PRE-TREATMENT OF FLAX FIBERS FOR USE IN ROTATIONALLY MOLDED BIOCOMPOSITES

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by

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ABSTRACT

Flax fibers can be used as environmentally friendly alternatives to conventional reinforcing fibers (e.g., glass) in composites. The interest in natural fiber-reinforced polymer composites is growing rapidly due to its high performance in terms of mechanical properties, significant processing advantages, excellent chemical resistance, low cost and low density. These advantages place natural fiber composites among the high performance composites having economic and environmental advantages. In the field of technical utilization of plant fibers, flax fiber-reinforced composites represent one of the most important areas. On the other hand, lack of good interfacial adhesion and poor resistance to moisture absorption make the use of natural fiber-reinforced composites less attractive. In order to improve their interfacial properties, fibers were subjected to chemical treatments, namely, mercerization, silane treatment, benzoylation, and peroxide treatment. Selective removal of non-cellulosic compounds constitutes the main objective of the chemical treatments of flax fibers to improve the performance of fiber-reinforced composites. The objective of this study was to determine the effects of pre-treated flax fibers on the performance of the fiber-reinforced composites.

Short flax fibers were derived from Saskatchewan-grown flax straws, for use in fiberreinforced composites. Composites consisting of high-density polyethylene (HDPE) or linear low-density polyethylene (LLDPE) or HDPE/LLDPE mix, chemically treated fibers and additives were prepared by the extrusion process. Extrusion is expected to improve the interfacial adhesion significantly as opposed to simple mixing of the two components. The extruded strands were then pelletized and ground. The test samples were prepared by rotational molding. The fiber surface topology and the tensile fracture surfaces of the composites were characterized by scanning electron microscopy to determine whether the modified fiber-matrix interface had improved interfacial bonding. Mechanical and physical properties of the composites were evaluated. The differential scanning calorimetry technique was also used to measure the melting point of flax fiber and composite.

Overall, the scanning electron microscopy photographs of fiber surface characteristics and fracture surfaces of composites clearly indicated the extent of fiber-matrix interface adhesion. Chemically treated fiber-reinforced composites showed better fiber-matrix interaction as observed from the good dispersion of fibers in the matrix system. Compared to untreated fiber-reinforced composites, all the treated fiber-reinforced composites had the same tendency to slightly increase the tensile strength at yield of composites. Silane, benzoylation, and peroxide treated fiber-reinforced composites offered superior physical and mechanical properties. Strong intermolecular fiber-matrix bonding decreased the high rate of water absorption in biocomposites. The incorporation of 10% untreated or chemically treated flax fibers also increased the melting point of composites. Further investigation is required to address the effect of increase in fiber content on the performance of composites.

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DEDICATION

I dedicate this thesis to my loving family who encouraged me to excel in a scholarly career. I wish to express my gratitude to my parents for the love and support during my studies.

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1. INTRODUCTION

Agriculture is an important sector in Western Canadian economy. Traditionally, agricultural materials have been shipped away for processing, or disposed of postharvest. Diversification of the industry is crucial in encouraging economic stability and growth. Value-added processing within Western Canada helps in agricultural diversification. Flax is an oilseed crop grown mostly in Saskatchewan and Manitoba. While the seed is processed for its high oil content, the biomass left behind tends to be a problem as it has strong fibers which breakdown very slowly under natural conditions. Traditionally, flax straw has been burned by farmers. Thus, the goal of this research has been to find a more environmentally responsible use for flax straw.

Traditional plastic materials are reinforced by glass fibers, which are both expensive and harmful to the environment. A flax-based biocomposite material contains polymers reinforced with flax fiber. There are a number of advantages of using flax fibers in biocomposites, among which are: a) flax fiber will make the material partially biodegradable; b) glass fiber is relatively expensive to make; c) flax is currently disposed of by burning; and d) flax has high tensile strength. Over the past decade, cellulosic fillers have been of greater interest as they give composites improved mechanical properties compared to those containing non-fibrous fillers. In recent years, thermoplastic materials have been increasingly used for various applications (Folkes

1982). Natural fiber-reinforced thermoplastic composites form a new class of materials which seem to have good potential in the future as a substitute for wood-based material in many applications. However, lack of good interfacial adhesion and poor resistance to moisture absorption makes the use of natural fiber-reinforced composites less attractive. Various fiber surface treatments like mercerization, isocyanate treatment, acrylation, latex coating, permanagante treatment, acetylation, silane treatment and peroxide treatment have been carried out which may result in improving composite properties. Research on a cost effective modification of natural fibers is necessary since the main attraction for today's market of biocomposites is the competitive cost of natural fiber. Interfaces play an important role in the physical and mechanical properties of composites. Reinforcing fibers are normally given surface treatments to improve their compatibility with the polymer matrix. This thesis attempts to address the following question: do chemical treatments of flax fibers have any influence on the composite properties?

The creation of fiber-reinforced composites is a multi-step process. First the flax fiber is collected as straw and then, chemical and/or physical treatments are used to reduce it to its fibrous form. Next it is chopped to appropriate size and combined with synthetic polymer materials. A series of steps including extrusion and plastic molding techniques are used to develop the final product.

Flax fiber-reinforced composite has material properties similar to that of conventional plastic products. The environmental benefits of the fiber-reinforced composite are

appealing to producers, consumers, and industry alike. The goal of this study is to determine the effects of pre-treated flax fibers on the performance of fiber-reinforced LLDPE, HDPE and HDPE/LLDPE composites. The vision is to develop an industry centered on the processing of flax fiber.

The objectives of this thesis follow the introduction. The chapter three contains a review of published literature relating to this subject. The literature review was used to guide this thesis in terms of its design, analysis, and expected trends. The methodology for the experimental work conducted is detailed in chapter four. The results and discussion on the effects of pre-treated flax fiber on the performance of composites follows. This thesis ends with a discussion on the results including conclusions and recommendations for future work. Data of tests appear in the Appendices.

2. OBJECTIVES

Most of the research reviewed indicated that only a limited work had been done on silane, benzoylation and peroxide-treatment of natural fibers for use in composites. The overall goal of the project was to focus on these three chemical treatments of flax fibers for use in fiber-reinforced LLDPE, HDPE and HDPE/LLDPE biocomposites to achieve improved properties of composites. The specific objectives of this study were to:

- investigate suitable fiber pre-treatment methods such as silane, benzoylation and peroxide treatment on their effects of flax fiber property;
- 2. explore a new method of manufacturing fiber-reinforced composites with the use of extrusion and rotational molding; and
- 3. study the effects of fiber surface modifications on the performance of flax fiberreinforced composites.

3. LITERATURE REVIEW

This chapter presents a review of literature on pre-treatment of flax fibers for use in composites. In addition, the details of different chemical treatments of natural fibers, characteristics of thermoplastic, composite manufacturing processes and the effects of fiber modification on the composites is discussed.

3.1 Properties of Natural Fibers

To better understand the properties of natural fiber-reinforced composite materials, it is necessary to know the physical and chemical properties of natural fibers.

3.1.1 Physical properties of natural fibers

Depending on their origin, natural fibers can be grouped into seed, bast, leaf and fruit qualities. The bast and leaf (the hard fibers) types are the most commonly used in composite applications (Williams and Wool 2000). Examples of bast fibers include hemp, jute, flax, ramie and kenaf. Leaf fibers include sisal and banana.

Flax fiber is an important bast fiber from the dicotyledonous *Linum usitatissimum* plant native to the Middle East. Its color is pale cream to brown. It has been used for centuries in the manufacture of fine linens (Edwards et al. 1997). Flax is a plant with a single stem, nearly one meter in height. The diameter at the base varies between 1 to 2 mm.

The length of a fibril is around 15-20 mm. Vegetable fibers have density of about half that of glass fibers. These fibers can withstand processing temperatures up to 250°C (Sreekala et al. 2000). They are fully combustible without the production of either noxious gases or solid residues.

The strength characteristics of fiber depend on the properties of the individual constituents, the fibrillar structure and the lamellae matrix (Joseph et al. 2000). For an understanding of the mechanical properties and durability of fibers, the structural components of the fibers (cellulose, hemicellulose and lignin) will be examined. Additional characteristics include fiber strength, fiber fitness, the polymerization of the cellulose, cleanness or purity and homogeneity of the sample, but these are not essentially genotype dependent.

Jähn et al. (2002) reported that plant fiber properties directly influence the physical parameters of the fiber-reinforced composites. Flax fiber properties are controlled by the molecular fine structure of the fibers which is affected by growing conditions and the fiber processing technique used. Flax fibers possess moderately high specific strength and stiffness and can be used as reinforcement in polymeric resin matrix to make useful composite materials.

Natural fibers exhibit considerable variation in diameter along with the length of individual filaments. Quality and other properties of fibers depend on factors such as size, maturity and processing methods adopted for the extraction of fiber (Mohanty et al.

2001). Properties such as density, electrical resistivity, ultimate tensile strength and initial modulus are related to the internal structure and chemical composition of fibers (Mohanty et al. 2001). Desirable properties for fibers include excellent tensile strength and modulus, high durability, low bulk density, good moldability and recyclability. Natural fibers have an advantage over glass fibers in that they are less expensive, abundantly available from renewable resources and have a high specific strength. Table 3.1 shows a comparison of properties of natural fibers and conventional man-made fibers.

Fiber	er Density Diameter (g/cm³) (μm)		Tensile Strength (MPa)	Young's Modulus (GPa)	Elongation at Break (%)	
Jute	1.3-1.45	20-200	393-773	13-26.5	7-8	
Flax	1.5	-	345-1100	27.6	2.7-3.2	
Hemp	-	-	690	-	1.6	
Ramie	1.5	-	400-938	61.4-128	1.2-3.8	
Sisal	1.45	50-200	468-640	9.4-22	3-7	
PALF	-	20-80	413-1627	34.5-82.51	1.6	
Cotton	1.5-1.6	-	287-800	5.5-12.6	7-8	
Coir	1.15	100-450	131-175	4-6	15-40	
E-glass	2.5	-	2000-3500	70	2.5	
S-glass	2.5	-	4570	86	2.8	
Aramid	1.4	-	3000-3150	63-67	3.3-3.7	
Carbon	1.7	-	4000	230-240	1.4-1.8	

Table 3.1. Comparative properties of natural fibers and conventional man-made fibers (Mohanty et al. 2000a).

3.1.2 Chemical composition of natural fibers

Natural fibers are complex in structure. They are generally lignocellulosic, consisting of helically wound cellulose micro fibrils in an amorphous matrix of lignin and hemi-cellulose. Table 3.2 shows natural fibers and their chemical and structural composition.

Mechanical properties are determined by the cellulose content and microfibril angle. A high cellulose content and low microfibril angle are desirable properties of a fiber to be used as reinforcement in polymer composites (Williams and Wool 2000).

Fiber	Cellulose (%)	Hemi- cellulose (%)	Lignin (%)	Extra- ctives (%)	Ash (%)	Pectin (%)	Wax (%)	Microfibril /spiral angle (°)	Moisture content (% w. b.)
BAST									
Jute	61-71.5	13.6-20.4	12-13	-	-	0.2	0.5	8.0	12.6
Flax	71-78.5	18.6-20.6	2.2	2.3	1.5	2.2	1.7	10.0	10.0
Hemp	70.2-74.4	17.9-22.4	3.7-5.7	3.6	2.6	0.9	0.8	6.2	10.8
Ramie	68.6-76.2	13.1-16.7	0.6-0.7	-	-	1.9	0.3	7.5	8.0
Kenaf	31-39	15-19	21.5	3.2	4.7	-	-	-	-
LEAF									
Sisal	67-78	10-14.2	8-11	-	-	10.0	2.0	20.0	11.0
PALF	70-82	-	5-12	-	-	-	-	14.0	11.8
Henequen	77.6	4-8	13.1	-	-	-	-	-	-
SEED									
Cotton	82.7	5.7	-	-	-	-	0.6	-	-
FRUIT									
Coir	36-43	0.15-0.25	41-45	-	-	3-4	-	41-45	8.0
WOOD									
Soft	40-44	25-29	25-31	5	0.2	-	-	-	-
Hard	43-47	25-35	16-24	2-8	0.4	-	-	-	-

Table 3.2. Chemical composition and structural parameters of natural fibers (Mohanty et al. 2000a).

The cells of flax fiber consist mostly of pure cellulose, being cemented as fascicle bundles by means of non-cellulosic incrusting such as lignin, hemicellulose, pectin, protein or mineral substances, resins, tannins, dyers and a small amount of waxes and fat (Mustată 1997). A mature flax cell wall consists of about 70% to 75% cellulose, 15% hemicellulose and pectic materials. Cellulose is a natural polymer with high strength and stiffness per weight, and it is the building material of long fibrous cells.

Selective removal of non-cellulosic compounds constitutes the main objective of fiber

chemical treatment. Both the hemicellulosic and pectic materials play important roles in fiber bundle integration, fiber bundle strength and individual fiber strength as well as water absorbency, swelling, elasticity and wet strength. The production of individual fibers without the generation of kink bands will generate fibers with much higher intrinsic fiber strength which is very useful for composite application (Mooney et al. 2001).

3.1.3 Moisture absorption characteristics

Natural fibers are hygroscopic in nature and they absorb or release moisture depending on environmental conditions. A major limitation of using natural fibers in durable composite applications is their high moisture absorption and poor dimensional stability (swelling) (Panigrahi et al. 2002). Swelling of fibers can lead to micro-cracking of the composite and degradation of mechanical properties. This problem can be overcome by treating these fibers with suitable chemicals to decrease the hydroxyl group in the fibers.

Stamboulis et al. (2000) reported that the moisture absorption and swelling of the treated flax fiber composites is approximately 30% lower than that of composites based on untreated flax fibers.

Strong intermolecular fiber-matrix bonding decreases the rate of moisture absorption in biocomposite. To increase the interface adhesion between the fiber and matrix, the fiber surface must be cleaned and chemically modified and the surface roughness must be increased (Yuan et al. 2002).

3.2 Chemical Treatment of Natural Fibers

Natural fibers are amenable to modification as they bear hydroxyl groups from cellulose and lignin. The hydroxyl groups may be involved in the hydrogen bonding within the cellulose molecules thereby reducing the activity towards the matrix. Chemical modifications may activate these groups or can introduce new moieties that can effectively interlock with the matrix.

3.2.1 Theoretical perspectives

In principle, natural fiber-reinforced composites could offer specific properties comparable to those of conventional fiber composites; however, low interfacial properties between fiber and polymer matrix often reduce the potential of natural fibers as reinforcing agents (Mohanty et al. 2001). Interfaces play an important role in the physical and mechanical properties of composites (Joseph et al. 2000). In order to improve natural fiber-matrix adhesion, the matrix should be commonly modified to better match fiber surface properties. Conversely, simple chemical treatments can be applied to the fibers with the aim of changing surface tension and polarity through modification of fiber surface (Scandola et al. 2000).

Several classes of compounds are known to promote adhesion, by chemically coupling the adhesive to the material. Silane coupling agents are one of many ingredients in commercial sizing that are applied to fibers. The chemical composition of coupling agents allows them to react with the fiber surface and forms a bridge of chemical bonds between the fiber and matrix (Al-Moussawi et al. 1993). Generally, coupling agents are molecules possessing two functions. The first function is to react with OH groups of cellulose and the second is to react with functional groups of the matrix. The selection of a coupling agent that can combine both strength and toughness to a considerable degree is important for a composite material to facilitate the optimum stress transfer at the interface between fiber and matrix. Several processes have been developed to modify polymers and fiber surfaces including chemical treatments, photochemical treatments, plasma treatments and surface grafting.

3.2.2 Surface chemical modifications of natural fibers

Several studies have shown the influence of various type of chemical modification on the performance of natural fiber and fiber-reinforced composites. The different surface chemical modifications of natural fibers such as mercerization, isocyanate treatment, acrylation, latex coating, permanganate treatment, acetylation, silane treatment and peroxide treatment with various coupling agents and others, have achieved various levels of success in improving fiber strength, fiber fitness and fiber-matrix adhesion in natural fiber composites. Brief descriptions of some important fiber chemical modifications are summarized in the following sub-sections.

3.2.2.1 Mercerization (alkali treatment)

Alkali treatment of cellulosic fibers, also called mercerization, is the usual method to produce high quality fibers (Ray et al. 2001). Alkali treatment improves the fiber-matrix adhesion due to the removal of natural and artificial impurities (Mishra et al. 2001a). Moreover, alkali treatment leads to fibrillation which causes the breaking down of the

composite fiber bundle into smaller fibers. In other words, alkali treatment reduces fiber diameter and thereby increases the aspect ratio. Therefore, the development of a rough surface topography and enhancement in aspect ratio offer better fiber-matrix interface adhesion and an increase in mechanical properties (Joseph et al. 2000). Alkali treatment increases surface roughness resulting in better mechanical interlocking and the amount of cellulose exposed on the fiber surface. This increases the number of possible reaction sites and allows better fiber wetting.

The following reaction takes place as a result of alkali treatment:

Fiber-OH + NaOH — Fiber-O'Na⁺ + H_2O

Jähn et al. (2002) found that the cellulosic fine structure of the flax fibers was directly influenced by mercerization treatment. Moreover, alkali treatment influenced the chemical composition of the flax fibers, degree of polymerization and molecular orientation of the cellulose crystallites due to cementing substances like lignin and hemicellulose which were removed during the mercerization process. Consequently, mercerization or more general alkali treatment had a lasting effect on the mechanical behavior of flax fibers, especially on fiber strength and stiffness (Gassan and Bledzki 1999). Several other studies were conducted on alkali treatment (Mishra et al. 2002; Joseph et al. 2000; Sreekala et al. 2000). They reported that mercerization led to the increase in the amount of amorphous cellulose at the expense of crystalline cellulose and the removal of hydrogen bonding in the network structure.

3.2.2.2 Silane treatment

Coupling agents usually improve the degree of cross-linking in the interface region and offer a perfect bonding result. Silane coupling agents were found to be effective in modifying the natural fiber-matrix interface. Various silanes were effective in improving the interface properties of wood-polypropylene (Coutinho et al. 1997), mineral-filled elastomers (González et al. 1997), fiber-reinforced epoxies (Culler et al. 1986) and phenolics composites (Ghatge and Khisti 1989). Alkoxy silanes are able to form bonds with hydroxyl groups. Coupling agents such as toluene dissocyanate and triethoxyvinyl silane were tested in fiber treatment in order to improve the interface properties. Silanes undergo hydrolysis, condensation and bond formation stage. Silanols can form polysiloxane structures by reaction with hydroxyl group of the fibers. The reaction schemes are given in Figure 3.1 and 3.2.

In the presence of moisture, hydrolyzable alkoxy group leads to the formation of silanols.

$$CH_2=CH-Si-OC_2H_5 \xrightarrow{H_2O} CH_2=CH-Si-O-H$$

Figure 3.1 Hydrolysis of silane (Sreekala et al. 2000).

