Research Article

Effect of Chemical Treatment on Physical, Mechanical and Thermal Properties of Ladies Finger Natural Fiber

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Received 2 May 2013; Accepted 4 September 2013

1. Introduction

Natural fibers are gaining progressive account as renewable, environmentally acceptable, and biodegradable starting material for industrial applications, technical textiles, composites, pulp, and paper, as well as for civil engineering and building activities. Natural fibers reinforced composites combine acceptable mechanical properties with a low density [1]. In recent years, wide range of research has been carried out on fiber reinforced polymer composites [2–11]. Typical reinforcements for composites with plastic matrix are various synthetic fibers such as glass, graphite, boron, metallic, and ceramic materials. These materials are heavy, expensive, and harmful to environment. The replacement of inorganic fibers with comparable natural fibers provides weight and cost reduction. The advantages of using natural fibers also include high specific stiffness and mechanical strength, availability, reduced energy consumption, low hardness which minimizes the wear of processing equipment, renewability, recyclability, nonhazard, and biodegradability. Natural fiber reinforced composites are suitably applicable for aerospace, leisure, construction, sport, packaging, and automotive industries [12–14]. However, the main drawback of natural fiber reinforced polymer composites is the incompatibility between the hydrophilic natural fiber and hydrophobic matrices. This leads to undesirable properties of the composites. It is therefore necessary to modify the fiber surface by employing chemical modifications to improve the adhesion between the fiber and matrix [15–18]. The objective of present research is to characterize raw and chemically treated ladies finger natural fibers by finding out their physical, mechanical, and thermal properties. The properties of raw ladies finger fiber are also compared with those of chemically treated ones.

2. Experimental Procedure

2.1. Chemical Treatment of Fiber. Three types of chemical treatments were performed on the ladies finger fiber. Fiber was extracted from the stem of a ladies figure tree (Figure 1). For alkali treatment, the fiber was treated with 2% NaOH solution at 70°C for about two and a half hours. A single stage treatment (SST), which is basic chromium sulfate solution treatment (pH 4), was performed by a 3-hour shaking of fiber...
in 4% Cr₂(SO₄)₃ • 12(H₂O) solution. The double stage treatment (DST) was performed by continuing the 4% Cr₂(SO₄)₃ • 12(H₂O) treatment with the addition of NaHCO₃ solution for another 2 hours.

2.2. Tensile Test. Tensile test of single ladies finger fiber was carried out by varying span length at 5, 15, 25, and 35 mm using a tensile testing machine. The fibers were glued in a paper frame (Figure 2) to ensure a good gripping and straight direction to the test clamps. Diameter of single fiber was measured by using scanning electron microscopy (SEM). The paper frame was carefully placed in between the jaws of the load cell. Load cell of 50N was used, and crosshead speed was 4 mm/min.

2.3. FTIR Spectroscopy. The infrared spectra of fiber were recorded on a Nicolet 380 spectrophotometer with coaddition of 32 scans. Powdered sample was taken for FTIR spectroscopy. Then potassium bromide (KBr), which acts as a reagent, was mixed (KBr: sample = 100:1) with them in a mortar pestle. The mixture was then taken in a dice of specific dimensions. The pellet was formed by pressing with a hand press machine and was placed on the sample holder. The IR spectrum obtained is presented in the Results and Discussion section.

2.4. Scanning Electron Microscopy. Surface morphology of the raw and chemically treated fibers was observed under a scanning electron microscope (Philips XL 30). The fiber surface was initially made conductive by applying gold coating using a sputtering machine. The fiber was then taken inside SEM, vacuum was created, and micrographs were taken. The diameter of the fiber was also measured using the same microscope.

2.5. Thermogravimetric Analysis. Thermogravimetric analysis was carried out for determining thermal stability of ladies finger fiber. TGA method used was based on continuous measurement of weight on a sensitive balance (called a thermobalance) as sample temperature was increased in an inert atmosphere. This is referred to as nonisothermal TGA. Data were recorded as a thermogram of weight versus temperature.

3. Results and Discussion

3.1. Tensile Properties. Tensile properties (Young’s modulus and tensile strength) of both raw and chemically treated single fibers were obtained from stress/strain curves. In order to remove the grip effect during tensile test, the Young’s modulus values were corrected using a method described in [19]. The variation of Young’s modulus and tensile strength is shown in Figures 3 and 4, respectively. The tensile strength decreased, while Young’s modulus increased with increase in span length [20, 21]. As mentioned by Bledzki and Gassan [22], the longer the stressed distance of the natural fiber, the more inhomogeneities (flaws) in the stressed fiber segment that weaken the structure. Thus increasing fiber length inevitably reduces the strength of the fiber.

During alkali treatment, hemicellulose and lignin were removed. The interfibrillar region is likely to be less dense and less rigid making the fibrils rearrange themselves along the direction of tensile loading. When fibers were stretched, such arrangements among the fibrils resulted in better load sharing and hence higher stress development in the fiber [23].

