Fabrication of Unidirectional Coir Fiber Reinforced Nonwoven Melt-Blown Glass Fabric Composite by Compression Molding

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Abstract
The unidirectional composites of Coir Fiber with nonwoven melt blown glass fabric (C-G) were fabricated using compression molding. The reinforcement fibers were chemically treated with NaOH and silane coupling agent and then parallel placed with each other to prepare the unidirectional preforms. The parallel fibers were placed onto the metallic frame and dried before molding. The C-G layers was placed in heated mold for consolidation to produce the unidirectional composite specimens. The heated mold was cooled by the cooling system of the compression machine. Consolidation process involved three stages. At first, under certain temperature, the solid fibrous matrix materials became softened and then the matrix in the liquid form soaked and infiltrated the reinforcing fibers and finally during cooling stage, the matrix turned into solid form to hold the fibers in a definite position. The molding time affected the mechanical properties of the unidirectional composite. The C-G composite were characterized in the term of mechanical properties, dynamic mechanical analysis, and Fourier transformed infrared spectroscopy. The optimum compression molding parameters was NaOH treated coir fiber at processing time 8 min having processing temperature 170°C for fabricating unidirectional composite.

Introduction
FRP usually referred to as fibre reinforced polymer composite is a quite new material in the various applications such as construction and building industry compared to concrete and steel [1]. Composites possess the desired and preferred properties by coalescing dissimilar constituents in a cautious and judicious way. Generally, they possess higher specific modulus and high specific strength enabling them as a valuable material in huge number of industrialized requests which requires such features [2]. Carbon fibres and glass fibres integrated in polyester resin are the traditional and conventional fibre reinforced composite materials. These composite materials have excellent mechanical properties, but these materials cause environmental pollution due to the non-degradability of fibres [3]. Natural fibre reinforced composites found to be an alternative solution to the ever-depleting petroleum sources thus they receive greater attention, attraction from research scientist and community. Manufacturer and scientists attracted towards natural fibre based composites due to its biodegradability, light in weight, nontoxic and relatively stronger and consider being virtuous products which can be used in construction industry, automotive industry and for furniture production [4]. Natural fibre composites have better formability, abundant, renewable, cost effective, possess tool wearing rates, thermal insulation properties, acoustic properties, sufficient energy requirements and safer towards health [5]. Many innumerable demerits such as hydrophilic in nature, poor fibre/matrix interfacial adhesion and poor thermal stability of natural fibres can be overcome by chemical treatment or compatibilizer which amended the adhesion between the fibre and matrix. Composite of polymers and Coir fibre possess the variances and incomparability in terms of their polarity structures [6]. Based on the origin natural fibres are categorized as animal based and plant based. Animal-based fibres are wool, silk,
etc. and natural fibres based on plant includes sisal, coir, ramie, jute, bamboo, pineapple and many more [2]. Lignocellulosic fibres possess many compensations of being financially reasonable to manufacture such as lightweight, eco-friendly, harmless to health, high stiffness and specific strength which provides a probable substitute to the synthetic or artificial fibres [7,8]. The reinforcing capability of the fibres mainly influenced by various aspects such as polarity of the fibre, mechanical strength of the fibres, surface appearances, and existence of reactive centers [9]. Moreover, many of the natural fibres properties are governed by several factors such as climate, harvest, maturity, variety, de- ecorations, retting degree, disintegration (steam explosion treatment, mechanical), fibre modification, technical and also textile processes (spinning and carding) [9]. In spite of these promising features shown by natural fibres certain major drawbacks are also underlined like water absorption, strength degradation, lack in thermal stability lowered impact properties [10,11] but it has been found that these can be improved and overcome by hybridization with either natural or synthetic fibre. Bast fibres derived from natural fibres such as coir, hemp, flax, kenaf and jute have high specific strength, low density and are extremely concerned in several industrial applications [12]. Coir fibres are gratifying increasingly widespread throughout the world as a significant natural materials source contributing towards the development of eco-friendly assets for the automotive, sports industries, food packaging and furniture [13], textiles, paper pulp, and fibre boards-based industries [14]. Inferior thermal resistance is displayed by coir as compared to artificial or synthetic fibres such as (aramid, glass fibres) like all other natural fibres [1]. Coir is in an advantageous position when compared with other lignocellulosic fibre crops since it has a short plantation cycle, flexibility to environmental conditions and requires relatively lowered quantity of pesticides and herbicides [15]. Coir fibres receive much attention owing to its prospective probability as polymer reinforcements in the natural fibre composite industry. Researchers claimed that mechanical strength and thermal properties of coir composite are superior to other type of natural fibre polymer composites, thus regarded as a suitable applicant for high-performance natural fibre polymer composites [14]. The coir fibre as reinforcement materials are full fledged noticeably from the past decade along with other products such as extruded plastic fencing, decking, and furniture padding [3]. Coir composites can eventually, supplement and substitute petroleum based composite materials in many of the known industrial applications, and thus proposing innovative environmental, agricultural, manufacturing and consumer profits formulation methods [16,17].