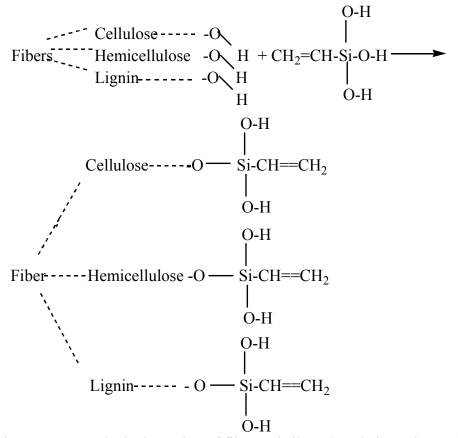


Figure 3.2 Hypothetical reaction of fiber and silane (Sreekala et al. 2000).

González et al. (1997) investigated the effect of silane coupling agent on the interface performance of henequen fiber-reinforced high-density polyethylene composites. The fiber-surface silanization resulted in better interfacial load transfer efficiency but did not improve the wetting of the fiber. Hydrogen and covalent bonding mechanisms could be found in the natural fiber-silane system. It was assumed that the hydrocarbon chains provided by the silane application influenced the wettability of the fibers, thus improving the chemical affinity to polyethylene.

Silane treatment of cellulosic fibers can increase the interfacial strength and therefore the mechanical properties of the composite (George et al. 1998; Bataille et al. 1989). Silane treatment also enhanced the tensile strength of the composite (Joseph et al. 2000).

3.2.2.3 Benzoylation

Manikandan Nair et al. (1996) reported that benzoylation of the fiber improved fiber matrix adhesion, thereby, considerably increasing the strength of composite. Joseph et al. (2000) studied the benzoylation treatment on the surface of fibers. A fixed amount of washed fiber (35g) was soaked in 18% NaOH solution for 0.5 h, filtered and washed with water. The treated fiber was suspended in 10% NaOH solution and agitated with 50 ml benzoyl chloride. The reaction between the cellulosic –OH group of sisal fiber and benzoyl chloride is shown in Figure 3.3 as follows:

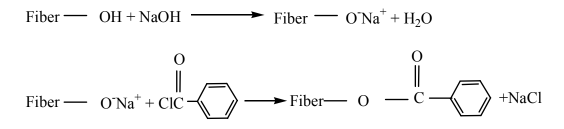


Figure 3.3 A possible reaction between cellulosic-OH groups and benzoyl chloride (Joseph et al. 2000).

3.2.2.4 Peroxide treatment

The decomposition of the peroxide and the subsequent reaction at the interface is expected at the time of curing of composites. Figure 3.4 shows the decomposition of the peroxides.



RO. + Cellulose – H \longrightarrow R – OH + Cellulose.

Figure 3.4 Peroxide treatment reaction (Sreekala et al. 2000).

Peroxide-induced adhesion in cellulose fiber-reinforced thermoplastic composites has attracted the attention of various researchers due to easy processability and improvement in mechanical properties. Sapieha et al. (1990) indicated that the addition of a small amount of benzoyl peroxide or dicumyl peroxide to cellulose-polymer (LLDPE) systems during processing improved the composite mechanical properties. The improvement of mechanical properties is attributed to the peroxide-induced grafting of polyethylene onto cellulose surfaces.

Joseph et al. (2000) investigated benzoyl peroxide treatment on short sisal fiberreinforced polyethylene composites. They reported that peroxide-treated composites showed an enhancement in tensile properties due to the peroxide-induced grafting. Sreekala et al. (2000) also studied benzoyl peroxide treatment on oil palm fiberreinforced phenol formaldehyde composites. Fibers were coated with benzoyl peroxide from acetone solution after alkali pre-treatments. High temperature was favored for decomposition of the peroxide. They reported that peroxide-treated fiber composites could withstand the tensile stress to higher strain level.

3.2.2.5 Other chemical treatment methods

Several interface modification methods were reported in literature. Acetylation of natural fibers is a well-known esterification method to introduce plasticization to cellulosic fibers. Acetylation has been extensively applied to wood cellulose to stabilize the cell wall, improving dimensional stability and environmental degradation. One of the modification techniques employed by the Okura Company in Japan was to produce esterified woods (Mohanty et al. 2001), which would be molded into plastic sheets by hot pressing.

Another effective method of surface chemical modification of natural fibers is graft copolymerization. Optimized vinyl grafted natural fibers, consisting of the orderly arrangement of grafted moieties, act as compatible reinforcing fibers with several resin systems to obtaining better fiber-matrix adhesion of the resulting biocomposites (Mohanty et al. 2001; Ghosh and Ganguly 1993).

Isocyanate has a functional group -N=C=O which is very susceptible to reaction with the hydroxyl group of cellulose and lignin in the fibers and forms strong covalent bonds, thereby creating better compatibility with the binder resin in the composites. Kokta et al. (1990a) and Raj et al. (1988) pointed to the performance of isocyanate as a coupling agent. Isocyanates provided better interaction with thermoplastics resulting in superior properties. Isocyanates could act as a promoter or as an inhibitor of interaction.

The radical enhances the chemical interlocking at the interface. Permanganate treatment was carried out to improve the bonding at the fiber-polymer interface. Joseph et al. (2000) and Sreekala et al. (2000) investigated the fibers which were pre-treated with alkali and then dipped in permanganate solution in acetone. Tensile strength values of the composite showed a marginal increase with permanganate treatment.

Mustată (1997) studied the sodium chlorite treatment on the surface of bleached flax.

The experiment focused on the links of fibers formed between lignin and carbohydrates. The stability of pluricellular fibers were subjected to mechanical stresses. Removal of a part of the flax fibers' noncellulosic compounds by chemical treatments was reflected in the mechanical and physical characteristics of the surface state, as well as in the fibers' behavior during processing and wearing.

Acrylation treatment, maleated polypropylene/maleic anhydride treatment and titanate treatment of cellulosic fibers have also been reported (Sreekala et al. 2000; Mohanty et al. 2001; Monte and Sugerman 1984). Acrylation treatment resulted in high strain values of the composites. The composites ability to withstand the applied flexural stress is manifested by higher strain values, which indicate the elastic nature of the material. Maleated polypropylene or maleic anhydride grafted polypropylene (MAPP) has been widely used as a coupling agent or a compatibilizer in natural fiber reinforced polypropylene composites. The treatment of natural fibers with MAPP copolymer provides covalent bonds across the interface. Through such treatment, the surface energy of the fibers is increased, thereby providing better wettability and high interfacial adhesion. Many other compounds such as chromium complexes and titanates can be used as coupling agents. The processing of composites with titanate coupling agents found that the deposition of a monolayer of organ functional titanate eliminated the water of hydration. This enhanced the dispersion and compatibility at the interface.

3.3 Development of Natural Fiber Composites

Since 1941, the study on composites, particularly natural fiber-reinforced plastics has

gained increasing attention of researchers and manufacturers (Joseph et al. 2000). The increased interest in natural fiber-reinforced composites is due to the high performance in mechanical properties, significant processing advantages, excellent chemical resistance, low cost and low density. They have long served many useful purposes but the application of material technology for the utilization of natural fibers as reinforcement in polymer matrix has taken place in recent years. Biocomposite consists of a polymer as the matrix material and a natural fiber as the reinforcing element. The use of fibers derived from annually renewable resources, such as reinforcing fibers, provide positive environmental benefits with respect to ultimate disposability and raw material utilization.

Recent studies indicate that plant-based natural fibers can be used as reinforcement in polymer composites, replacing the expensive and non-renewable synthetic fibers such as glass, because of their potential for recyclability (Mohanty et al. 2001). Vegetable fibers can serve as excellent reinforcing agent for plastics because of their moderately high specific strength and stiffness which is used as reinforcing materials in polymeric resin matrices to make useful structural composite materials (Joseph et al. 2000). Cellulose-based natural fibers are a potential resource for making low cost composite materials. Cellulosic fillers of a fibrous nature have been of greater interest, because they would give composites with improved mechanical properties compared to those containing non-fibrous fillers. Lignocellulosic fibers like jute, sisal, coir, and pineapple have been reportedly used as reinforcements in polymer matrix (Joseph et al. 2000).

Published data (Zaini et al. 1996; Woodhams et al. 1984) show that various commercial wood fibers have good potential as reinforcements in thermoplastics. Biocomposites can make more value-added products from bioplastics and will get more attention in the future.

3.4 Characteristics of Thermoplastic Polymers

Thermoplastic polymers constitute an important class of materials with a wide variety of applications (Kokta et al. 1983). They are capable of being re-melted without any change in chemical structure or properties. Polyethylene, polypropylene, polystyrene and polyvinyl chloride are the most common thermoplastic polymers and are frequently called commodity polymers (Crawford and Throne 2000). As long as processing does not mechanically damage the thermoplastic polymer structure, these polymers are considered recyclable. Besides the use of "pure" polymer for structural purposes, it is also used as a matrix for fiber-reinforced composites. These composites are mostly based on the traditional reinforcement fibers such as glass fibers. However, natural fibers can also be used as reinforcement. Polymers have a different affinity towards the fiber owing to the difference in their chemical structure. In the search for the most suitable thermoplastic matrix for a flax fiber-reinforced composite, density and temperature related properties seems to be limiting criteria (Mohanty et al. 2001).

Low-density polyethylene (LDPE), high-density polyethylene (HDPE) and linear lowdensity polyethylene (LLDPE) will be used as major matrix for reinforcement. These thermoplastic polymers are widely used and have a melting point compatible with natural fibers. Their low melting points also allow processing below the degradation temperature of the fibers. Table 3.3 lists typical properties of three major thermoplastics.

Polymer Type	Density (g/cm³)	Degree of Crystallinity	Glass Transition Temperature (°C)	Crystal Melting Temperature (°C)	Tensile Strength (MPa)	Elongation at Break (%)	Flexural Modulus (GPa)
High-density polyethylene (HDPE)	0.95-0.97	high	-120	137	20-30	10-1,000	1-1.5
Low-density polyethylene (LDPE)	0.92-0.93	moderate	-120	110	8-20	100-650	0.25-0.35
Linear low- density polyethylene (LLDPE)	0.91-0.94	high	-74	122-124	20	100-500	0.35

Table 3.3. Typical properties of three major thermoplastics (www.azom.com).

3.4.1 High-density polyethylene (HDPE)

High-density polyethylene (HDPE), also known as linear polyethylene or low-pressure polyethylene, is the preferred polyethylene for chemical containers of all sizes primarily due to its exceptional environment stress crack resistance (Crawford and Throne 2000). It has excellent stiffness from room temperature to the boiling point of water. Even though HDPE is frequently called linear polyethylene, it still has some short chain branching. Nevertheless, its linear nature and its high backbone mobility allow it to crystallize from 75% to 90% of theoretical. The crystalline structure causes the product to have a milky, translucent appearance. Since the crystallite is more ordered and more tightly packed than the amorphous phase, the density of HDPE is typically around 960 kg/m³ approaching the theoretical value of 1000 kg/m³. Many HDPEs are formulated for extrusion and blow molding applications (Crawford and Throne 2000).

3.4.2 Linear low-density polyethylene (LLDPE)

LLDPE has side chains similar to those of LDPE but with proper catalysts and coreactive agents, the chains are dramatically reduced in length (Crawford and Throne 2000). LLDPE has a density range of 910 kg/m³ to about 940 kg/m³, and is 65% to 75% crystalline at room temperature. Competitive with LDPE, the 'linear low' materials have found rapid acceptance because of their high toughness (at low, normal and high temperatures), improved stiffness, chemical resistance, tensile strength, elongation at break and puncture resistance. However, it has somewhat poorer impact strength when compared with LDPE and MDPE. Suppliers have emphasized more specifically the improved resistance to environmental stress cracking (Brydson 1989).

In Lee and Joo's study (1999), a thermoplastic LLDPE resin was used as the matrix for fiber composites. Its low processing temperature (less than 130°C) made composite fabrication possible without partial melting or annealing of the fibers. The high toughness of LLDPE yielded a good impact-resistant composite and had advantages of thermoplastic composite processing, such as short processing time, unlimited storage time and solvent free processing.

3.4.3 Low-density polyethylene (LDPE)

Low-density polyethylene (LDPE), also referred to as high-pressure polyethylene or branched polyethylene, has extensive side chains, up to about 100 ethylene units in length. The long branches tend to inhibit molecular organization during cooling. As a result, LDPEs typically have relatively low densities of 910 kg/m³ to 925 kg/m³ and relatively low crystallinity of 45% to 66%. LDPEs are relatively soft polyethylene with flexural modulus ranging between 0.24 and 0.35 GPa. Owing to the high number of tertiary hydrogen, LDPE does not have good environmental stress crack resistance (ESCR). Nevertheless, LDPEs mold well at low temperatures that accurately replicate mold surfaces (Crawford and Throne 2000).

3.5 Composites Manufacturing

A systematic study of the process variables for composites based on different natural fibers like flax, jute, sisal, ramie, and pineapple has been made with a view to determine the most suitable processing conditions for such composites by some researchers. Some major methods of molding are rotational molding, compression molding, injection molding and extrusion. Alternative processing methods of natural fiber-reinforced composites are an important advancement necessary for their increased use.

3.5.1 Rotational molding

Rotational molding is a process for manufacturing hollow plastic products. Rotational molding involves power mixing, melting, sintering and melt solidification. Various aspects of the rotational molding process have been studied by several researchers (Throne 1979; Crawford 1992). Fundamental research on rotational molding has been directed to reduce the molding cycle time and to optimize the mechanical properties of final parts. Polyethylene accounts for more than 80% of the total production (Bellehumeur and Vlachopoulos 1998). Rotational molding has particular advantages in terms of relatively low levels of residual stresses and inexpensive molds. Rotational

molding also has few competitors for the production of large (>2 m^3) hollow objects in one piece. Currently, the rotational molding industry is in its exciting development. Important new market sectors are opening up as rotational molders are able to deliver high quality parts at competitive prices.

3.5.2 Extrusion

The extrusion process basically consists of continuously shaping a fluid polymer through the orifice of a suitable tool (die), and subsequently solidifying it into a product (Hensen 1997). In the case of thermoplastics, the feed material, in powder or pellet form, is now most commonly heated to a fluid state and pumped into the die. Oladipo et al. (1999) investigated the composite (aspen wood fiber/HDPE) manufacturing process. The components were fed at pro-determined mass flow rates, based on the desired wood fiber mass fraction in the composite, into a ZSK-30 Werner & Pfleiderer extruder (Werner & Pfleiderer Ltd., Marple, Cheshire, UK) having 28 mm co-rotating twin screws. The extruder was operated at a working temperature of 150°C and a screw speed of 100 rpm. This temperature ensured that while the polymer was fully melted (melting point is 120-135°C), the wood fibers were not burned.

3.6 Effects of Fiber Surface Modifications on Fibers

The chemical modification directly influences the cellulosic fine structure of natural fiber. This section reviews the effects of fiber modification on the stress-strain behaviour and tensile properties of flax fibers.

3.6.1 Stress-strain behavior

The mechanical performance of fibers is dependent upon its chemical composition, chemical structure and cellular arrangement. Sreekala et al. (2000) performed tensile stress-strain test for untreated and modified oil palm fiber. Each individual fiber was composed of fibrils held together by non-cellulosic substances, such as lignin and pectin. Failure of the fiber was gradual upon the application of tensile stress. It showed intermediate behavior between brittle and amorphous. As stress gradually increased, some of the fibrils may have slipped out. The total of the stress was then sheared by fewer cells. Further increase of stress led to the rupture of cell walls and decohesion of cells. This resulted in a catastrophic failure of the fiber. Modifications led to major changes on the fibrillar structure of the fiber. It removed the amorphous components. This changed the deformation behavior of the fibers. The brittleness of the fiber was substantially reduced upon treatments.

3.6.2 Tensile properties of flax fibers

Sreekala et al. (2000) measured the tensile properties of untreated and modified fibers, such as tensile strength, Young's modulus and elongation at break. Many of the modifications decreased the strength properties due to the breakage of the bond structure, and also due to the disintegration of the non-cellulosic materials. Some of the treatments, like silane and acrylation, led to strong covalent bond formation and thereby the strength was enhanced marginally. Optimum mechanical performance was observed for silane-treated and acrylated fiber. The reinforcing ability of the fibers did not just depend upon the mechanical strength of the fibers but on many other features, such as

polarity of the fiber, surface characteristics and presence of reactive centers. These factors control interfacial interaction. The Young's modulus of the fibers improved upon acrylation, alkali and silane treatment. The improved stiffness of the fibers was attributed to the crystalline region (cellulosic) of the fiber. The fiber also showed very good elongation properties, with values increasing upon modifications. Lower elongation of the untreated fiber may be due to the three dimensionally cross-linked networks of cellulose and lignin. Treatment broke this network structure giving the fiber higher elongation and lower strength properties.

Mishra et al. (2001b) investigated the tensile properties of untreated, chemically modified and AN-grafted sisal fibers. Chemically modified fibers showed an appreciable decrease in the tensile properties. This decrease was attributed to the substantial delignification and degradation of cellulosic chains during chemical treatment. The extension at break of these fibers did not change much. In all the cases of grafting, it has been found that the tensile strengths were higher than that of untreated fiber.

3.7 Effects of Fiber Surface Modifications on Composite Properties

Chemical treatments will be necessary to strengthen the interface between fiber and matrix. Several studies have been conducted on the influence of various types of chemical modifications on the properties of natural fiber-reinforced thermoplastic composites (Mansour et al. 1983; Manrich and Agnelli 1989; Kenaga et al. 1962). This section reviews the effects of fiber modification on the mechanical properties, thermal properties and macro-mechanical properties of composites.

3.7.1 Mechanical properties of composites

The mechanical properties of a natural fiber-reinforced composite depend on many parameters, such as fiber strength, modulus, fiber length and orientation, in addition to the fiber-matrix interfacial bond strength. A strong fiber-matrix interface bond is critical for high mechanical properties of composites. A good interfacial bond is required for effective stress transfer from the matrix to the fiber whereby maximum utilization of the fiber strength in the composite is achieved (Karnani et al. 1997). Modification to the fiber also improves resistance to moisture-induced degradation of the interface and the composite properties (Joseph et al. 2000). In addition, factors like processing conditions/techniques have significant influence on the mechanical properties of fiber-reinforced composites (George et al. 2001).

Sapieha et al. (1989; 1990) have found that by the addition of a small amount of dicumyl peroxide or benzoyl peroxide into the cellulosic fiber-polymer (LDPE) systems during processing significantly improved the mechanical properties of the composite. Kokta et al. (1990a; 1990b) have extensively studied the effect of different chemical modifications, such as silane treatment and grafting, on the mechanical properties and dimensional stability of cellulosic fiber-thermoplastic composites. They found that the chemically modified cellulosic fiber-reinforced thermoplastic composites offered superior physical and mechanical properties under extreme conditions even after recycling. Ray et al. (2001) have employed the technique on jute and found that the improvements occurred on the fiber properties. Münker and Holtmann (1998) studied different natural fibers (flax, ramie, curaua) and matrices (polyester, polypropylene).

Their findings showed that mechanical properties of natural fiber-reinforced composites could be improved by the use of different coupling agents.

3.7.1.1 Tensile properties of composites

Natural fiber-reinforced composites often show enhancement in tensile properties upon different modifications owing to the increased fiber-matrix adhesion. Tensile properties can be explained on the basis of the changes in chemical interactions at the fiber-matrix interface. The tensile strength of flax fiber-reinforced composites is determined both by the tensile strength of the fibers and the presence of weak lateral fiber bonds.