During SST, the fibers were shaken for 3 hours in a 4% basic chromium sulfate (Cr₂(SO₄)₃ • 12(H₂O)) solution. The pH of the solution was maintained about 2–2.5. After 3 hours shaking, a thin coating layer was formed on the fiber surface due to chemical reaction between Cr₂(SO₄)₃ • 12(H₂O) and fiber (Figure 5).

Again for DST, the same procedure was followed with an additional 2 hours shaking with NaHCO₃ in the solution. In this case an even thicker coating was formed on the fiber surface due to chemical reaction between Cr₂(SO₄)₃ • 12(H₂O) and NaHCO₃ and fiber (Figure 6).

The chemical reactions between cellulose of ladies finger fiber and Cr₂(SO₄)₃ • 12(H₂O) and NaHCO₃ occurred in the following three stages.

(i) The chrome complexes reacted with the fiber cellulose carboxyl groups. During the first reaction, the solution was very strong and pH was around 2. The chromium held the first OH group:

\[
[Cr]^{3+} + OH^- = [Cr-OH]^{2+} \quad 33\% \text{ basicity pH } 2. \tag{1}
\]

(ii) As pH of the solution was increased, sulfate associated with the chromium became displaced by the hydroxyl
groups. Here the second OH group entered into the reaction with Cr at pH 3. During these reactions acidic nature of solution decreased and pH increased:

\[ [\text{Cr} \cdot \text{OH}]^{++} + \text{OH}^{-} \]
\[ = [\text{Cr} \cdot (\text{OH})_{2}]^{+} \quad 66\% \text{ basicity pH 2} \rightarrow \text{pH 4}. \]

(iii) As pH was increased to 4, the hydroxyl groups became shared by chromium atoms. When pH was increased to 8-9, the reaction was completed:

\[ [\text{Cr} \cdot (\text{OH})_{2}]^{+} + \text{OH}^{-} \]
\[ = [\text{Cr} \cdot (\text{OH})_{2}]^{0} \downarrow \quad 100\% \text{ basicity pH 4} \rightarrow \text{pH 8}. \]

After SST, the activity of chromium still remained incomplete. As a result, the tensile properties of ladies fiber increased less compared to the raw ones. However, after the DST, chromium became fully occupied in reducing hydroxyl groups, which in turn increased the tensile properties of ladies finger fiber compared to both raw and SST fibers.

3.2. FTIR Spectroscopic Analysis. FTIR spectra of raw and treated ladies finger fibers are shown in Figure 7. The IR spectrum for raw ladies finger fiber (Figure 7(a)) clearly shows the strong and broad characteristics band of (–OH) at the regions of 3600–3200 cm\(^{-1}\), lignin and hemicelluloses at about 1731.5 cm\(^{-1}\), and (C–H) aromatic rings and alkanes at 3292.5 cm\(^{-1}\) [24]. The alkali treated fiber (Figure 7(b)) shows the characteristics band of (–OH) of high concentration at around 3452.2 cm\(^{-1}\) and aromatic rings and alkanes at around 2921.8 cm\(^{-1}\). Peaks for lignin and hemicelluloses are not very significant. The basic Cr\(_2\)SO\(_4\) treatment showed absorption peak at wave number of 3446.7 cm\(^{-1}\) (Figure 7(c)). Double stage basic Cr\(_2\)SO\(_4\) and NaHCO\(_3\) treated fiber showed (–OH) stretching vibration at 3473.6 cm\(^{-1}\) wave number (Figure 7(d)).

3.3. Surface Morphology. Surface morphology of raw, alkali treated, SST, and DST ladies finger fiber was observed under SEM and is shown in Figure 8. The surface of the alkali treated ladies finger fiber was found more rough compared to the raw fiber due to the removal of hemicellulose and lignin from the fiber surface. During single stage treatment and double stage treatment, the surface roughness seemed to be a bit
3.4. Thermogravimetric Analysis. Thermogravimetric analysis was performed on the raw, alkali treated, SST, and DST ladies finger fiber in order to find the effect of chemical treatment on the thermal stability of the fiber. TGA curves of raw and treated ladies finger fibers are shown in Figure 9. There was a huge weight change at around 250°C in all four cases indicating the start of thermal decomposition of ladies finger fiber. Thus no significant difference in thermal stability was observed among the raw, alkali treated, SST, and DST fiber of ladies finger.

4. Conclusion

Present research demonstrated that chemical treatment clearly affected the mechanical, structural, and thermal properties of fiber obtained from ladies finger plant. The tensile strength of both raw and treated fibers decreased, while Young's modulus increased with span length. The surface of raw ladies finger fiber was rough. Alkali treatment increased the surface roughness by removing hydrophilic hemicellulose. Both single stage and double stage treatments seemed to
decrease the surface roughness by forming a coating layer on the fiber surface and improved chemical bonding with cellulose of fiber. As a result all chemical treatment increased the tensile properties of fiber. However, double stage chemical treatment showed better properties compared to alkali treatment and single stage treatment. No significant effect of treatment on thermal stability of fiber was observed from TGA.
References


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