**Experimental**

**Materials and methods**

**Materials**: Coir fiber has been collected from Gazipur District (Bangladesh). After collection, the fibers were washed and dried in order to remove impurities like sand, leaves etc. Melt-blown nonwoven glass fiber fabric was collected from Shanghai Macklin Biochemical Co. Ltd.

**Chemical modification of coir fiber**: Coir fibers were soaked in a solution of NaOH and silane coupling agent (7% NaOH and 7% silane coupling, 7 g/100 mL of water) for maximum 3h at room temperature. After 3h, ramie fibers were rinsed in tap water with a drop of acetic acid until excess NaOH was washed way.

**Composite fabrication**: The melt-blown glass sheets were cut to the desired size (12 cm × 12 cm) for composite manufacturing. Coir fibers were cut into the length of 12 cm. Two layers of unidirectional fibers were inserted between three sheets of pre-weighted glass sheets during the fabrication of composite. Fibers volume ratio was chosen to be 3:7. The fibers were embedded randomly in between glass sheets. By this way, a sandwich is formed which was then employed between two steel plates under a pressure of 5 MPa for 8 min with keeping the temperature of 170°C. The thickness of the resultant composites was kept 2 mm. Testing specimens were prepared from the composite sheet by cutting with grinding machine carefully. Figure 1 shows the schematic fabrication layout of composite laminates (Table 1). Represents the sample specifications of different composites.

**Mechanical Tests**

Tensile strength was investigated by following the DIN 53455 standard method using a Hounsfield S series Universal Testing Machine, model: H 50 KS-0404. The cross-head speed was set 10 mm/min during testing and the gauge length was 20 mm. The
geometry of the test specimen was maintained 60mm × 15mm × 2mm. The data was recorded at room temperature (20°C) and constant relative humidity of 65%.

**Dynamic mechanical analysis (DMA)**

The dynamic mechanical analysis test was performed in a tension mode on DMA METTLER TOLEDO (DMA star system, made in the USA). The size of the specimen was cut to a dimension of 15 mm × 5 mm × 3 mm and testing was performed in a temperature range of 30–200°C with a heating rate of 3 m/min.

**Surface analysis**

The Fourier transform infrared spectroscopy, FTIR (Perkin Elmer) analysis was used to evaluate the changes of surfaces happened to ramie fibers before and after treatments. Before testing, the prepared samples were mixed with Potassium Bromide (KBr) powder. Then, the FTIR spectra were taken for all samples in a range of 500–4000 cm⁻¹.

**Thermal analysis**

The Thermogravimetric Analysis (TGA4000) was taken to analyze the thermal degradation procedure of C-G composite within a temperature range of 30–800°C. TGA characterization was carried out in a nitrogen gas media at a heating rate of 20°C min⁻¹, in a sealed vessel.

**Result and Discussion**

**Mechanical tests**

Suitable working parameters are quite essential to produce outstanding products with the required physical and chemical qualities wanted. In this case, the combined processing condition of C2 is more significant. The tensile strength of composite materials is straightly depending on the strength and modulus of the reinforcing fibers, orientation and length of the fiber, fiber loading,
as well as fiber-matrix interfacial adhesion. These variable factors can explain the circumstances that the mechanical properties obtained for C-G composite at higher processing temperature are lower. Other reason can be tensile strength variation of melt-blown glass sheet as it melts and gets brittle at higher temperature. The effect of alkali, silane can be seen from the results of tensile strength. Whatever the processing condition, the modification of alkali or silane treatment has greatly improved the tensile strength of C-G composites. Figure 3 represents the tensile modulus of C-G composites of different processing conditions.