Sreekala et al. (2000) performed one of the pioneering studies on the mechanical performance of treated oil palm fiber-reinforced composites. They studied the tensile stress-stain behavior of composites having 40% by weight fiber loading. Isocyanante-, silane-, acrylated, latex coated and peroxide-treated composite withstood tensile stress to higher strain level. Isocyanate treated, silane treated, acrylated, acetylated and latex coated composites showed yielding and high extensibility. Tensile modulus of the composites at 2% elongation showed slight enhancement upon mercerization and permanganate treatment. The elongation at break of the composites with chemically modified fiber was attributed to the changes in the chemical structure and bondability of the fiber. Alkali treated (5%) sisal-polyester biocomposite showed about 22% increase in tensile strength (Mishra et al. 2002). Ichazo et al. (2001) found that adding silane-treated wood flour to PP produced a sustained increase in the tensile modulus and tensile strength of the composite. Joseph and Thomas (1993) studied the effect of chemical

treatment on the tensile and dynamic mechanical properties of short sisal fiberreinforced low density polyethylene composites. It was observed that the CTDIC (cardanol derivative of toluene diisocyanate) treatment reduced the hydrophilic nature of the sisal fiber and enhanced the tensile properties of the sisal-LDPE composites. They found that peroxide and permanganate treated fiber-reinforced composites showed an enhancement in tensile properties. They concluded that with a suitable fiber surface treatment, the mechanical properties and dimensional stability of sisal-LDPE composites could be improved. Mohanty et al. (2000b) studied the influence of different surface modifications of jute on the performance of the biocomposites. More than a 40% improvement in the tensile strength occurred as a result of reinforcement with alkali treated jute. Jute fiber content also affected the biocomposite performance and about 30% by weight of jute showed optimum properties of the biocomposites.

3.7.1.2 Impact properties of composites

Fibers have a significant effect on the impact resistance through the principle of stress transfer. When an impact load is applied perpendicular to the reinforcing fibers, good fiber-matrix adhesion is required for even moderate impact strength (Nielsen 1974). The impact properties of the polymeric materials are directly related to the overall toughness of the material (Shah 1998). Toughness is defined as the ability of the polymer to absorb applied energy. Impact resistance is the ability of a material to resist breaking under a shock loading or the ability to resist fracture under stress applied at high speed.

Detailed studies have already been done on the impact resistance of short fiber-

reinforced composites (Kau 1990; Jang et al. 1990; Reed and Bevan 1993). The impact resistance of fiber-reinforced composite depends on fiber rigidity, interfacial stress resistance and fiber aspect ratio. The strength of the matrix, the weakest part of the material, should be related to the failure process. The involvement of fibers in the failure process is related to their interaction with the crack formation in the matrix and their stress transferring capability. The total energy dissipated in the composite before final failure occurs is a measure of its impact resistance. The total energy absorbed by the composite is the sum of the energy consumed during plastic deformation and the energy needed for creating new surfaces.

3.7.2 Macro-mechanical properties of composites

The macro-mechanical properties of composites are attributed to an increase in the interfacial shear strength (ISS) of the modified composites. The ability to control the chemical and mechanical properties of the fiber-matrix interphase is crucial. Interface studies of untreated and surface treated sisal-polyester composites has been investigated to determine fiber splitting, fiber pullout, debonding, matrix cracking and fiber-matrix interaction using scanning electron microscopy (Mishra et al. 2002). Scanning electron micrographs of the tensile fracture of the composites revealed the failure mechanisms and impact fracture morphology. Fiber breakage was the main failure criteria observed. They also reported that in the untreated sisal composite, a clean pullout of fibers without any adhering resin matrix was observed. This proved that there was very poor adhesion between fiber and matrix. The 5% alkali-treated sisal composite showed better fiber-matrix interaction as observed from the good dispersion of fibers in the matrix system

predicting micropores at the interface. Morphological studies showed that the MAPP and silane treatment improved the polymer-wood flour (WF) adhesion and the dispersion of the particles while the alkaline treatment only improved the dispersion (Ichazo et al. 2001).

3.7.3 Thermal properties of composites

A quick method for determining the threshold values for processing temperature is done by thermal analysis. Thermogravimetric (TG) analyses are carried out with a thermal balance. The thermo gravimetric degradation curve provides information about the thermal stability of a material (Shah 1998).

Wielage et al. (1999) used differential scanning calorimetry (DSC) to determine the melting point of the flax-reinforced polypropylene and to collect caloric data. DSC is a thermo-analytical technique in which the heat flow is measured as a function of temperature or time. They subjected the flax-reinforced polypropylene to a defined temperature regime under controlled atmosphere and reported that the melting range of the polymer matrix was displayed as an endothermic peak. An increased heating rate leads to a displacement of the melting range to higher temperatures. Powell et al. (2002) considered the effect of the matrix on the heating of flax fiber-reinforced composites. They reported that pure HDPE showed degradation beginning at approximately 410-430°C. Therefore, the matrix absorbed a great amount of heat, taking the thermal stress off of the flax fiber.

3.8 Summary

Most research reviewed indicated the effect of alkali treatment, isocyanate treatment, acrylation, latex coating, permanganate treatment, acetylation on the fiber-reinforced composite. Only few studies in literature were related to silane treatement, benzoylation and peroxide treatment in improving fiber strength, fiber-matrix adhesion and the performance of the natural fiber composites. The results of these studies were limited. This research was conducted on three chemical modifications. There is a lack of measurement of fiber bundle tensile strength. The method used in this research involved measuring the thermal properties of the composite using the differential scanning calorimetry. A multi-step process of production of fiber-reinforced composites by rotational molding was reviewed. In this research, the effect of chemical modification on the performance of the fiber-reinforced composites was studied.

4. MATERIALS AND METHODS

In this chapter, the materials and devices, experimental procedures and data acquisition used to complete the experiments are presented. Data used throughout the development of this thesis to obtain the tables and figures presented can be found in its original form in the Appendices.

4.1 Materials

Flax fibers were derived from linseed flax grown in Saskatchewan and decorticated on a standard scutching mill at Durafiber in Canora, SK. The fibers were first washed thoroughly with 2% detergent water and dried in an air oven at 70°C for 24 h. The dried fibers were designated as untreated fibers. Flax fibers were then subjected to sequential extraction with 1:2 mixture of ethanol and benzene for 72 h at 50°C, followed by washing with double distilled water and air drying to remove waxes and water soluble ingredients prior to chemical treatments. Reagent grade chemicals were used for fiber surface modifications, namely, sodium hydroxide (NaOH), benzoyl chloride, ethanol, dicumyl peroxide, acetone and alcohol. The structure of coupling agent, triethoxyvinylsilane (Aldrich Chemical Co. Ltd.) is shown in Figure 4.1.

In this series of experiments, high-density polyethylene, linear low-density polyethylene (HDPE 8761.27 and LLDPE 8460.29, Exxon Mobil, Toronto, ON) and LLDPE/HDPE

25087 (NOVA Chemicals Ltd., Calgary, AB) were used as polymer matrix materials.

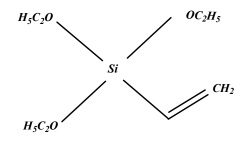


Figure 4.1 Structure of triethoxyvinylsilane (Aldrich Chemical Co. Ltd.).

4.2 Fiber Surface Treatment

Generally, the first step in chemical treatment is the mercerization process (pretreatment process) of all the fiber samples which cause changes in the crystal structure of cellulose. Fibers were soaked in 5-18% NaOH (silane treatment: 5%, benzoylation: 18%, peroxide treatment: 10%) for about half an hour in order to activate the OH groups of the cellulose and lignin in the fiber. The appropriate concentration of NaOH solution used in mercerization before each type of chemical treatment was completed in the initial work. Sreekala et al. (2000) indicated that a 10-30% sodium hydroxide solution produced the best effects on natural fiber properties. Flax fibers were soaked into 2.5, 5, 10, 13, 15, 18, 20, 25, or 30% NaOH solutions before the chemical treatment. It was found that 5%, 18% or 10% of sodium hydroxide solution were the appropriate concentrations for mercerization before silane, benzoylation or peroxide treatment, respectively. The fibers were then washed many times in distilled water and finally dried. The different chemicals can then be used on the surface in order to improve their interfacial properties. **Silane treatment:** The pre-treated fibers were dipped in an alcohol water mixture (60:40) containing triethoxyvinylsilane coupling agent. The pH of the solution was maintained between 3.5 and 4, using the METREPAK Phydrion buffers and pH indicator strips. Fibers were washed in double distilled water and dried in the oven at 80°C for 24 h.

Benzoylation: The pre-treated fibers were suspended in 10% NaOH solution and agitated with benzoyl chloride. The mixture was kept for 15 min, filtered, washed thoroughly with water and dried between filter papers. The isolated fibers were then soaked in ethanol for 1 h to remove the benzoyl chloride and finally was washed with water and dried in the oven at 80°C for 24 h.

Peroxide treatment: Fibers were coated with dicumyl peroxide from acetone solution after alkali pre-treatments. Saturated solution of the peroxide in acetone was used. Soaking of the fibers in the solution was conducted at a temperature of 70°C for 30 min. High temperatures were favored for decomposition with the peroxide. The chemically treated fibers were washed with distilled water and placed in an oven at 80°C for 24 h.

4.3 Composite Preparation

The silane coupling agent was processed in the lab. The liquid form of triethoxyvinylsilane was blended with the thermoplastic powder and fed to the laboratory mixing extruder (LME) (Dynisco, Franklin, MA). The extruded strands were pelletized and ground into powder form. Then the silane coupling agent was added

during the dry compounding of fiber/thermoplastic at a rate of 5% by mass as "resin additive".

Pre-treated and untreated fibers were ground by the grinding mill (Falling Number, Huddinge, Sweden) and oven dried at 80°C for 24 h to reduce the moisture content to less than 2%. Mixtures of thermoplastic powder and 10% by weight of flax fibers were prepared by using a food blender (Waring Products Corporation, New York, NY). This was done to aid in the homogeneous mixing of fibers and polymer matrix during the extrusion process. The blend was fed into the twin-screw extruder (Werner & Pfleiderer Engineers, Ramsey, NJ) located at the Centre for Agri-Industrial Technology (CAIT) in Edmonton, AB using a barrel to die temperature profile of 175°C, a screw speed of 125 rpm and feed rate to the extruder of 20 kg/hr. Blends prepared in this manner were extruded using a six-hole strand die (Figure 4.2). Figure 4.3 shows the material being extruded. Extruded strands were then pelletized. The pellets were ground using a grinding mill (Retsch GmbH 5657 HAAN, West Germany) and the ground product was used in rotational molding.

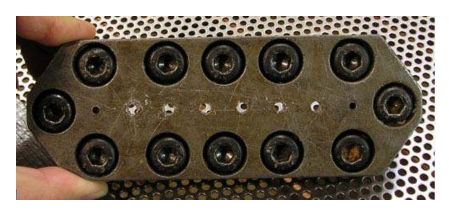


Figure 4.2 Six-hole strand die.



Figure 4.3 The material being extruded.

In the initial stage of this research, two types of extruders were compared. The singlescrew extruder (CW Brabender Instruments Inc., South Hackensack, NJ) at the Northern Alberta Institute of Technology (NAIT) in Edmonton, AB was used in the extrusion process to compare with the output from a twin-strew extruder, as previously detailed. Swelling of extruded strands was observed when the single-strew extruder was used. Therefore, the twin-screw extruder offers a superior mixing and compounding compared to the single-screw extruder.

4.4 Biocomposites Manufacturing by Rotational Molding

The powder of fiber/thermoplastic was dried in an air-circulating oven for 24 h at 70°C before rotational molding. Test samples were prepared from ground extruded strands using a rotational molding machine located at Norwesco Canada Ltd. in Saskatoon, SK. It is a carousel-type molding machine with four separate arms that can each rotate at two separate axes, while completely enclosed in an oven at 250°C for 30 min. Single large

mold cavities can be placed on each arm or many smaller shapes. Rotational molding manufacturing process is shown in Figure 4.4.

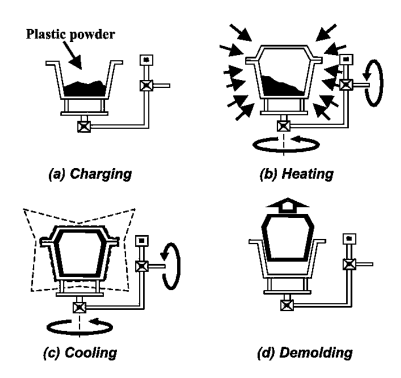


Figure 4.4 Rotational molding manufacturing process (Beall 1998).

4.5 Experimental Design and Data Analysis

The study was broken into two phases. The first phase focused on the effect of chemical modifications on the flax fibers, while the second phase focused on the effect of pre-treated flax fibers on the performance of the fiber-reinforced composite.

4.5.1 Experimental design

The experimental design is a factorial arrangement of treatments conducted in a randomized design. Table 4.1 shows the outline of the experimental design for three types of treated fibers.

Main treatment: T₁ (Untreated); T₂ (Silane treatment);

 T_3 (Benzoylation); T_4 (Peroxide treatment) = 4

Sub-treatment: S₁ (LLDPE); S₂ (HDPE); S₃ (HDPE/LLDPE) = 3

Treatment combination = 4 types of fiber \times 3 types of thermoplastic = 12

 $(T_1S_1, T_1S_2, T_1S_3, T_2S_1, T_2S_2, T_2S_3, T_3S_1, T_3S_2, T_3S_3, T_4S_1, T_4S_2, T_4S_3)$

Total tests conducted = 12×5 (replicates) = 60 tests

Fiber	Polymer	Fiber Content	Silane Coupling Agent (%)
I luture te d	LLDPE	10	5
Untreated (2% detergent water)	HDPE	10	5
(276 delergent water)	HDPE/LLDPE	10	5
Silane treated (triethoxyvinylsilane coupling agent)	LLDPE	10	5
	HDPE	10	5
	HDPE/LLDPE	10	5
Democrate tion treated	LLDPE	10	5
Benzoylation treated (benzoyl chloride)	HDPE	10 10 10 10 10	5
(belizbyr chioride)	HDPE/LLDPE	10	5
Demoniale transfer l	LLDPE	10	5
Peroxide treated	HDPE	10	5
(dicumyl peroxide)	HDPE/LLDPE	10	5

Table 4.1. Experimental design for three types of treated fibers.

Data for each test appear in Appendices A-G. In each test, the experimental parameters and conditions are listed. The number of replicates in each test is listed in Table 4.2. According to the appropriate ASTM standard, five samples were replicated for tensile strength at yield test of composites.

This research studies the effect of fiber surface modifications on the performance of composites focused on four properties, namely, morphological characterization, mechanical, physical and thermal property. Table 4.2 shows the property tests performed

on each composite sample and the number of replicates.

Property	Test	Replicates
Morphological	Fiber surface topology (SEM)	1
Characterization	Composite microstructure (SEM)	1
	Fiber bundle tensile strength test (Instron)	50
Mechanical	Tensile strength at yield of composites (Instron) ASTM: D638	5
	Tensile-impact strength of composites (ASTM: D1822-93)	5
	Durometer hardness of composites (ASTM: D2240-97)	10
Physical	Moisture absorption of flax fibers (Environmental test chamber)	3
5	Water absorption of composites (ASTM: D570-99)	3
Thermal	Melting point of composites (DSC)	1

Table 4.2. Property tests conducted on sample composites.

4.5.2 Data analysis

The number of replicate specimens varied for each test type. The data constitute a sample of certain observations from the population of all loads obtainable by these techniques. Deviation in the observed properties arises from variability in the manufacturing of the composites, in the machine administering the load, in the experimenter's technique, and in a host of other, possibly unknown factors. It is clear that we cannot hope to obtain a mean (μ) and standard deviation (σ) for this abstract population, but we can estimate them. A one-way analysis of variance (ANOVA) was used to study the difference between the tests and between treatments and the results are shown in Appendices A-G. The Duncan's multiple range test was also used to compare the paired means.

In each of the test, the sample mean value (\overline{Y}), standard deviation (s), the standard error

of the mean (s_Y) , the coefficient of variation (CV) and 95% confidence interval of the mean were calculated. The description of the statistical analysis of each test is also listed in the Appendices. All the data analysis was generated using the program in SPSS statistical software (SPSS Inc., Chicago, IL).

4.6 **Experimental Procedures**

Morphological characterization test of fiber and the composites were carried by a scanning electron microscope (SEM). Mechanical property test focused on tensile strength of fiber bundle, tensile strength of composites, tensile-strength test of composites and Durometer hardness test of composites. Physical property test focused on moisture absorption test of fibers and water absorption test of composites. Thermal property measurement was conducted by a differential scanning calorimetry (DSC).

4.6.1 Morphological characterization

The treated fiber and the composites were examined by a scanning electron microscopy (SEM505 Philips Electron Optics, Eindhoven, the Netherlands). The sample surfaces were vacuum coated with a thin layer of gold on the surface using an Edwards S150B sputter coater (BOC Edwards, Wilmington, MA) to provide electrical conductivity and did not significantly affect the resolution. One set of sample from each type of fibers/composite was examined. Figure 4.5 shows the SEM used in this study located in the Department of Biology at the University of Saskatchewan.

As a supplementary tool, the microscopic examination of treated and untreated fiber

surface was carried out with a scanning electron microscope at the accelerating voltage of 10 KV. The microstructure of the fiber-matrix interface of composites was examined at the accelerating voltage of 30 KV.



Figure 4.5 Scanning electron microscope.

4.6.2 Mechanical properties

Specimens were conditioned for 7 days at standard laboratory atmosphere prior to performing mechanical tests. The appropriate ASTM methods were followed. At least five replicate specimens were tested and the results were presented as an average of tested specimens. The tests were conducted at a standard laboratory atmosphere of 23°C and 50% relative humidity.

4.6.2.1 Tensile strength of fiber bundle

Tensile test is a measurement of the ability of material to withstand forces that tend to pull it apart. It determines to what extent the material stretches before breaking. Flax fiber bundle tensile strength tests were preformed by a computer-controlled Instron Model 1011 (Instron Corporation, Canton, MA) with a gauge length of 40 mm and at a crosshead speed of 5 mm/min. Figure 4.6 shows how the apparatus appeared once it has been clamped and ready for testing. The round bars were covered with surgical glove fingers, and the flax was clamped at the top and bottom. The fiber bundle was wrapped one revolution around each of the two bars and was spread out over the entire gauge length in a parallel. For each set of chemical treatment, a minimum of fifty specimens were tested for determining the fiber tensile strength. According to ASTM standard D1294 and D1445 (1995), linear density or more commonly tex is calculated by:

$$D = \frac{W}{L}$$
(4.1)

Where: D = linear density or tex (mg/m)

W = mass of fibers (mg) L = length of fiber (m)

Also the unit break was calculated by:

$$UB = \frac{F}{D}$$
(4.2)

Where: F = maximum breaking load (mN)

D = linear density or tex (mg/m)

UB = unit break (mN/tex)

The Instron was set up to display a force-deformation curve at loading and to read the load at maximum or the break point. By inspecting the shape of this curve, one could check the accuracy of each individual test. If the sample was tensioned unevenly, more than one peak will appear on the graph. This is caused when some fibers are initially tensioned more than others. These fibers will break first, then the remaining fibers will break, causing an invalid test with two or more break points or graph peaks. Figure 4.7 shows how the graph can distinguish the valid results. With a valid test, there is one distinct peak that shows the true break force.



