses, and lignin. [27] However if the soaking time in NaOH was too long, as shown by tensile test that the strength of the composite became lower due to the degradation of cellulose structure. [21]



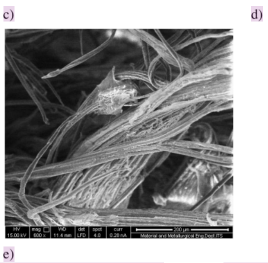


Figure 6 SEM micrographs of bagasse fibers (a) untreated fibers (c) fibers after NaOH treatmentfor 2 hours (d) fibers after NaOH treatment for 4 hours (e) fibers after NaOH treatment for 6 hours; (b) schematic structure of a natural fiber before and after a treatment [27]

Structure of Fracture Surface

To support the strength data from tensile test, SEM analysis on the fracture surface was performed. Fracture surface of samples A_i and A_j was studied to study the change in interfacial properties as an effect of NaOH treatment of the bagasse fibers. As it has been identified by tensile test that the increase of bagasse fibers more than 25 wt.% caused the decrease in tensile strength of the composite. Thus the effect of bagasse content on strength needs to be investigated further. Therefore SEM observation was also done on fracture surface of samples B_i and C_i to get a better understant on the relationship between the quantity of bagasse fibers in the composite with its strength. SEM images of fracture surface of composite samples A_i which the bagasse fibers used were untreated and had the lowest strength (Figure 5) is shown in Figure 7a. This low strength resulted from poor interface adhesion between the bagasse fibers with PP matrix. Fiberpull out is the evidence of that less intimate interfacial contact due to hydrophilicity of bagasse fibers which results in moisture absorption and weak adhesion to PP materials. Low content of bagasse fibers (20 wt.%) in sampe A_i caused the thickness obtained was also low compared to samples B_i and C_i (Figure 4, 7c, 7d).

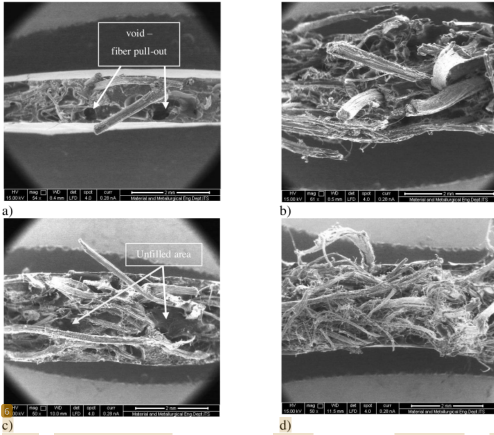


Figure 7 SEM micrographs of samples fracture surface cross section of a) sample A_i, b) sample A_i, c) sample B_i, and d) sample C_i

Composite's strength made from alkali treated fibers was superior compared to the composite made from untreated fibers. There was 43% improvement in tensile strength found in sample A_j (229,29 MPa) compared to sample A_i (160,81 MPa). Figure 7b shows the rough surface of fibers due to the elimination of the external layer of the fibers during alkali treatment therefore it provides better adhesion between fibers and matrix. Splitting of fibers in multiple area was evidence of this improved interface properties.

evidence of this improved interface properties.

There was no significant difference between SEM images of fracture surface of sample A_i (Figure 7a) and sample B_i (Figure 7c) except from higher bagasse fibers content in sample B_i. It was observed that there was an empty area which was not filled or wetted by PP resin due to bagasse hydrophilic properties. The strength of sample B_i is 161,01 MPa which the strength did not change significantly by adding more bagasse fibers from 20 to 25 wt.% unless alkali treatment was applied. The increment of bagasse fibres to 30 wt.% was not a solution to improve composite strength. The strength obtained from tensile test on sample C_i (133,11 MPa) decreases 21% compared to the strength of sample Bi when bagasse fibres used was increased to 30 wt.%. This happened because of the shortage of PP resin which bonded the bagasse fibers as shown from the SEM 22 crograph in Figure 7d. Therefore the maximum strength in this work was obtained when the weight ratio of bagasse fibers:PP is 25/75.

CONCLUSION

- 1. Increase soaking time of bagasse fibers in 10% NaOH solution results in the reduction of composite thickness. A decrease in thickness follows the increase of the amount of bagasse fibers in the composite as well as the increase of soaking time of bagasse in NaOH solution. The biggest reduction in thickness (23%) takes place in sample C₁ which the bagasse: PP weight ratio is 30/70 and the bagasse was soaked for 6 hours. The reduction in thickness was due to alkali treatment resulted in the loss of existing pitch surface thus reducing the diameter of fibers.
- 2. The highest tensile strength (290.30 MPa) was obtained in sample B_j with bagasse:PP weight ratio of 25/75 and the lowest tensile strength of 133.11 MPa was found with sample C_i. An increase of the amount of bagasse fibers more than 25 wt.% was not a solution to improve tensile strength of the composite. A treatment using 10% NaOH with a duration of phours performed on bagasse fibers before mixing them with PP fibers was a good way to improve good adhesion between bagasse fibers and PP matrix due to the removal of hemicellulose and lignin which resisted the wetting of PP on the fibers.

ACKNOWLEDGEMENT

The authors are grateful for the assistance of Is from Material and Metallurgical Engineering Department of Technological Institute of Sepuluh November (ITS) in performing SEM observation.

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