**Dynamic mechanical analysis (DMA)**

DMA test results provide the effect of elevated temperature on the stiffness of coir fibers reinforced glass composite. Here, the information can be obtained in the form of storage modulus, loss modulus, and tan δ. The temperature dependency (dynamic mechanical properties) of C-G composites are presented in Figure 4.

It can be clearly seen that the dynamic mechanical properties of the samples nearly showed a similar pattern. The storage moduli of tested samples closely mean to define the load bearing capacity of C-G composite. The storage modulus of C2 composite was higher than others. The higher value indicates the higher stiffness and load bearing capacity of coir fibers in the reinforcement system. Similar to the storage modulus, the loss modulus illustrates the maximum heat dissipation of C-G composite. The loss modulus of all samples showed a decrease with increasing temperature, and there was a significant fall in a region of 60-80°C. As it can be seen in from Figure 5, the maximum heat dissipation occurs at a maximum temperature where loss modulus was highest, indicating the glass transition (Tg) of the material. The ratio of loss modulus to storage modulus could illustrate the mechanical loss factor (tan δ) of C-G composites.

From Figure 6, the impact resistances of C-G composites were significantly improved by reinforcement of coir fibers.

**Surface analysis**

Figure 7 represents the FT-IR spectra of different composite samples. From the spectra study, it is evident that the absorption peaks of 3447 cm⁻¹ for all samples were allocated to OH stretching of α-cellulose [18,19]. Meanwhile, the absorption band at 2946 cm⁻¹ were attributed to CH and CH2 stretching vibrations [19]. Furthermore, vibration peaks observed at 1734 cm⁻¹ in the spectrum of the bast which denotes C=O stretching of methyl ester and carboxylic acid or the acetyl group in hemicelluloses was drastically decreased for C2 and C5 sample due to the successful removal of the pectin and hemicelluloses during their treatment with NaOH and silane [19]. On the other hand, the absorption band at 1636 cm⁻¹ of the C8 was drastically decreased and shifted to 1629 cm⁻¹ for C2 which is attributed to antisymmetric COO-stretching which proves the efficiency of NaOH treatment method to remove lignin from the bast fibers [18]. Moreover, absorption
peaks observed at 1375 cm⁻¹, which denotes CH₂ and CH symmetric bending mode, respectively [19]. Besides, absorption peaks in the fingerprint regions at 1164 cm⁻¹ and 1058 cm⁻¹ attributed to cellulose structure [18,19]. In addition, absorption peaks at 1311 cm⁻¹ and 898 cm⁻¹ were associated with C-H scissoring and C-C bending [20,21]. And, the small peaks at 832 cm⁻¹ ascribed to aromatic C-H out-of-plane vibration in the lignin was decreased in intensity after the degumming process [22]. Consequently, it very well may be reasoned that the absorption peaks of all the non-cellulose substances, for instance, hemicellulose, pectin, lignin, and wax-like substances, were missing or radically decreased for C2 than that of C5 and C8.

Figure 7: FT-IR spectra of composites.

**Thermal analysis**

![TGA curves of composites](image)

Figure 8 shows the thermogravimetric analysis curves of C5, C2, and C8 composite samples. It is evident from the curves that all samples depict two stages of thermal degradation patterns. Initial weight loss due to evaporation of water and other volatile compounds within the samples takes place between the temperature range 60-150°C. Cellulose decomposition for C8, C2, and C5 starts from 296.4°C, 313.9°C, and 325.3°C, respectively. The thermogravimetric analysis (TGA) results confirmed that the chemical treatment of coir fibers and addition of treated fibers to the nonwoven glass fabric increased the thermal resistance of the composite.

**Conclusion**

In this study, a composite containing coir and nonwoven melt blown glass fabric was manufactured by hot compression molding. Alkali treated modified coir fibers composite showed better mechanical properties with a significant improvement of surface compatibility over silane treated and untreated composites. Nine processing conditions were prepared to find the optimization properties of the composites. Accordingly, a suitable processing condition was selected as sample C2 with fibers volume ratio of 3:7, molding temperature of 170°C, a molding pressure of 5 MPa and 8 min of processing time, respectively.

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None.

**Conflict of Interest**

No conflict of interest.
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