Figure 4.6 Set up for the tensile strength of fiber bundle.

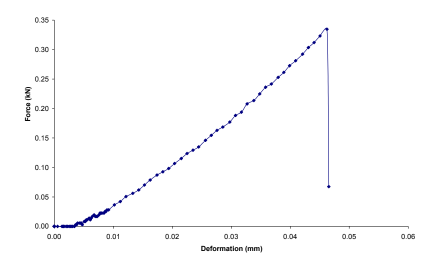


Figure 4.7 Force-deformation curve of a valid individual test.

4.6.2.2 Tensile strength of composites

Tensile property data are more useful in preferential selection of a particular type of plastic from a large group of plastic materials. The familiar dog-bone shape of the rotationally molded sample was utilized in the testing procedure. This type I specimen is the preferred specimen and should be used where sufficient material having a thickness of 7 mm or less is available. An Instron Universal testing machine (SATEC Systems, Inc., Grove City, PA) (Figure 4.8) was used to perform the tensile strength test at a crosshead speed of 5 mm/min as described in ASTM procedure D638-99 (ASTM 1999), and each test was performed until tensile failure occurred. The maximum (peak) load value (force) (F_{max}) was recorded by the instrument, which can be recalled after the completion of the test. The tensile strength at yield (σ_{ty}) is calculated from the following:

$$\sigma_{\rm ty} = \frac{F_{\rm max}}{A} \tag{4.3}$$

Where A is the cross sectional area.



Figure 4.8 Tensile strength test of a composite sample using the Instron Universal testing machine.

4.6.2.3 Tensile-impact strength of composites

The impact properties of the polymeric materials are directly related to the overall toughness of the material. The tensile-impact strength test was developed to overcome the deficiencies of flexural impact tests. Tensile-impact energy is the energy required to break a standard tension-impact specimen in tension by a single swing of a standard calibrated pendulum under a set of standard conditions.

Tensile-impact strength test was conducted according to ASTM D1822-93 (ASTM 1993). The tensile impact testing machine consists of a rigid massive base with a suspending frame. The pendulum is specially designed to hold the dumbbell-shaped specimen so that the specimen is not under stress until the moment of impact. Figure 4.9 shows the tensile impact tester (Tinius Olsen Testing Machines Co., Willow Grove, PA) used in the test.



Figure 4.9 Tensile-impact strength tester.

The specimens were prepared by rotational molding to the desired shape from a sheet. The type L (long) specimen extension is comparatively high. Type L specimens provide a greater differentiation between materials. Type L (long) specimens were prepared using the shape shown in Figure 4.10.



Figure 4.10 Type L tensile-impact specimens.

4.6.2.4 Durometer hardness of composites

Hardness is defined as the resistance of a material to deformation, particularly permanent deformation, indentation, or scratching. The Durometer hardness test is used for measuring the relative hardness of soft materials. The test method is based on the penetration of a specified indentor forced into the material, under specified conditions. Higher Durometer hardness readings are considered positive results.

Durometer readings were performed according to ASTM D2240-97 (ASTM 1998). The Durometer hardness tester (Shore Instrument and MFG Co., Freeport, NY) consists of a pressure foot, an indentor, and an indicating device. Two types of durometers are most commonly used: Type A and Type D. Due to the slightly harder sample being examined, the Type D gauge was used. The test was carried out by first placing a specimen on a hard, flat surface. The pressure foot of the instrument was pressed on to the specimen, making sure that it was parallel to the surface of the specimen. The Durometer hardness was read within 1 s after the pressure foot was in firm contact with the specimen. Each specimen was subjected to ten Durometer hardness readings, at designated positions on the sample bases. Values for these readings were then averaged. The Durometer hardness measuring instrument is shown in Figure 4.11.



Figure 4.11 Durometer hardness testing.

4.6.3 Physical properties

Water absorption is generally considered to be disadvantages, especially in composites. Migration of water through the polymer can lead to a disturbance of the fiber-matrix interface, reducing the overall strength and resulting in the dimensional instability of composites.

4.6.3.1 Moisture absorption of fibers

Prior to testing, the fibers were dried in an oven at 70°C for 24 h. Each sample was placed in the conditioning chamber for 72 h. Conditioning was conducted in the environmental test chamber (Angelantoni, ACS, Massa Martana, Italy) at 23°C and relative humidity values of 33, 66 and 100%, respectively. The mass of fibers was measured at different time intervals and the moisture absorption was calculated by the

Increase in mass (percent) =
$$\left(\frac{\mathbf{M}_{t} - \mathbf{M}_{0}}{\mathbf{M}_{0}}\right) \times 100$$
 (4.4)

mass difference. The percent increase in mass was calculated according to the following equation:

Where: M_t = mass of the sample after conditioning (g) (wet weight)

 M_0 = mass of the sample before conditioning (g) (dry weight)

4.6.3.2 Water absorption of composites

Water absorption characteristics of composites are altered by the addition of additives such as flax fibers because these additives showing a greater affinity to water. Rectangular specimens were cut from each sample with dimensions of 25.4 mm x 76.2 mm. The samples were dried in an oven at 50°C for 24 h, cooled in a desiccator, and immediately weighed to the nearest 0.001 g. In order to measure the water absorption of composites, all samples were immersed in water for about 24 h at room temperature as described in ASTM procedure D570-99 (ASTM 1999). Excess water on the surface of the samples was removed before weighing. The percentage increase in mass during immersion, was calculated to the nearest 0.01% as follows:

Increase in mass(percent) =
$$\frac{\text{Conditioned mass} - \text{Dry mass}}{\text{Dry mass}} \times 100$$
 (4.5)

The sample during water absorption test of composites is shown in Figure 4.12. Three replicate specimens were tested and the results were presented as average of the tested specimens.



Figure 4.12 Water absorption test of composites.

4.6.4 Melting points of composites

DSC is a thermoanalytical technique in which heat flow is measured as a function of temperature or time. Thermal analysis on pure materials, as well as on composites was performed using a DSC. The thermograms were then analyzed for any changes in the thermal behavior of the fibers. The DSC instrument is shown in Figure 4.13. A Perkin-Elmer DSC system (TA instruments, New Castle, DE) was used in this test.

Untreated and treated fiber composite sample weighing between 6 to 10 mg were placed in an aluminum pan and sealed with the crucible sealing press. The DSC system was operated in a dynamic mode with a heating scheme of -50 to 400°C, heating rate of 10°C/min and a chart of heat flow versus temperature was produced.



Figure 4.13 DSC melting point test of composites.

5. RESULTS AND DISCUSSION

The effects of three chemical treatments, namely, silane, benzoylation, and peroxide treatment on the performance of flax fibers and fiber-reinforced composites are discussed in this chapter.

5.1 Morphological Characterization

The possibility of forming mechanical bonding at the surface is mainly dependent on the surface topology of the fibers. It is important to mention that the changes of surface topography affect the interfacial adhesion. Fiber-matrix interface plays an important role in composite properties.

5.1.1 Fiber surface topology

Scanning electron microscopic analysis examined the surface topology of untreated and treated fibers. The removal of surface impurities on plant fibers is advantageous for fiber-matrix adhesion as it facilitates both mechanical interlocking and the bonding reaction due to the exposure of the hydroxyl groups to the chemicals used in treatment. Figure 5.1 shows the SEM photographs of fiber surfaces after chemical treatment. A porous structure is observed for untreated fibers. Figure 5.1b, c, and d shows the SEM photographs of fiber surfaces after silane treatment, benzoylation and dicumyl peroxide treatment.

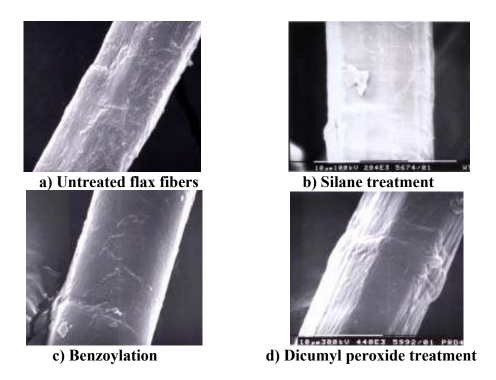


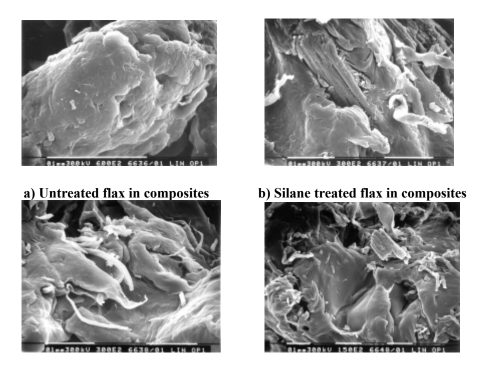
Figure 5.1 SEM photographs of fiber surfaces after chemical treatment.

These imagines indicate that after chemical treatment, the surfaces of the fibers became rougher enhancing the mechanical interlocking with resins. There is strong evidence that physical microstructure changes occurred at the fiber surface. The untreated flax fibers are in separated bundles with a smooth surface (Figure 5.1a). It is observed that silane treatment gave surface coating to the fibers, and surface features of fibers were not clearly visible. Since flax fibers exhibited micropores on theirs surface, the coupling agent penetrated into the pores and formed a mechanically interlocked coating on their surface. Benzoylation treatment led to major changes on the fiber surface. Smooth fiber surface is observed due to the substances deposited on the surface of the fiber. The surface topography is entirely modified after dicumyl peroxide treatment. The fibrillar structure of the individual ultimate fibers is revealed from the photograph and may be due to the leaching out of waxes and pectic substances. Micropores, particles adhering to

the surface, groove like portions and protruding structures made the fiber surface very rough. These effects on natural fibers are of particular importance for fiber-matrix adhesion and the creation of high fiber surface area required for the optimization of fiber-resin reinforcement. Therefore, the modification of cellulose fibers develops into changes in morphology and increase in hydroxyl groups. These changes will effectively result in improved surface tension, wetting ability, swelling, adhesion and compatibility with polymeric materials (Mohanty et al. 2001).

5.1.2 Composite microstructure

When it comes to using natural fibers as reinforcement in composite materials, many problems occur at the interface due to imperfect bonding. Interfacial properties of flax/polymer composites are largely determined by the strength and nature of secondary interactions that are established across the phase boundary. Interfacial stress transfer is therefore limited to relatively weak dispersion forces (Joseph et al. 2000). A strong fiber-matrix interface bond is critical for high mechanical properties of composites. When manufacturing composite materials, compatibility of the matrix and the fibers is also a problem. Therefore, modification of the fibers by chemical treatments is conducted to improve compatibility. These chemical reactions modify the properties of the fiber, and one of the roles of the cellulose fibers in composites is to give stiffness and strength to the polymeric matrix.



c) Benzoylation treated flax in composites d) Peroxide treated flax in composites Figure 5.2 SEM micrographs of LLDPE with 10% treated flax in composites.

Figure 5.2 shows the SEM photographs of the fiber-matrix interaction of untreated and surface treated flax/LLDPE composites. Scanning electron micrographs of the cross-session of the fractured tensile specimen for selected composites were carried out as an attempt to evaluate the fiber distribution within the composite which examined the failure mechanisms. Fiber breakage was the main failure criteria observed. Indeed, the untreated flax composite presented a very poor dispersion of the fiber which also exhibited some fiber agglomeration, small fiber breakage and very poor adhesion between fiber and matrix. On the other hand, chemically treated flax composites showed better fiber-matrix interaction. This is a result of a more uniform dispersion of fibers within the polymer matrix, thereby predicting micropores at the interface. The fibers were less agglomerated, showing the presence of some fibers dissociated into a matrix

polymer. It is important to mention that dissociation of the fiber into a matrix polymer may increase the surface area of the fibers, which contributes to a greater stress transfer area from the matrix to the fiber upon stress solicitation. The compatibility can be improved by grafting a matrix-compatible polymer onto the fiber surface.

The SEM micrographs of the cross-session of HDPE and LLDPE/HDPE flax fiberreinforced composites are found in Appendix H.

5.2 Mechanical Properties

Lignocellulosic fillers offer attractive properties, but are used only to a limited extent in industrial practice (Mishra et al. 2002). Natural fibers are strongly hydrophilic materials and moisture absorption leads to a significant deterioration of their mechanical properties. Furthermore, most polymers are hydrophobic and due to this divergent behaviour, the interface in natural fiber composites is rather poor. Any alteration of the characteristics of the cell wall, either chemical or morphological, has an effect on the mechanical properties of the fibers. By limiting the substitution reaction to the fiber surface, the good mechanical properties are reserved and a degree of biodegradability is maintained.

5.2.1 Tensile strength of fiber bundle

Flax fiber properties are controlled by the molecular fine structure of fibers. The chemical processing directly influences the cellulosic fine structure of plant fiber. Consequently, the chemical treatments have a lasting effect on the mechanical behavior

of flax fibers, especially on fiber strength and stiffness. It is very important to know the strength of the fibers before being combined into the thermoplastic matrix to understand better how the final composite behaves. Flax fiber tensile properties can be obtained by testing either single fibers or fiber bundles. Test results from the two methods are substantially different. Depending on the mechanism of the bundle breakage, this difference may be caused by the variations in fiber breaking elongations, breaking strengths and fiber crimps. From a fundamental point of view, the single or bundle fiber strength testing is not well understood. This is due, in part, to inadequate testing regimes, and because problems exist with the variability of mechanical properties of natural cellulosic fibers such as flax and hemp and are particularly, because it is difficult to measure. The fiber bundle fails within the hemicellulose and pectin layers that connect the single fibers together.

The single fiber testing method is too time-consuming and expensive compared to bundle test method. A method is developed in this thesis to test the tensile strength properties of flax fiber bundle. The average unit break of fiber bundle was tested based on fifty tests and the results are shown in Figure 5.3. The test was conducted on fiber bundle with a gauge length of 40 mm at standard laboratory atmosphere of 23°C and relative humidity of 51%. The data show that the higher strength (although not statistically significant) of silane- and peroxide-treated fibers compared to untreated fibers may be a result of the removal of surface imperfections after the treatment. The increased uniformity of the fibers would give an increase to strength, as points of unconformity are removed during the treatment and this changes the deformation behavior of the fibers (Mohanty et al. 2001). On the other hand, the average unit break reached a low of 107.33 mN/tex when using benzoylated samples, due to breakage of the bond structure. The testing procedure used linear density instead of the area to evaluate the unit break of fiber bundle. Thus, the results can not be reverse engineered to a MPa reading without making assumptions about the fiber bundle, or drastically increasing the testing time. Unfortunately, the results of this test do not lend themselves to comparison with other research results such as those listed in Table 3.1, where the tensile strength of flax was 345-1100 MPa (Mohanty et al. 2000a). This test also used numerous fibers and not individual fibers. This enables some of the errors associated with individual fiber testing to be eliminated, such as assuming the fibers are perfectly round and the breaking point is located at the same spot where the area was measured.

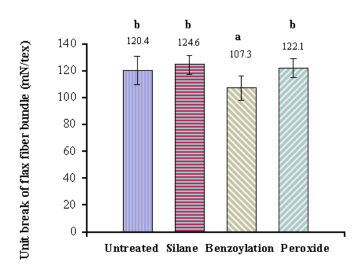
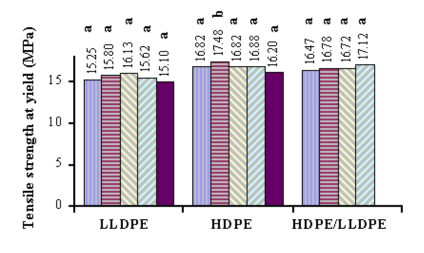


Figure 5.3 Average unit break of flax fiber bundle. Each bar shows the mean value with a range according to 95% confidence interval. Means with the same letter designation are not significantly different at P=0.05, n=50.

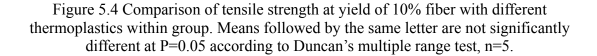
5.2.2 Tensile strength of composites

Certain plant fibers, notably flax and hemp possess tensile properties which make them

potentially attractive for use as reinforcement in polymer matrix composites. The effect of chemical treatment on the tensile properties of flax fiber-reinforced composites was investigated. Treatments using chemicals such as triethoxyvinylsilane coupling agent, benzoyl chloride and dicumyl peroxide were carried out to improve the bonding at the fiber-polymer interface. Figure 5.4 shows the tensile strength at yield of fiber-reinforced LLDPE, HDPE and LLDPE/HDPE composites. Compared to the untreated fiber composite having 10% by weight fiber loading, the pre-treatments slightly enhanced the tensile strength of the composites, but not in a statistically significant amount. For silane-treated fibers with LLDPE, the tensile strength increased from 15.10 to 15.80 MPa compared to 15.25 MPa for the untreated fiber composite processed in the same manner. This may be due to the increased fiber-matrix adhesion, their rough surface topography and the peroxide-induced grafting. The tensile strength of flax fiberreinforced composites is determined both by the tensile strength of the fiber and by the presence of weak lateral fiber bonds (Mohanty et al. 2001). The variations in the tensile strength at yield of the composites using different fiber treatments were attributed to the changes in the chemical structure and bondability of the fiber. Tests with different flax fiber-reinforced biodegradable matrix polymers showed that the tensile strength of these biocomposites was clearly influenced by the particular matrix and the adhesion between fiber and matrix. In comparison with the virgin polymer, all the treatments showed the same tendency to slightly increase (although not statistically significant) the tensile strengths of modified composites. The processing employed also played an important role on the tensile properties of the composites.



🔲 untreated 📋 silane treatment 🔝 Benzoylation 📝 Peroxide treatment 💼 Resin



5.2.3 Tensile-impact strength of composites

The tensile-impact strength of composites with 10% pre-treated flax fibers compared to composites with 10% untreated flax fibers composites with different types of thermoplastic matrix is shown in Figure 5.5. The influence of chemical modification on the tensile-impact strength of composites is also represented in Figure 5.5. It is observed that reinforcement of composites with treated flax fiber slightly enhanced (although not statistically significant) the tensile-impact strength of the resulting composite.

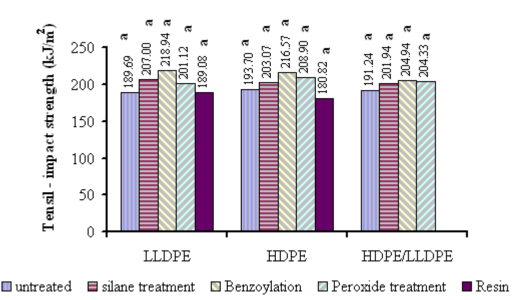


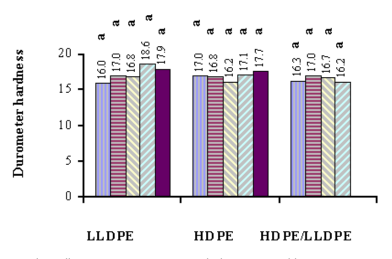
Figure 5.5 Tensile-impact strength of 10% pre-treated flax fibers compared to untreated flax fibers composites within group. Means followed by the same letter are not significantly different at P=0.05 according to Duncan's multiple range test, n=5.

5.2.4 Durometer hardness of composites

The data and numerical results from each Durometer hardness test are presented in this section. Figure 5.6 shows the hardness of 10% flax fiber composites with different types of thermoplastic matrix. The hardness of plastics is measured by the Shore (Durometer) test. This method measures the resistance of plastics to indentation and provides an empirical hardness value that does not correlate well to other properties or fundamental characteristics. The hardness value is determined by the penetration of the Durometer indenter foot into the sample. The results obtained from this test are a useful measure of relative resistance to indentation of various grades of polymers.

Ten readings were taken for each specimen, as material properties were expected to vary with location on the sample. For the 10% fiber-based composites, chemically treated

flax fibers did not increase the hardness of specimens. When compared across fiber pretreatment types, composites containing chemically treated fibers with LLDPE had higher hardness (although not statistically significant) than composites containing untreated fibers.



🔲 Untreated 🚍 Silane treatment 🔄 Benzoylation 📨 Peroxide treatment 💼 Resin

Figure 5.6 Durometer Hardness of 10% flax fibers composites with different thermoplastics within group. Means followed by the same letter are not significantly different at P=0.05 according to Duncan's multiple range test, n=10.

5.3 **Physical Properties**

The hydrophilic nature of fibers leads to biocomposites having high water absorption characteristics. Before making the composite, the moisture absorption of flax fibers should be reduced. During chemical treatment of the flax fiber, the hemicellulose and lignin were separated and cellulose was used for the biocomposite. Pre-treatment of the flax fiber replaced some of the hydroxyl groups in the cell wall of the flax molecule, which reduced the hygroscopic nature of the flax fiber-reinforced composites (Sreekala et al. 2000).

5.3.1 Moisture absorption of fibers

Figure 5.7 shows the moisture absorption of untreated and treated flax fibers at different relative humidities. The moisture absorption of the chemically treated flax fiber was lower than that of untreated flax fibers. This result shows that chemical treatments can decrease the moisture absorption of the fibers and, hopefully it may also lead to the biocomposites having low moisture absorption characteristics.

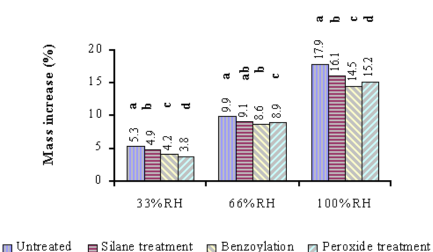
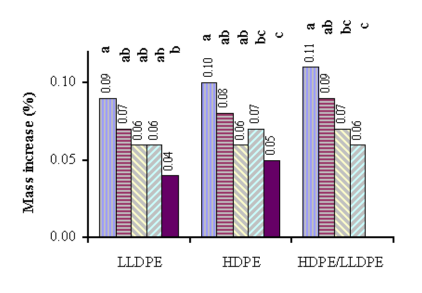


Figure 5.7 Moisture absorption of untreated and treated flax fibers at different relative humidity. Means followed by the different letter are statistically different from the untreated sample at P=0.05 according to Duncan's multiple range test, n=3.

5.3.2 Water absorption of composites

The water absorption of untreated and chemically modified flax fiber-based composites is presented in Figure 5.8 as a percentage of dry weight after 24 h immersion in water. The results show that the water absorption of the chemically treated flax fiber-based composites was lower than that of the untreated fiber-based composites. The untreated composites absorbed the most water and the peroxide-treated composites absorbed the least, suggesting that changes in surface chemistry have reduced the affinity of fibers to moisture. Strong intermolecular fiber-matrix bonding decreased the rate of moisture absorption in biocomposites. It shows that chemical treatments of flax fiber can decrease the water absorption of the biocomposites.



🔲 Untreated 📋 Silane treatment 🔝 Benzoylation 🖾 Peroxide treatment 🔳 Resin

Figure 5.8 Water absorption of 10% fiber-based composites within group. Means followed by the different letter are statistically different from the untreated sample at P=0.05 according to Duncan's multiple range test, n=3.

5.4 Melting Points of Composites

Analyzing the DSC thermograms showed that the melting range of flax fiber-reinforced composites is displayed as an endothermic peak. DSC analysis enables the identification of chemical activity occurring in the fiber as heat is applied. DSC was used to determine not only the melting point (T_m) but also the melting range of the polymer. The glass transition temperature (T_g) could not be observed. The T_g of pure polyethylene is usually below -100°C. Composites containing untreated fiber and silane-, benzoyl- and

peroxide-treated fiber also did not display T_g . Table 5.1 shows the melting points of pure LLDPE, pure HDPE, untreated fiber-polymer composites and treated fiber-polymer composites. The melting point of pure LLDPE and HDPE was found to be 129.9°C and 130.7°C respectively. Changing in the melting point temperature of the polymers due to the fiber incorporation was observed. The addition of untreated/treated fiber in LLDPE increased the T_m of LLDPE compared to that of pure LLDPE. The increase of T_m may be attributed to the plasticization effect of the fiber that diffuses or dissolves into the polymer. The incorporation of 10% chemically treated flax fiber in LLDPE/HDPE composites increased the T_m of untreated fiber composites. The extent of increase in T_m is more pronounced in the case of silane-treated fiber. The increased melting point of composites meant that thermal resistance increased as elucidated by the DSC method.

Variable	Melting Point (°C)
LLDPE + U	130.6
LLDPE + S	130.3
LLDPE + B	130.2
LLDPE + P	129.2
LLDPE (pure)	129.9
HDPE + U	133.4
HDPE + S	133.1
HDPE + B	132.9
HDPE + P	132.7
HDPE (pure)	130.7
HDPE/LLDPE + U	130.0
HDPE/LLDPE + S	131.6
HDPE/LLDPE + B	130.4
HDPE/LLDPE + P	130.2

Table 5.1. Melting points of composites.

There is only one replicate in these tests.

DSC thermograms of untreated and chemically treated flax fiber-reinforced composites are in Appendix G.

6. SUMMARY AND CONCLUSION

In this chapter, the conclusions based on results obtained from the experiments are illustrated. Statistically significant results were also obtained. The data presented are only representative of those that chose to participate in the study. The major objective of this thesis was to focus on three chemical treatments of flax fibers for use in fiber-reinforced composite to achieve improved properties of composites.

Renewable fibers like flax can be used as reinforcing materials for low cost composites, due to the economic and environmental advantages of such materials. However, flax fiber is highly hydrophilic due to the presence of hydroxyl groups from cellulose and lignin. Chemical treatment can reduce the hydrophilicity of the fiber by treating these fibers with suitable chemicals to decrease the hydroxyl groups in the fibers. The first objective of this thesis was to investigate suitable fiber pre-treatment methods on the effects of flax fiber properties. The tensile strength and moisture absorption data were used to address this objective. In terms of moisture absorption of fibers, all three pretreatments were effective in reducing the hydrophilic nature of fibers. In terms of the fiber bundle strength, silane treatment slightly improved the unit break of fiber bundle (although not statistically significant) more than the other fiber pre-treatments.

The composite properties were greatly influenced by the technique as well as the

processing method used. Processing produces a great variation on the dimension and dispersion of the fiber within the composite. Understanding the relationship between processing and the properties of the composites is of key importance to obtain materials with optimized performance. From initial studies comparing single-screw and twinscrew extruders, the latter worked better in producing good quality strand of composites. Thus, it can be concluded that compounding fiber and polymer-matrix by using a twinscrew extruder could be an effective processing method prior to rotational molding. Chemically treated flax fiber is used as filler to plastics and reinforcement in thermoplastic matrix in rotational molding process. The addition of fibers can potentially lower material costs, as the fibers are available at a cheaper price than glass fibers or other inorganic additives.

The final objective was to study the effects of fiber surface modifications on the performance of the flax fiber-reinforced composites. Morphological study, mechanical properties data, physical properties data and thermal properties data were used to address this objective. Morphological and structural changes of the fibers were investigated by using scanning electron microscopy. The coupling agents were found to be effective in improving the surface properties of flax fiber, forming a mechanically interlocked coating on its surface. Therefore, physical microstructure changes occurred to the fiber surface by chemical treatment. Fiber-matrix interface plays an important role in composite properties. The ability to control the chemical and mechanical properties of the fiber-matrix interface is crucial. Morphological studies showed that the pre-treatments improved the fiber-matrix adhesion and the dispersion of the particles.

A method was developed to test quickly and accurately the tensile strength properties of flax fiber. This involved a procedure where the fibers were parallelized, conditioned and tested with the Instron testing machine. The fiber bundles break in the gauge length zone and not beside the clamping areas showing that fiber strength was not influenced by the testing apparatus. Silane and peroxide treatment on flax fiber bundle lead to a higher (although not statistically significant) tensile strength than that of the untreated fiber bundle. Chemically treated samples with a high tensile strength were considered to be quite acceptable for a biological material. Compared to the untreated fiber-based composite, tensile properties were improved with a suitable fiber surface treatment. Silane-, benzoyl-, and peroxide-treated fiber composites offered superior physical and mechanical properties. Mechanical properties of natural fiber-reinforced plastic composites could be improved by the use of a silane coupling agent.

The hydrophilic nature of biofibers leads to biocomposites with high water absorption characteristics that can be overcome by treating these fibers with suitable chemicals to decrease the hydroxyl groups of the fibers. The water absorption and swelling of the treated flax fiber composites were lower than those of untreated flax fiber composites.

A validation of the method was made on the samples with differential scanning calorimetry (DSC). The incorporation of chemically treated/untreated flax fiber into LLDPE or HDPE considerably increased the melting point (T_m) of the composites. Incorporation of 10% chemically treated flax fiber in LLDPE/HDPE composites increased the T_m of untreated fiber composites. The T_m increase is more pronounced in

the case of silane treated fiber.

The flax fiber is already being produced and can be obtained at a relatively low cost compared to glass fiber reinforcements. Flax fiber has a very promising future in the rotational molding industry. This research presently being conducted at the University of Saskatchewan will hopefully benefit flax growers.

7. RECOMMENDATIONS

New materials derived from biofibers and thermoplastic polymers which may substitute for wood flour under particular circumstances of local supplies and cost have attracted considerable attention in recent years. The development of this novel composite type has presented new versions of common problems, such as material handling and processing. Substantial progress has been made toward overcoming the technological problems associated with compounding and extrusion of fiber-reinforced composites.

Normal methods of modifying the interface are usually not applicable in natural fibers for many reasons, cost being the most important. Natural fibers cannot compete in terms of strength with man-made fibers, but their main advantage is their low cost. Therefore, a cost effective modification method is needed to enhance the interface in natural fiber composites.

From the results of these experiments, it is quite evident that flax fiber has a very promising future and can be used as a substitute for glass fibers. Surface modifications of hydrophilic natural fibers have achieved some degree of success in making a superior interface, mechanical properties and thermal properties, but lower cost surface modification needs to be emphasized for biocomposites to replace glass fiber composites in many applications in the future. Natural fiber-reinforced composites should be developed and characterized so as to produce cost-competitive biocomposites for industrial applications. A thorough study is needed to evaluate the optimum levels and to make general conclusions to commercialize the use of biorenewable fibers like flax.

The exploitation of natural fibers in industrial applications provides challenges to come up with effective ways of both analyzing the modes of deformation of such materials, and in effecting the adhesion between matrix and fiber. In conjunction with this, there is a need to understand fully the basic structural components of the fibers, and their effect on the mechanical properties. The interface is one such area that has attracted a lot of interest, and new and cost effective ways of treating cellulose fibers, with the desirable end-properties, can only benefit exploitation. In the future, combination of modern testing techniques, such as Raman and miniature tensile testing, and the modification of fiber surface will lead to a better knowledge of desirable properties. The development of better processing for releasing the fibers could be another focus for the research community. Finally, to understand and measure properly the role of the interface, it is thought that a combination of techniques may be the only way to fully achieve this goal.

Processing conditions and surface treatments need to be further optimized to achieve improved properties of fiber-reinforced composites.

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APPENDICES

APPENDIX A

Tensile Strength of Fiber Bundle

Experimental conditions for the tensile strength of fiber bundle presented in Tables A1 to A4.

Temperature (F): 73 Humidity (%): 50 Version : 1.08 Version date : 30 May 1991 Machine : 1100 Operator : bei Bar type : yarn Entry dimens : YES Units type : SI Gauge length of 40mm Test speed of 5 mm/min

Table A1. Tensile strength of untreated fiber bundle.

No.	Load at break (N)	Unit Break (mN/tex)	Mass of fiber (mg)	Length of fiber (m)	D (tex) =W/L
1	1.59	119.14	3.60	0.27	13.33
2	1.04	79.75	2.60	0.20	13.00
3	2.76	149.37	3.70	0.20	18.50
4	2.85	157.63	3.80	0.21	18.10
5	1.51	114.77	2.90	0.22	13.18
6	1.13	87.56	2.70	0.21	12.86
7	1.45	103.61	2.80	0.20	14.00
8	1.11	80.09	2.50	0.18	13.89
9	0.77	61.29	2.40	0.19	12.63
10	2.96	171.57	3.80	0.22	17.27
11	1.95	142.92	3.00	0.22	13.64
12	2.64	162.98	3.40	0.21	16.19
13	1.85	121.64	3.50	0.23	15.22
14	1.33	100.59	2.90	0.22	13.18
15	4.05	188.34	4.30	0.20	21.50
16	1.64	110.93	3.10	0.21	14.76
17	0.77	59.56	2.60	0.20	13.00
18	1.36	102.12	2.80	0.21	13.33
19	2.49	150.75	3.30	0.20	16.50
20	1.35	105.94	2.80	0.22	12.73
21	1.51	105.90	3.00	0.21	14.29
22	0.84	71.70	2.10	0.18	11.67
23	1.30	96.25	2.70	0.20	13.50
24	1.93	116.77	3.30	0.20	16.50
25	1.40	94.95	3.10	0.21	14.76

Table A1	continued				
26	2.79	161.52	3.80	0.22	17.27
27	1.00	83.43	2.40	0.20	12.00
28	1.48	103.96	2.70	0.19	14.21
29	2.95	182.22	3.40	0.21	16.19
30	1.49	112.95	2.50	0.19	13.16
31	2.83	172.67	3.60	0.22	16.36
32	2.11	147.95	3.00	0.21	14.29
33	3.09	166.22	3.90	0.21	18.57
34	2.07	115.88	3.40	0.19	17.89
35	1.45	100.04	2.90	0.20	14.50
36	1.86	122.35	3.20	0.21	15.24
37	1.20	84.54	2.70	0.19	14.21
38	3.46	182.20	3.80	0.20	19.00
39	3.95	207.45	4.00	0.21	19.05
40	0.84	75.29	2.00	0.18	11.11
41	1.84	114.07	2.90	0.18	16.11
42	2.89	155.50	3.90	0.21	18.57
43	2.01	121.90	3.30	0.20	16.50
44	1.40	102.43	2.60	0.19	13.68
45	2.47	167.60	3.10	0.21	14.76
46	0.86	65.84	3.00	0.23	13.04
47	2.09	143.48	3.20	0.22	14.55
48	1.36	113.47	2.40	0.20	12.00
49	0.94	71.98	3.00	0.23	13.04
50	1.35	86.98	3.10	0.20	15.50
SUM		6018.06			
Average (UB _{ave})		120.36			
Standard Deviation (S)		37.31			
Coef. Var. (CV)		31.00			
95% confidence interval		Upper	Lower		
		131.07	109.65		

Linear Density or more commonly tex is calculated by:

$$D = \frac{W}{T}$$

L Where: D = linear density or tex (mg/m) W = mass of fibers (mg) L = length of fiber (m)

Unit Break is calculated by:

$$UB = \frac{F}{D}$$

Where: F = maximum breaking load (mN) D = linear density or tex (mg/m) UB = unit break (mN/tex) UB_{ave} = average unit break (mN/tex)

No.	Load at break (N)	Load at break (N) Unit Break (mN/tex)		Length of fiber (m)	D (tex) =W/L
1	2.07	162.00	3.20	0.25	12.80
2	1.64	131.00	3.00	0.24	12.50
3	1.94	115.36	3.70	0.22	16.82
4	1.85	115.07	3.70	0.23	16.09
5	1.55	138.83	2.90	0.26	11.15
6	2.84	199.49	3.70	0.26	14.23
7	1.16	106.46	2.40	0.22	10.91
8	1.96	130.87	3.00	0.20	15.00
9	1.55	114.70	2.70	0.20	13.50
10	1.39	109.08	2.80	0.22	12.73
11	1.44	110.19	3.00	0.23	13.04
12	2.85	176.17	3.40	0.21	16.19
13	2.18	166.82	3.00	0.23	13.04
14	1.86	128.58	2.90	0.20	14.50
15	2.01	129.76	3.10	0.20	15.50
16	1.02	102.34	2.00	0.20	10.00
17	2.16	156.60	2.90	0.21	13.81
18	1.56	124.95	2.50	0.20	12.50
19	1.25	105.03	2.50	0.21	11.90
20	1.98	131.71	3.30	0.22	15.00
21	1.28	106.42	2.40	0.20	12.00
22	1.56	126.15	2.60	0.21	12.38
23	1.17	107.26	2.30	0.21	10.95
24	0.77	93.73	1.90	0.23	8.26
25	1.33	108.05	2.70	0.22	12.27
26	1.16	101.62	2.40	0.21	11.43
27	0.97	92.58	2.00	0.19	10.53
28	2.83	206.15	3.70	0.27	13.70
29	1.15	120.49	2.20	0.23	9.57
30	1.10	125.61	2.10	0.24	8.75
31	1.59	120.47	2.90	0.22	13.18
32	1.64	116.96	2.80	0.20	14.00
33	1.66	138.68	3.00	0.25	12.00
34	1.17	122.58	2.30	0.24	9.58
35	1.91	127.56	3.30	0.22	15.00
36	1.66	109.21	3.20	0.21	15.24
37	1.56	120.92	3.10	0.24	12.92
38	1.68	116.92	3.30	0.23	14.35
39	1.15	100.22	2.30	0.20	11.50
40	1.51	145.24	2.50	0.24	10.42
41	1.76	127.60	2.90	0.21	13.81
42	1.34	104.82	2.30	0.18	12.78
43	1.13	97.23	2.20	0.19	11.58
44	0.94	103.77	1.90	0.21	9.05
45	1.37	143.23	2.40	0.25	9.60
46	1.25	104.20	2.40	0.20	12.00

Table A2. Tensile strength of silane treated fiber bundle.

Table A2 co	Table A2 continued							
47	1.02	94.16	2.50	0.23	10.87			
48	1.33	126.84	2.30	0.22	10.45			
49	1.59	146.64	2.60	0.24	10.83			
50	1.49	118.90	2.50	0.20	12.50			
SUM		6229.17						
Average (UB _{ave})	124.58						
Standard D	Deviation (S)	24.86						
Coef. Var. (CV)		19.95						
95% confi	dence interval	Upper	Lower					
9370 confidence interval		131.72	117.45					

No.	Load at break (N)	Unit Break (mN/tex)	Mass of fiber (mg)	Length of fiber (m)	D (tex) =W/L
1	1.40	128.95	2.50	0.23	10.87
2	1.15	137.20	2.10	0.25	8.40
3	0.93	112.04	1.90	0.23	8.26
4	0.81	114.96	1.70	0.24	7.08
5	0.81	104.05	1.80	0.23	7.83
6	1.05	125.02	2.10	0.25	8.40
7	1.10	120.90	2.00	0.22	9.09
8	0.56	80.60	1.60	0.23	6.96
9	1.01	110.28	2.30	0.25	9.20
10	1.03	107.38	2.10	0.22	9.55
11	0.69	90.52	1.60	0.21	7.62
12	0.55	80.93	1.50	0.22	6.82
13	0.56	82.23	1.50	0.22	6.82
14	1.02	114.00	1.70	0.19	8.95
15	0.78	71.50	2.40	0.22	10.91
16	1.11	155.40	1.50	0.21	7.14
17	0.59	47.65	2.60	0.21	12.38
18	0.67	63.65	2.00	0.19	10.53
19	0.85	58.62	2.90	0.20	14.50
20	0.87	130.50	1.40	0.21	6.67
21	1.20	120.00	1.90	0.19	10.00
22	0.66	60.00	2.20	0.20	11.00
23	0.59	72.88	1.70	0.21	8.10
24	1.00	100.00	1.80	0.18	10.00
25	1.04	93.60	2.00	0.18	11.11
26	0.98	171.50	1.20	0.21	5.71
27	0.96	101.05	1.90	0.20	9.50
28	0.74	74.00	1.90	0.19	10.00
29	0.83	79.39	2.30	0.22	10.45
30	0.96	138.00	1.60	0.23	6.96
31	1.02	106.86	2.10	0.22	9.55
32	0.95	118.75	2.00	0.25	8.00
33	1.04	199.33	1.20	0.23	5.22
34	0.87	116.00	1.50	0.20	7.50
35	1.06	111.89	1.80	0.19	9.47
36	0.69	90.79	1.90	0.25	7.60
37	0.59	71.83	2.30	0.28	8.21
38	0.79	82.29	2.40	0.25	9.60
39	0.89	97.90	2.00	0.22	9.09
40	1.20	184.00	1.50	0.23	6.52
40	0.83	72.17	2.30	0.20	11.50
42	0.94	119.07	1.50	0.19	7.89
42	0.94	152.43	1.40	0.22	6.36
43 44	0.76	104.50	1.60	0.22	0.30 7.27
44 45	0.93	103.33	1.80	0.22	9.00
43 46	1.01	132.89	1.80	0.20	9.00 7.60

Table A3. Tensile strength of benzoylation treated fiber bundle.

Table A.	3 continued				
47	1.09	103.55	2.00	0.19	10.53
48	0.85	115.36	1.40	0.19	7.37
49	0.96	114.29	2.10	0.25	8.40
50	0.99	122.29	1.70	0.21	8.10
SUM		5366.32			
Average	(UB _{ave})	107.33			
Standard	Deviation (S)	31.61			
Coef. Va	r. (CV)	29.45			
95% confidence interval		Upper	Lower		
		116.40	98.25		

No.	Load at break (N)	Unit Break (mN/tex)	Mass of fiber (mg)	Length of fiber (m)	D (tex) =W/L
1	2.25	174.31	3.10	0.24	12.92
2	0.77	91.09	1.70	0.20	8.50
3	1.02	113.12	1.90	0.21	9.05
4	1.55	137.12	2.60	0.23	11.30
5	1.33	126.84	2.30	0.22	10.45
6	2.14	172.61	3.10	0.25	12.40
7	1.29	123.01	2.30	0.22	10.45
8	1.02	102.34	1.90	0.19	10.00
9	1.73	147.07	2.70	0.23	11.74
10	0.77	95.64	1.70	0.21	8.10
11	1.80	146.84	2.70	0.22	12.27
12	1.90	142.50	2.80	0.21	13.33
13	1.79	139.13	2.70	0.21	12.86
14	2.22	170.57	3.00	0.23	13.04
15	1.51	128.01	2.60	0.22	11.82
16	1.09	124.86	2.00	0.23	8.70
17	1.49	130.04	2.40	0.21	11.43
18	0.55	73.57	1.50	0.20	7.50
19	1.48	135.91	2.50	0.23	10.87
20	1.35	134.83	2.40	0.24	10.00
21	2.15	157.61	3.00	0.22	13.64
22	1.94	145.51	2.80	0.21	13.33
23	0.57	75.34	1.60	0.21	7.62
24	1.55	126.17	2.70	0.22	12.27
25	0.68	79.57	1.70	0.20	8.50
26	1.50	118.71	2.40	0.19	12.63
27	1.65	117.92	2.80	0.20	14.00
28	0.96	105.73	2.00	0.22	9.09
29	2.07	153.85	3.10	0.23	13.48
30	1.44	143.73	2.40	0.24	10.00
31	1.21	116.20	2.30	0.22	10.45
32	1.62	138.35	2.70	0.23	11.74
33	1.56	144.62	2.70	0.25	10.80
34	0.85	94.43	1.80	0.20	9.00
35	1.19	124.46	2.10	0.22	9.55
36	1.30	124.03	2.20	0.21	10.48
37	0.81	105.38	1.70	0.22	7.73
38	0.66	95.31	1.60	0.23	6.96
39	0.70	87.33	1.60	0.20	8.00
40	0.94	113.66	1.90	0.23	8.26
41	1.28	121.90	2.20	0.21	10.48
42	1.02	113.12	1.90	0.21	9.05
43	0.99	119.58	1.90	0.23	8.26
44	1.01	121.74	2.00	0.24	8.33
45	1.26	126.37	2.20	0.22	10.00
46	0.91	100.82	1.90	0.21	9.05

Table A4. Tensile strength of peroxide treated fiber bundle.

Table A4	4 continued				
47	1.08	117.94	2.10	0.23	9.13
48	1.08	118.45	2.00	0.22	9.09
49	0.59	69.75	1.60	0.19	8.42
50	1.01	116.67	2.00	0.23	8.70
SUM		6103.64			
Average	(UB _{ave})	122.07			
Standard	Deviation (S)	24.57			
Coef. Var. (CV)		20.13			
95% confidence interval		Upper	Lower		
		129.13	115.02		

Variable	N	Mean (mN/tex)	Std. Deviation	Std. Error	95% Confidence Interval for Mean (mN/tex)		UB (m	N/tex)
		(mi (itex)	(mN/tex) ⁽ⁿ	(mN/tex)	Lower	Upper	Minimum	Maximum
Untreated	50	120.36	37.31	5.28	109.76	130.96	59.56	207.45
Silane	50	124.58	24.86	3.52	117.52	131.65	92.58	206.15
Benzoylation	50	107.33	31.42	4.44	98.40	116.26	47.65	199.33
Peroxide	50	122.07	24.57	3.47	115.09	129.06	69.75	174.31
Total	200	118.59	30.52	2.16	114.33	122.84	47.65	207.45

Table A5. Statistical analysis of tensile strength of fiber bundle.

One way Analysis of Variance

	df	df Sum of Squares		Computed	Computed		
	ui	Sum of Squares	Mean Square	F	Sig. (0.05)		
Between Groups	3	8903.49	2967.83	3.30*	0.022		
Within Groups	196	176438.09	900.19				
Total	199	185341.58					
^{ns} = not significant, *= significant at 5% level, ** = significant at 1% level.							

Multiple Comparison	S

	(I)	(J)	Mean Difference	Std. Com	Computed	95% Confidence Interval for Mean		
	(1)	(9)	(I-J)	Error	Sig. (0.05)	Lower	Upper	
LSD	Untreated	Silane	-4.22	6.00	0.482 ^{ns}	-16.0577	7.6105	
		Benzoylation	13.03(*)	6.00	0.031*	1.2003	24.8685	
		Peroxide	-1.71	6.00	0.776^{ns}	-13.5465	10.1217	
	Silane	Benzoylation	17.26(*)	6.00	0.004^{**}	5.4239	29.0921	
		Peroxide	2.51	6.00	0.676^{ns}	-9.3229	14.3453	
	Benzoylation	Peroxide	-14.75(*)	6.00	0.015*	-26.5809	-2.9127	
. The	mean difference	is significant at P	=0.05 level.					

APPENDIX B

Tensile Strength of Composites

Experimental conditions for the tensile strength of composites presented in Tables B1 and B2.

Temperature (°C): 23 Humidity (%): 50 ASTM procedure: D638-99 (ASTM 1999) Type I specimen: familiar dog-bone shape Machine : Instron Universal testing machine (SATEC Systems, Inc., Grove City, PA) Operator : bei Units type : SI Test speed of 5 mm/min

Composite	Load at Yield (N)	Width (mm)	Thickness (mm)	Cross Sectional Area (mm ²)	$\sigma_{tv} = F/A$ (MPa)
LLDPE+U	875	0.0129	0.00449	5.78761E-05	15.12
LEDIE	900	0.0129	0.00461	5.87775E-05	15.31
	925	0.0120	0.00455	0.000059696	15.50
	900	0.0127	0.00485	6.16435E-05	14.60
	875	0.0124	0.00449	5.56311E-05	15.73
LLDPE+S	900	0.0121	0.00456	5.75472E-05	15.64
LEDIE	875	0.0120	0.00430	5.34252E-05	16.38
	825	0.0127	0.00422	5.34204E-05	15.44
	800	0.0125	0.00421	4.87939E-05	16.40
	1000	0.0110	0.00421	0.000066101	15.13
LLDPE+B	800	0.0113	0.00449	5.13207E-05	15.59
	875	0.0114	0.00453	5.40429E-05	16.19
	950	0.0113	0.00504	6.18912E-05	15.35
	850	0.0123	0.00458	5.12044E-05	16.60
	850	0.0112	0.00438	5.02875E-05	16.90
LLDPE+P	975	0.0115	0.00458	5.28532E-05	18.45
	975	0.0115	0.00534	0.000067284	14.49
	925	0.0120	0.00508	0.00006223	14.49
	900	0.0123	0.00513	6.07392E-05	14.82
	900	0.0118	0.00513	0.000063	14.82
HDPE+U	870	0.0123	0.00304	5.00332E-05	17.39
IIDI E+O	850	0.0117	0.00428	5.46116E-05	15.56
	900	0.0124	0.00439	5.05908E-05	17.79
	900	0.0120	0.00423	5.52568E-05	16.47
	920	0.0116	0.00478	0.000054516	16.88
HDPE+S	920 780	0.0116	0.00472	0.000034318	10.88
HDPE+5	825	0.0108	0.00430	4.86245E-05	17.05
	890 825	0.0120	0.00394	0.000047083	18.90
	825	0.0116	0.00411	4.76349E-05	17.32

Table B1. Tensile strength at yield (σ_{ty}) of composites (10% fiber).

Table B1 continued					
	740	0.0107	0.00402	4.30542E-05	17.19
HDPE+B	895	0.0129	0.00433	5.60302E-05	15.97
	750	0.0116	0.00373	4.31188E-05	17.39
	740	0.0118	0.00363	4.27977E-05	17.29
	760	0.0116	0.00393	4.57452E-05	16.61
	775	0.0120	0.00385	0.000046046	16.83
HDPE+P	905	0.0117	0.00451	5.29474E-05	17.09
	900	0.0124	0.00431	5.32716E-05	16.89
	860	0.0125	0.00409	5.10023E-05	16.86
	875	0.0129	0.00414	5.34888E-05	16.36
	925	0.0125	0.00429	5.37537E-05	17.21
HDPE/LLDPE+U	860	0.0123	0.00484	0.00005929	14.50
	805	0.0112	0.00424	4.75728E-05	16.92
	875	0.0120	0.00441	5.30082E-05	16.51
	825	0.0114	0.00420	0.00004767	17.31
	850	0.0117	0.00423	4.96602E-05	17.12
HDPE/LLDPE+S	875	0.0122	0.00433	5.29992E-05	16.51
	930	0.0116	0.00454	5.24824E-05	17.72
	940	0.0129	0.00434	5.58558E-05	16.83
	875	0.0123	0.00418	5.12468E-05	17.07
	975	0.0123	0.00502	6.18966E-05	15.75
HDPE/LLDPE+B	925	0.0131	0.00420	0.000055062	16.80
	850	0.0120	0.00446	0.00005352	15.88
	875	0.0118	0.00450	0.00005301	16.51
	815	0.0115	0.00411	4.71828E-05	17.27
	825	0.0117	0.00412	4.81628E-05	17.13
HDPE/LLDPE+P	935	0.0117	0.00474	5.55528E-05	16.83
	990	0.0119	0.00506	0.000059961	16.51
	900	0.0116	0.00437	5.08231E-05	17.71
	985	0.0120	0.00465	5.55675E-05	17.73
	975	0.0118	0.00491	5.78889E-05	16.84

The tensile strength at yield (σ_{ty}) is calculated from the following:

$$\sigma_{ty} = \frac{F_{max}}{A}$$

Where: F_{max} = the maximum (peak) load value (force)

A = the cross sectional area (m^2)

Composite	Load at Yield (N)	Width (mm)	Thickness (mm)	Cross Sectional Area (mm ²)	$\sigma_{tv} = F/A$ (MPa)
LLDPE	900	0.0113	0.00470	0.000053157	16.93
	850	0.0112	0.00512	0.000057344	14.82
	825	0.0118	0.00489	5.76042E-05	14.32
	975	0.0117	0.00509	0.000059553	16.37
	825	0.0125	0.00506	0.00006325	13.04
HDPE	875	0.0119	0.00464	0.000055216	15.85
	900	0.0116	0.00456	0.000052896	17.01
	815	0.0129	0.00427	0.000055083	14.80
	985	0.0123	0.00455	0.000055965	17.60
	875	0.0118	0.00471	5.55309E-05	15.76

Table B2. Tensile strength at yield (σ_{ty}) of composites (pure thermoplastic).

Table B3. Summary of tensile strength at yield (σ_{ty}) of composites.

Composite	Average	Standard Deviation	Coef. Var.	95% Confidence	e Interval (MPa)
Composite	(MPa)	(MPa)	(%)	Upper	Lower
LLDPE + U	15.25	0.43	2.81	15.85	14.66
LLDPE + S	15.80	0.57	3.60	16.59	15.01
LLDPE + B	16.13	0.66	4.07	17.04	15.21
LLDPE + P	15.62	1.62	10.38	17.87	13.37
HDPE + U	16.82	0.86	5.12	18.01	15.62
HDPE + S	17.48	0.80	4.60	18.60	16.37
HDPE + B	16.82	0.57	3.40	17.61	16.03
HDPE + P	16.88	0.33	1.93	17.34	16.43
HDPE/LLDPE + U	16.47	1.14	6.91	18.05	14.89
HDPE/LLDPE + S	16.78	0.73	4.32	17.78	15.77
HDPE/LLDPE + B	16.72	0.63	3.79	17.60	15.84
HDPE/LLDPE + P	17.12	0.56	3.26	17.90	16.35
LLDPE (pure)	15.10	1.57	10.41	17.28	12.92
HDPE (pure)	16.20	1.11	6.84	17.74	14.66

Variable	N	Mean	Std. Deviation	Std. Error		nce Interval for (MPa)	σ_{ty} ((MPa)
		(MPa)	(MPa)	(MPa)	Lower	Upper	Minimum	Maximum
LLDPE + U	5	15.25	0.43	0.19	14.72	15.78	14.60	15.73
LLDPE + S	5	15.80	0.57	0.26	15.09	16.51	15.13	16.4
LLDPE + B	5	16.13	0.66	0.29	15.31	16.94	15.35	16.9
LLDPE + P	5	15.62	1.62	0.73	13.61	17.63	14.49	18.45
LLDPE	5	15.58	1.57	0.70	13.14	17.05	13.04	16.93
Total	25	15.58	1.07	0.21	15.14	16.02	13.04	18.45
HDPE + U	5	16.82	0.86	0.39	15.75	17.89	15.56	17.79
HDPE + S	5	17.49	0.80	0.36	16.49	18.48	16.97	18.9
HDPE + B	5	16.82	0.57	0.26	16.11	17.53	15.97	17.39
HDPE + P	5	16.88	0.33	0.15	16.48	17.29	16.36	17.21
HDPE	5	16.20	1.11	0.49	14.83	17.58	14.80	17.6
Total	25	16.84	0.82	0.16	16.50	17.18	14.80	18.9
HDPE/LLDPE + U	5	16.47	1.14	0.51	15.05	17.89	14.50	17.31
HDPE/LLDPE + S	5	16.78	0.73	0.32	15.88	17.68	15.75	17.72
HDPE/LLDPE + B	5	16.72	0.55	0.25	16.03	17.41	15.88	17.27
HDPE/LLDPE + P	5	17.12	0.56	0.25	16.43	17.82	16.51	17.73
Total	20	16.77	0.76	0.17	16.42	17.13	14.50	17.73

Table B4. Statistical analysis of tensile strength at yield of composites.

One way Analysis of Variance

		df	Sum of	Moon Squara	Computed	Computed
		aı	Squares	Mean Square	F	Sig. (0.05)
LLDPE	Between Groups	4	3.45	0.86	0.71 ^{ns}	0.593
	Within Groups	20	24.17	1.21		
	Total	24	27.61			
HDPE	Between Groups	4	4.12	1.03	1.69 ^{ns}	0.191
	Within Groups	20	12.18	0.61		
	Total	24	16.30			
HDPE/LLDPE	Between Groups	3	1.08	0.36	0.59 ^{ns}	0.63
	Within Groups	16	9.80	0.61		
	Total	19	10.88			
^{ns} = not significan	t, *= significant at 5%	evel, ** = sign	nificant at 1% leve	1.		

Homogeneous Subsets

	Fiber Treatment	N	$\mathbf{P}=0.05$
	Fiber Treatment	IN	Mean (MPa)
LLDPE	Pure	5	15.580 ^a
	U	5	15.252 ^a
	Р	5	15.620 ^a
	S	5	15.798 ^a
	В	5	16.126 ^a

Table B4 continued				
HDPE	Pure	5	16.204 ^a	
	В	5	16.818 ^a	
	U	5	16.818 ^a	
	Р	5	16.882 ^a	
	S	5	17.486 ^b	
HDPE/LLDPE	U	5	16.472 ^a	
	В	5	16.718 ^a	
	S	5	16.776 ^a	
	Р	5	17.124 ^a	

Means for groups in homogeneous subsets are displayed. Uses Harmonic Mean Sample Size = 5.000.

APPENDIX C

Tensile-Impact Strength of Composites

Experimental conditions for the tensile-impact strength of composites presented in Tables C1 and C2.

Temperature (°C): 23 Humidity (%): 50 ASTM procedure: D1822-93 (ASTM 1993) Type L (long) specimen: dumbbell-shaped Machine : tensile impact tester (Tinius Olsen Testing Machines Co., Willow Grove, PA) Operator : bei Units type : SI

Composite	Width (in)	Depth (in) -	Impact	Energy	Tensile-Imp	act Strength
Composite	width (ill)	Deptii (iii) -	(in·lbf)	(N·m)	(ft·lbf/in ²)	(kJ/m ²)
LLDPE+U	0.1020	0.1305	14.00	1.59	87.65	184.15
	0.1050	0.1495	20.00	2.26	106.17	223.07
	0.1335	0.1305	18.50	2.09	88.49	185.92
	0.1160	0.1280	15.00	1.69	84.19	176.88
	0.1025	0.1340	14.00	1.59	84.94	178.46
LLDPE+S	0.0925	0.1345	15.50	1.75	103.82	218.13
	0.0945	0.1545	14.50	1.64	82.76	173.88
	0.0985	0.1420	16.00	1.80	95.33	200.28
	0.0750	0.1390	14.50	1.64	115.91	243.52
	0.0900	0.1465	15.00	1.69	94.80	199.18
LLDPE+B	0.0950	0.1395	15.00	1.69	94.32	198.17
	0.0930	0.1350	10.00	1.13	66.37	139.45
	0.0850	0.1180	14.00	1.59	116.32	244.38
	0.1010	0.1220	20.50	2.32	138.64	291.28
	0.0820	0.1350	14.00	1.59	105.39	221.42
LLDPE+P	0.0965	0.1375	16.00	1.80	100.49	211.12
	0.0945	0.1425	14.00	1.59	86.64	182.02
	0.0975	0.1315	15.00	1.69	97.49	204.84
	0.0970	0.1320	16.00	1.80	104.13	218.79
	0.0990	0.1405	15.00	1.69	89.87	188.81
HDPE+U	0.1170	0.1480	20.00	2.26	96.25	202.22
	0.1150	0.1420	17.00	1.93	86.75	182.27
	0.1170	0.1450	20.50	2.32	100.70	211.57
	0.1110	0.1450	14.00	1.59	72.49	152.29
	0.1035	0.1460	19.00	2.14	104.78	220.14
HDPE+S	0.1035	0.1335	15.00	1.69	90.47	190.07
	0.1035	0.1330	14.00	1.59	84.75	178.07
	0.0950	0.1385	15.00	1.69	95.00	199.60
	0.0965	0.1320	16.00	1.80	104.67	219.92
	0.1110	0.1420	20.50	2.32	108.38	227.71

Table C1. Tensile-impact strength of composites (10% fiber).

Table C1 continued						
HDPE+B	0.1095	0.1390	21.00	2.37	114.98	241.57
	0.1110	0.1415	20.50	2.32	108.77	228.52
	0.1070	0.1400	22.00	2.48	122.39	257.13
	0.1035	0.1455	14.50	1.64	80.24	168.58
	0.1115	0.1385	16.50	1.86	89.04	187.07
HDPE+P	0.1225	0.1400	17.00	1.93	82.60	173.55
	0.1165	0.1290	16.00	1.80	88.72	186.40
	0.1140	0.1255	13.00	1.46	75.72	159.09
	0.1105	0.1220	22.00	2.48	135.99	285.72
	0.1115	0.1310	20.00	2.26	114.10	239.73
HDPE/LLDPE+U	0.1240	0.1120	15.50	1.75	93.01	195.41
	0.1250	0.1175	14.00	1.59	79.43	166.89
	0.1270	0.1115	17.50	1.98	102.99	216.37
	0.1220	0.1195	15.00	1.69	85.74	180.14
	0.1295	0.1130	16.50	1.86	93.96	197.42
HDPE/LLDPE+S	0.1210	0.1100	17.00	1.93	106.44	223.62
	0.1190	0.0975	13.50	1.52	96.96	203.72
	0.1285	0.1005	14.50	1.64	93.57	196.58
	0.1230	0.1045	14.50	1.64	94.01	197.51
	0.1295	0.1185	16.50	1.86	89.60	188.25
HDPE/LLDPE+B	0.1125	0.1180	15.00	1.69	94.16	197.83
	0.1265	0.1190	17.00	1.93	94.11	197.72
	0.1145	0.1135	15.50	1.75	99.39	208.82
	0.1310	0.1195	16.50	1.86	87.83	184.54
	0.1100	0.1215	18.00	2.03	112.23	235.80
HDPE/LLDPE+P	0.1200	0.1215	23.00	2.60	131.46	276.19
	0.1280	0.1220	20.00	2.26	106.73	224.24
	0.1275	0.1225	15.00	1.69	80.03	168.15
	0.1300	0.1205	15.00	1.69	79.80	167.65
	0.1275	0.1185	16.00	1.80	88.25	185.41

The tensile-impact strength is calculated from the following:

Tensile – impact strength $(kJ/m^2) = \frac{Impact energy}{A}$

Where: A =the cross sectional area (in²)

C		Denth (in)	Impact	Energy	Tensile-Impact Strength		
Composite	Width (in)	Depth (in)	(in.·lbf)	(N·m)	(ft·lbf/in. ²)	(kJ/m^2)	
LLDPE	0.1025	0.1399	16.00	1.81	92.98	195.35	
	0.1045	0.1425	17.00	1.93	95.13	199.88	
	0.1075	0.1316	15.00	1.70	88.36	185.64	
	0.1170	0.1320	15.50	1.75	83.64	175.72	
	0.0990	0.1405	15.00	1.70	89.87	188.81	
HDPE	0.1035	0.1355	15.00	1.70	89.13	187.26	
	0.1015	0.1379	13.50	1.52	80.38	168.87	
	0.1050	0.1385	15.00	1.70	85.95	180.59	
	0.1165	0.1320	16.00	1.81	86.70	182.16	
	0.1110	0.1320	15.50	1.75	88.16	185.22	

Table C2. Tensile-impact strength of composites (pure thermoplastic).

Table C3. Summary of tensile-impact strength of composites.

Composite	Average (kJ/m ²)	Standard Deviation	Coef. Var.	95% Confidence Interval (kJ/m ²)		
•		(kJ/m^2)	(%)	Upper	Lower	
LLDPE + U	189.69	19.04	10.04	216.12	163.27	
LLDPE + S	207.00	25.79	12.46	242.80	171.20	
LLDPE + B	218.94	56.21	25.67	296.97	140.92	
LLDPE + P	201.12	15.35	7.63	222.42	179.81	
HDPE + U	193.70	27.09	13.99	231.31	156.09	
HDPE + S	203.07	20.60	10.14	231.67	174.48	
HDPE + B	216.57	37.37	17.25	268.45	164.70	
HDPE + P	208.90	52.65	25.20	281.99	135.81	
HDPE/LLDPE + U	191.24	18.73	9.79	217.25	165.24	
HDPE/LLDPE + S	201.94	13.31	6.59	220.42	183.46	
HDPE/LLDPE + B	204.94	19.28	9.41	231.70	178.18	
HDPE/LLDPE + P	204.33	46.30	22.66	268.60	140.06	
LLDPE (pure)	189.08	9.31	4.92	202.00	176.16	
HDPE (pure)	180.82	7.17	3.96	190.77	170.87	

Variable	N	Mean	Std. Deviation	Std. Error		ence Interval 1 (kJ/m ²)		act Strength /m ²)
		(kJ/m^2)	(kJ/m^2)	(kJ/m^2)	Lower	Upper	Minimum	Maximum
LLDPE + U	5	189.70	19.04	8.51	166.06	213.33	176.88	223.07
LLDPE + S	5	207.00	25.79	11.53	174.97	239.02	173.88	243.52
LLDPE + B	5	218.94	56.21	25.14	149.15	288.73	139.45	291.28
LLDPE + P	5	201.12	15.35	6.86	182.06	220.17	182.02	218.79
LLDPE	5	189.08	9.31	4.16	177.53	200.63	175.72	199.88
Total	25	201.17	29.70	5.94	188.91	213.43	139.45	291.28
HDPE + U	5	193.70	27.09	12.12	160.06	227.34	152.29	220.14
HDPE + S	5	203.07	20.60	9.21	177.50	228.65	178.07	227.71
HDPE + B	5	216.57	37.37	16.71	170.17	262.98	168.58	257.13
HDPE + P	5	208.90	52.65	23.55	143.52	274.27	159.09	285.72
HDPE	5	180.82	7.17	3.21	171.92	189.72	168.87	187.26
Total	25	200.61	32.51	6.50	187.20	214.03	152.29	285.72
HDPE/LLDPE + U	5	191.25	18.73	8.38	167.99	214.50	166.89	216.37
HDPE/LLDPE + S	5	201.94	13.31	5.95	185.41	218.47	188.25	223.62
HDPE/LLDPE + B	5	204.94	19.28	8.62	181.01	228.88	184.54	235.80
HDPE/LLDPE + P	5	204.33	46.29	20.70	146.85	261.81	167.65	276.19
Total	20	200.61	25.94	5.80	188.47	212.75	166.89	276.19

Table C4. Statistical analysis of tensile-impact strength of composites.

		36	Sum of	Maar Caraa	Computed	Computed
		df	Squares	Mean Square	F	Sig. (0.05)
LLDPE	Between Groups	4	3137.81	784.45	0.87 ^{ns}	0.50
	Within Groups	20	18035.41	901.77		
	Total	24	21173.22			
HDPE	Between Groups	4	3845.16	961.29	0.89 ^{ns}	0.49
HDPE	Within Groups	20	21513.93	1075.70		
	Total	24	25359.08			
HDPE/LLDPE	Between Groups	3	610.16	203.39	0.27 ^{ns}	0.85
	Within Groups	16	12171.08	760.69		
	Total	19	12781.24			

APPENDIX D

Durometer Hardness of Composites

Experimental conditions for the Durometer hardness of composites presented in Table D1.

Temperature (°C): 23 Humidity (%): 50 ASTM procedure: D2240-97 (ASTM 1998) Types of durometer : Type D gauge Machine : Durometer hardness tester (Shore Instrument and MFG Co., Freeport, NY) Operator : bei Units type : SI

Composite -				H	lardness	Test (SE)			
Composite	1	2	3	4	5	6	7	8	9	10
LLDPE+U	15	19	14	14	13	19	16	17	15	18
LLDPE+S	9	19	15	20	24	20	11	20	15	17
LLDPE+B	12	11	15	24	16	20	19	14	18	19
LLDPE+P	16	15	19	18	17	21	20	21	20	19
HDPE+U	19	17	16	19	15	14	18	17	20	15
HDPE+S	15	9	24	19	16	10	17	18	15	19
HDPE+B	19	16	15	20	17	19	15	13	18	10
HDPE+P	14	21	14	15	14	15	19	20	19	20
HDPE/LLDPE+U	14	16	13	15	20	17	19	20	14	15
HDPE/LLDPE+S	20	14	15	15	19	14	16	20	20	17
HDPE/LLDPE+B	16	19	15	16	18	16	19	17	17	14
HDPE/LLDPE+P	15	21	16	15	7	17	12	20	24	15
LLDPE	19	18	21	15	16	16	20	18	16	20
HDPE	15	22	17	19	21	16	17	17	18	15

 Table D1. Durometer hardness of composites (10% fiber and pure thermoplastic).

Composite	Average	Standard Deviation	Coef. Var.	95% Confidence Interval (SD)		
-	(SD)	(SD)	(%)	Upper	Lower	
LLDPE + U	16.0	2.0	11.8	18.5	15.5	
LLDPE + S	17.0	4.4	27.1	19.5	12.9	
LLDPE + B	16.8	3.1	19.0	18.5	13.9	
LLDPE + P	18.6	2.9	17.1	19.3	14.9	
HDPE + U	17.0	2.6	15.9	18.3	14.4	
HDPE + S	16.2	2.5	14.9	18.9	15.1	
HDPE + B	16.2	1.6	9.8	17.9	15.5	
HDPE + P	17.1	4.8	29.5	19.8	12.6	
HDPE/LLDPE + U	16.3	2.2	13.5	17.6	14.4	
HDPE/LLDPE + S	17.0	4.6	26.9	20.5	13.6	
HDPE/LLDPE + B	16.7	4.0	23.6	19.8	13.8	
HDPE/LLDPE + P	16.2	2.1	11.1	20.2	17.0	
LLDPE (pure)	17.9	2.1	11.6	19.5	16.3	
HDPE (pure)	17.7	2.4	13.3	19.5	15.9	

Table D2. Summary of Durometer hardness of composites.

Variable	N	Mean	Std. Deviation	Std. Error		ence Interval an (SD)	Durometer H	ardness (SD)
		(SD)	(SD)	(SD)	Lower	Upper	Minimum	Maximum
LLDPE + U	10	16.00	2.16	0.68	14.45	17.55	13	19
LLDPE + S	10	17.00	4.57	1.45	13.73	20.27	9	24
LLDPE + B	10	16.80	3.97	1.25	13.96	19.64	11	24
LLDPE + P	10	18.60	2.07	0.65	17.12	20.08	15	21
LLDPE	10	17.90	2.08	0.66	16.41	19.39	15	21
Total	50	17.26	3.16	0.45	16.36	18.16	9	24
HDPE + U	10	17.00	2.00	0.63	15.57	18.43	14	20
HDPE + S	10	16.20	4.39	1.39	13.06	19.34	9	24
HDPE + B	10	16.20	3.08	0.98	13.99	18.41	10	20
HDPE + P	10	17.10	2.92	0.92	15.01	19.19	14	21
HDPE	10	17.70	2.36	0.75	16.01	19.39	15	22
Total	50	16.84	2.99	0.42	15.99	17.69	9	24
HDPE/LLDPE + U	10	16.30	2.58	0.82	14.45	18.15	13	20
HDPE/LLDPE + S	10	17.00	2.54	0.80	15.18	18.82	14	20
HDPE/LLDPE + B	10	16.70	1.64	0.52	15.53	17.87	14	19
HDPE/LLDPE + P	10	16.20	4.78	1.51	12.78	19.62	7	24
Total	40	16.55	3.00	0.47	15.59	17.51	7	24

Table D3. Statistical analysis of Durometer hardness of composites.

		16	6	Maria	Computed	Computed	
		df	Sum of Squares	Mean Square	F	Sig. (0.05)	
LLDPE	Between Groups	4	40.72	10.18	1.02 ^{ns}	0.41	
	Within Groups	45	448.90	9.98			
	Total	49	489.62				
HDPE	Between Groups	4	16.52	4.13	0.44 ^{ns}	0.78	
	Within Groups	45	422.20	9.38			
	Total	49	438.72				
HDPE/LLDPE	Between Groups	3	4.10	1.37	0.14 ^{ns}	0.93	
	Within Groups	36	347.80	9.66			
	Total	39	351.90				

APPENDIX E

Moisture Absorption of Flax Fibers

Experimental conditions for the moisture absorption of flax fibers presented in Table E1.

Temperature (°C): 23 Humidity (%): 33, 66, 100 Duration: 72h Machine : Environmental test chamber (Anglelantoni, ACS, Massa Martana, Italy) Analytical balance Operator : bei Units type : SI

No.	Fiber	33% RH (Dry bulb temp:2	3°C, Wet bulb tem	p:13.33°C)	- Mass Increase
of Rep.	Treatment	Before Conditioning (g)	Feb 14, 03, 1:50 pm			(%)
	Untreated	24.242	25.237	25.462	25.477	5.09
1	Silane	24.715	25.710	25.951	25.954	5.01
1	Benzoylation	24.926	25.828	25.923	25.992	4.28
	Peroxide	24.627	25.029	25.320	25.523	3.64
	Fiber	33% RH (Dry bulb temp:2	3°C, Wet bulb tem	p:13.33°C)	- Mass Increase
	Treatment	Before Conditioning (g)	Feb 27, 03, 10:50 am	Feb 28, 03, 11:00 am	Mar 01, 03, 10:55 am	- Mass Increase (%)
2	Untreated	24.584	25.698	25.770	25.952	5.56
	Silane	24.691	25.251	25.598	25.865	4.75
	Benzoylation	24.712	24.972	25.002	25.711	4.04
	Peroxide	24.772	25.499	25.695	25.759	3.98
	1211	33% RH (p:13.33°C)	N# 1		
	Fiber - Treatment	Before Conditioning (g)	Mar 11, 03, 10:00 am	Mar 12, 03, 11:10 am	Mar 13, 03, 10:45 am	- Mass Increase (%)
3	Untreated	24.806	25.026	25.352	26.129	5.33
	Silane	24.775	25.041	25.477	25.985	4.88
	Benzoylation	24.612	24.759	25.009	25.637	4.16
	Peroxide	24.872	25.013	25.400	25.820	3.81
	5.11	66% RH	(Dry bulb temp:2	23°C, Wet bulb tem	ıр:18.6°С)	
	Fiber Treatment	Before Conditioning (g)	Feb 21, 03, 11:10 am	Feb 22, 03, 10:40 am	Feb 23, 03, 7:30 pm	- Mass Increase (%)
1	Untreated	24.269	25.922	26.350	26.585	9.54
	Silane	24.739	25.270	26.170	27.029	9.26
	Benzoylation	24.940	25.229	26.232	27.067	8.53
	Peroxide	24.256	26.640	26.642	26.443	9.02
	F ¹	66% RH	(Dry bulb temp:2	23°C, Wet bulb tem	р:18.6°С)	Manufation
2	Fiber Treatment	Before Conditioning (g)	Mar 03, 03, 10:10 am	Mar 04, 03, 11:50 am	Mar 05, 03, 12:00 pm	– Mass Increase (%)

Table E1. Moisture absorption of flax fibers.

Table	e E1 continued					
	Untreated	24.566	25.998	26.452	27.075	10.21
2	Silane	24.852	25.862	26.301	27.085	8.99
Z	Benzoylation	24.760	25.968	26.635	26.902	8.65
	Peroxide	24.699	26.001	26.052	26.861	8.75
	Fiber	66% RH	(Dry bulb temp:	23°C, Wet bulb tem	ıp:18.6°C)	– Mass Increase
	Treatment	Before Conditioning (g)	Mar 16, 03, 1:00 pm	Mar 17, 03, 1:50 pm	Mar 18, 03, 12:30 pm	(%)
3	Untreated	24.953	25.798	26.523	27.418	9.88
	Silane	24.641	25.995	26.465	26.891	9.13
	Benzoylation	24.882	25.887	26.265	27.020	8.59
	Peroxide	24.523	25.563	26.009	26.703	8.89
	5.1	100% RH	I (Dry bulb temp	:23°C, Wet bulb te	mp:23°C)	
	Fiber Treatment	Before Conditioning (g)	Feb 24, 03, 10:40 am	Feb 25, 03, 10:00 am	Feb 26, 03, 6:40 pm	– Mass Increase (%)
1	Untreated	24.177	27.365	28.300	28.447	17.66
	Silane	24.691	27.332	28.500	28.745	16.42
	Benzoylation	24.892	27.017	28.171	28.545	14.68
	Peroxide	24.143	27.407	27.496	27.850	15.35
		100% RH	I (Dry bulb temp	:23°C, Wet bulb te	mp:23°C)	
	Fiber Treatment	Before Conditioning (g)	Mar 07, 03, 11:15 am	Mar 08, 03, 1:00 pm	Mar 09, 03, 12:30 pm	– Mass Increase (%)
2	Untreated	24.875	27.589	28.542	29.368	18.06
	Silane	24.561	27.458	28.289	28.451	15.84
	Benzoylation	24.922	27.550	28.320	28.491	14.32
	Peroxide	24.530	27.693	27.879	28.227	15.07
	D *1	100% RH	I (Dry bulb temp	:23°C, Wet bulb te	mp:23°C)	
	Fiber Treatment	Before Conditioning (g)	Mar 20, 03, 11:25 am	Mar 21, 03, 1:05 pm	Mar 22, 03, 12:55 pm	– Mass Increase (%)
3	Untreated	24.757	27.854	28.411	29.178	17.86
	Silane	24.898	27.310	28.099	28.913	16.13
	Benzoylation	24.694	27.420	28.215	28.274	14.50
	Peroxide	24.901	27.369	27.799	28.688	15.21
RH: r	elative humidity					

The moisture absorption is calculated according to the following equation:

Increase in mass (percent) =
$$\left(\frac{M_t - M_0}{M_0}\right) \times 100$$

Where: M_t = mass of the sample after conditioning (g) (wet weight) M_0 = mass of the sample before conditioning (g) (dry weight)

RH	Fiber	Mean	Standard	Coef. Var.	95% Confiden	ce Interval (%)
КП	Treatment	(%)	Deviation (%)	(%)	Upper	Lower
	Untreated	5.33	0.24	4.41	6.04	4.61
33%	Silane	4.88	0.13	2.66	5.28	4.48
5570	Benzoylation	4.16	0.12	2.88	4.53	3.79
	Peroxide	3.81	0.17	4.46	4.33	3.29
	Untreated	9.88	0.33	3.39	10.89	8.86
66%	Silane	9.13	0.14	1.48	9.54	8.71
0070	Benzoylation	8.59	0.06	0.7	8.77	8.41
	Peroxide	8.89	0.13	1.52	9.3	8.47
	Untreated	17.86	0.2	1.12	18.47	17.25
100%	Silane	16.13	0.29	1.8	17.01	15.25
10070	Benzoylation	14.5	0.18	1.24	15.05	13.95
	Peroxide	15.21	0.14	0.92	15.64	14.78

Table E2. Summary of moisture absorption (mass increase, %) of flax fibers.

RH	Variable	N	Mean (%)	Std. Deviatio	Std. Error		ence Interval an (%)	Mass Increase (%)	
				n (%)	(%)	Lower	Upper	Min.	Max.
	Untreated	3	5.33	0.24	0.14	4.74	5.91	5.09	5.56
220/	Silane	3	4.88	0.13	0.08	4.56	5.20	4.75	5.01
33%	Benzoylation	3	4.16	0.12	0.07	3.86	4.46	4.04	4.28
	Peroxide	3	3.81	0.17	0.10	3.39	4.23	3.64	3.98
	Total	12	4.54	0.64	0.18	4.14	4.95	3.64	5.56
	Untreated	3	9.88	0.34	0.19	9.04	10.71	9.54	10.21
	Silane	3	9.13	0.14	0.08	8.79	9.46	8.99	9.26
66%	Benzoylation	3	8.59	0.06	0.03	8.44	8.74	8.53	8.65
	Peroxide	3	8.89	0.14	0.08	8.55	9.22	8.75	9.02
	Total	12	9.12	0.52	0.15	8.79	9.45	8.53	10.21
	Untreated	3	17.86	0.20	0.12	17.36	18.36	17.66	18.06
	Silane	3	16.13	0.29	0.17	15.41	16.85	15.84	16.42
100%	Benzoylation	3	14.50	0.18	0.10	14.05	14.95	14.32	14.68
	Peroxide	3	15.21	0.14	0.08	14.86	15.56	15.07	15.35
	Total	12	15.93	1.33	0.38	15.08	16.77	14.32	18.06

Table E3. Statistical analysis of moisture absorption (mass increase, %) of flax fibers.

RH		36	Sum of	Maan Samaaa	Computed	Computed
		df	Squares	Mean Square	F	Sig. (0.05)
33%	Between Groups	3	4.24	1.41	48.92**	0.0000
	Within Groups	8	0.23	0.03		
	Total	11	4.47			
66%	Between Groups	3	2.72	0.91	23.85**	0.0002
	Within Groups	8	0.30	0.04		
	Total	11	3.03			
100%	Between Groups	3	18.98	6.33	143.74**	0.0000
	Within Groups	8	0.35	0.04		
	Total	11	19.34			

 ns = not significant, *= significant at 5% level, ** = significant at 1% level.

Multiple Comparisons

Dependent	Mean (I) (J) Difference		Mean Difference	Std.	Computed		onfidence for Mean	
Variable				(I-J)	Error	Sig. (0.05)	Lower	Upper
33%RH	LSD	Untreated	Silane	0.45 (*)	0.14	0.012^{*}	0.13	0.77
			Benzoylation	1.17 (*)	0.14	0.000^{**}	0.85	1.49
			Peroxide	1.52 (*)	0.14	0.000^{**}	1.20	1.84
		Silane	Benzoylation	0.72 (*)	0.14	0.001^{**}	0.40	1.04
			Peroxide	1.07 (*)	0.14	0.000^{**}	0.75	1.39
		Benzoylation	Peroxide	0.35 (*)	0.14	0.036*	0.03	0.67
66%	LSD	Untreated	Silane	0.75 (*)	0.16	0.002^{**}	0.38	1.12

able E3 co	ontinued							
			Benzoylation	1.29 (*)	0.16	0.000^{**}	0.92	1.65
			Peroxide	0.99 (*)	0.16	0.000^{**}	0.62	1.36
		Silane	Benzoylation	0.54 (*)	0.16	0.010^{**}	0.17	0.90
			Peroxide	0.24	0.16	0.170 ^{ns}	-0.13	0.61
		Benzoylation	Peroxide	-0.30	0.16	0.100 ^{ns}	-0.66	0.07
100%	LSD	Untreated	Silane	1.73 (*)	0.17	0.000^{**}	1.33	2.13
			Benzoylation	3.36 (*)	0.17	0.000^{**}	2.96	3.76
			Peroxide	2.65 (*)	0.17	0.000^{**}	2.25	3.05
		Silane	Benzoylation	1.63 (*)	0.17	0.000^{**}	1.23	2.03
			Peroxide	0.92 (*)	0.17	0.001**	0.52	1.32
		Benzoylation	Peroxide	-0.71 (*)	0.17	0.003**	-1.11	-0.31

*. The mean difference is significant at the .05 level.

Homogeneous Subsets

ын	E:h T	N	P = 0.05	
RH	Fiber Treatment	N	Mean (%)	
	Peroxide	3	3.81 ^a	
220/	Benzoylation	3	4.16 ^b	
33%	Silane	3	4.88 ^c	
	Untreated	3	5.23 ^d	
(())	Benzoylation	3	8.59 ^a	
	Peroxide	3	8.89 ^{ab}	
66%	Silane	3	9.13 ^b	
	Untreated	3	9.88 ^c	
	Benzoylation	3	14.50 ^a	
1000/	Peroxide	3	15.21 ^b	
100%	Silane	3	16.13 ^c	
	Untreated	3	17.86 ^d	

Uses Harmonic Mean Sample Size = 3.000.

APPENDIX F

Water Absorption of Composites

Experimental conditions for the water absorption of composites presented in Table F1.

Temperature (°C): 23 Humidity (%): 50 Duration: 24h Machine : Desiccator, oven, Types of sample : Rectangular specimen (25.4 mm x 76.2 mm) ASTM procedure: D570-99 (ASTM, 1999) Operator : bei Units type : SI

No.	Comparito	Mas	s (g)	Water absorption	
of Rep.	Composite	Before Conditioning	After Conditioning	g (%)	
	LLDPE + U	6.4561	6.4620	0.09	
	LLDPE + S	6.9015	6.9075	0.09	
	LLDPE + B	7.3397	7.3453	0.08	
	LLDPE + P	8.1876	8.1943	0.08	
	HDPE + U	6.3861	6.3931	0.11	
	HDPE + S	6.1757	6.1820	0.10	
1	HDPE + B	7.8593	7.8650	0.07	
1	HDPE + P	7.8488	7.8541	0.07	
	HDPE/LLDPE + U	5.8606	5.8676	0.12	
	HDPE/LLDPE + S	6.5316	6.5383	0.10	
	HDPE/LLDPE + B	4.5737	4.5771	0.07	
	HDPE/LLDPE + P	6.9035	6.9075	0.06	
	LLDPE (pure)	7.5236	7.5260	0.03	
	HDPE (pure)	6.9876	6.9910	0.05	
	LLDPE + U	8.2918	8.2989	0.09	
	LLDPE + S	8.8344	8.8389	0.05	
	LLDPE + B	9.8284	9.8336	0.05	
	LLDPE + P	7.6031	7.6061	0.04	
	HDPE + U	6.6867	6.6929	0.09	
	HDPE + S	9.3302	9.3363	0.07	
2	HDPE + B	8.5258	8.5299	0.05	
2	HDPE + P	6.8826	6.8865	0.06	
	HDPE/LLDPE + U	7.9564	7.9650	0.11	
	HDPE/LLDPE + S	8.5545	8.5611	0.08	
	HDPE/LLDPE + B	8.1846	8.1892	0.06	
	HDPE/LLDPE + P	6.3480	6.3510	0.05	
	LLDPE (pure)	7.6658	7.6690	0.04	
	HDPE (pure)	7.0523	7.0563	0.06	

Table F1. Water absorption of composites (10% fiber and pure thermoplastic).

Table I	F1 continued			
	LLDPE + U	6.7799	6.7854	0.08
	LLDPE + S	7.7682	7.7730	0.06
	LLDPE + B	8.9982	9.0030	0.05
	LLDPE + P	7.6132	7.6180	0.06
	HDPE + U	6.9272	6.9332	0.09
	HDPE + S	4.0384	4.0418	0.08
2	HDPE + B	4.7376	4.7411	0.07
3	HDPE + P	3.9028	3.9056	0.07
	HDPE/LLDPE + U	6.9744	6.9810	0.09
	HDPE/LLDPE + S	8.0598	8.0680	0.10
	HDPE/LLDPE + B	8.1674	8.1750	0.09
	HDPE/LLDPE + P	6.5664	6.5710	0.07
	LLDPE (pure)	7.2543	7.2569	0.04
	HDPE (pure)	7.2153	7.2183	0.04

The percentage increase in weight during immersion was calculated according to the nearest 0.01% as follows:

Increase in mass (percent) =
$$\frac{\text{Conditioned mass} - \text{Dry mass}}{\text{Dry mass}} \times 100$$

Table F2. Summary of water absorption (mass increase, %) of composites.

Composito	Mean	Standard	Coef. Var.	95% Confiden	ce Interval (%)
Composite	(%)	Deviation (%)	(%)	Upper	Lower
LLDPE + U	0.09	0.005	5.98	0.10	0.07
LLDPE + S	0.07	0.018	27.75	0.12	0.01
LLDPE + B	0.06	0.013	21.99	0.10	0.02
LLDPE + P	0.06	0.021	34.55	0.13	0.00
HDPE + U	0.10	0.012	12.37	0.13	0.06
HDPE + S	0.08	0.018	21.84	0.14	0.03
HDPE + B	0.06	0.015	22.39	0.11	0.02
HDPE + P	0.07	0.008	11.91	0.09	0.04
HDPE/LLDPE + U	0.11	0.012	11.57	0.15	0.07
HDPE/LLDPE + S	0.09	0.014	15.39	0.14	0.05
HDPE/LLDPE + B	0.07	0.018	24.72	0.13	0.02
HDPE/LLDPE + P	0.06	0.011	19.52	0.09	0.02
LLDPE (pure)	0.04	0.005	13.58	0.05	0.02
HDPE (pure)	0.05	0.008	15.47	0.07	0.03

Variable	N	Mean	Std. Deviation	Std. Error		nce Interval for n (%)	Mass Increase (%)	
		(%)	(%)	(%)	Lower	Upper	Minimum	Maximum
LLDPE + U	3	16.00	0.01	0.00	0.07	0.10	0.08	0.09
LLDPE + S	3	17.00	0.02	0.01	0.01	0.12	0.05	0.09
LLDPE + B	3	16.80	0.02	0.01	0.02	0.10	0.05	0.08
LLDPE + P	3	18.60	0.02	0.01	0.01	0.11	0.04	0.08
LLDPE	3	17.90	0.01	0.00	0.02	0.05	0.03	0.04
Total	15	17.26	0.02	0.01	0.05	0.07	0.03	0.09
HDPE + U	3	17.00	0.01	0.01	0.07	0.13	0.09	0.11
HDPE + S	3	16.20	0.02	0.01	0.05	0.12	0.07	0.10
HDPE + B	3	16.20	0.01	0.01	0.03	0.09	0.05	0.07
HDPE + P	3	17.10	0.01	0.00	0.05	0.08	0.06	0.07
HDPE	3	17.70	0.01	0.01	0.03	0.07	0.04	0.06
Total	15	16.84	0.02	0.00	0.06	0.08	0.04	0.11
HDPE/LLDPE + U	3	16.30	0.02	0.01	0.07	0.14	0.09	0.12
HDPE/LLDPE + S	3	17.00	0.01	0.01	0.06	0.12	0.08	0.10
HDPE/LLDPE + B	3	16.70	0.02	0.01	0.04	0.11	0.06	0.09
HDPE/LLDPE + P	3	16.20	0.01	0.01	0.04	0.08	0.05	0.07
Total	12	16.55	0.02	0.01	0.07	0.10	0.05	0.12

Table F3. Statistical analysis of water absorption (mass increase, %) of composites.

		df	Sum of Squanag	Maan Squara	Computed	Computed
		ai	Sum of Squares	Mean Square	F	Sig. (0.05)
LLDPE	Between Groups	4	0.0038	0.0010	4.0000^{*}	0.0343
	Within Groups	10	0.0024	0.0002		
	Total	14	0.0062			
HDPE	Between Groups	4	0.0040	0.0010	7.8421**	0.0040
	Within Groups	10	0.0013	0.0001		
	Total	14	0.0052			
HDPE/LLDPE	Between Groups	3	0.0039	0.0013	7.3651*	0.0109
	Within Groups	8	0.0014	0.0002		
	Total	11	0.0053			
^s = not significan	t, *= significant at 5%	level, ** =	significant at 1% level.			

Multiple Comparisons

Dependent		(I) (J)		Mean Difference	Difference Std.		95% Confiden Me	
Variable				(I-J)	Error	Sig. (0.05)	Lower	Upper
LLDPE	LSD	U	S	0.020	0.013	0.145 ^{ns}	-0.008	0.048
			В	0.027	0.013	0.061 ^{ns}	-0.002	0.055
			Р	0.027	0.013	0.061 ^{ns}	-0.002	0.055
			Pure	0.050 (*)	0.013	0.003**	0.022	0.078

Table F3 continu	ied							
		S	В	0.007	0.013	0.610 ^{ns}	-0.022	0.035
			Р	0.007	0.013	0.610 ^{ns}	-0.022	0.035
			Pure	0.030 (*)	0.013	0.039^{*}	0.002	0.058
		В	Р	0.000	0.013	1.000 ^{ns}	-0.028	0.028
			Pure	0.023	0.013	0.095 ^{ns}	-0.005	0.052
		Р	Pure	0.023	0.013	0.095 ^{ns}	-0.005	0.052
HDPE	LSD	U	S	0.013	0.009	0.177 ^{ns}	-0.007	0.034
			В	0.033 (*)	0.009	0.005^{**}	0.013	0.054
			Р	0.030 (*)	0.009	0.009^{**}	0.010	0.050
			Pure	0.047 (*)	0.009	0.000^{**}	0.026	0.067
		S	В	0.020	0.009	0.055 ^{ns}	0.000	0.040
			Р	0.017	0.009	0.100 ^{ns}	-0.004	0.037
			Pure	0.033 (*)	0.009	0.005^{**}	0.013	0.054
		В	Р	-0.003	0.009	0.724 ^{ns}	-0.024	0.017
			Pure	0.013	0.009	0.177 ^{ns}	-0.007	0.034
		Р	Pure	0.017	0.009	0.100 ^{ns}	-0.004	0.037
HDPE/LLDPE	LSD	U	S	0.013	0.011	0.252 ^{ns}	-0.012	0.038
			В	0.033 (*)	0.011	0.015^{*}	0.008	0.058
			Р	0.047 (*)	0.011	0.003**	0.022	0.072
		S	В	0.020	0.011	0.101 ^{ns}	-0.005	0.045
			Р	0.033 (*)	0.011	0.015^{*}	0.008	0.058
		В	Р	0.013	0.011	0.252 ^{ns}	-0.012	0.038

^{*}. The mean difference is significant at the .05 level.

Iomogeneous Subsets							
	Fiber Treatment	N	P = 0.05				
	Fiber Treatment	1	Mean (%)				
LLDPE	Pure	3	0.037^{a}				
	В	3	0.060^{ab}				
	Р	3	0.060 ^{ab}				
	S	3	0.067^{ab}				
	U	3	0.087^{b}				
HDPE	Pure	3	0.050^{a}				
	В	3	0.063 ^{ab}				
	Р	3	0.067^{ab}				
	S	3	0.083 ^{bc}				
	U	3	0.097 ^c				
HDPE/LLDPE	Р	3	0.060^{a}				
	В	3	0.073 ^{ab}				
	S	3	0.093 ^{bc}				
	U	3	0.107^{c}				

Means for groups in homogeneous subsets are displayed. Uses Harmonic Mean Sample Size = 3.000.

APPENDIX G

Thermal Property by DSC

Experimental conditions for all the DSC thermal property presented in Table G1.

Temperature (°C): 23 Humidity (%): 50 Heating rate: 10 °C/min Machine: Perkin- Elmer DSC system (TA instruments, New Castle, DE) Types of sample : 6 to 10 mg Temperature range: -50-400 °C Operator : bei Units type : SI

Table G1. Melting point of composites (10% fiber and pure thermoplastic).

Variable	Mass (mg)	Melting Point (°C)
LLDPE + U	6.3	130.6
LLDPE + S	6.5	130.3
LLDPE + B	6.9	130.2
LLDPE + P	7.3	129.2
LLDPE (pure)	10.0	129.9
HDPE + U	8.7	133.4
HDPE + S	8.4	133.1
HDPE + B	7.7	132.9
HDPE + P	6.4	132.7
HDPE (pure)	7.5	130.7
HDPE/LLDPE + U	6.7	130.0
HDPE/LLDPE + S	6.9	131.6
HDPE/LLDPE + B	8.2	130.4
HDPE/LLDPE + P	8.2	130.2

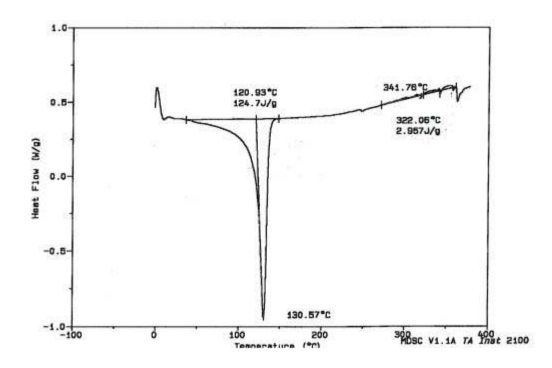
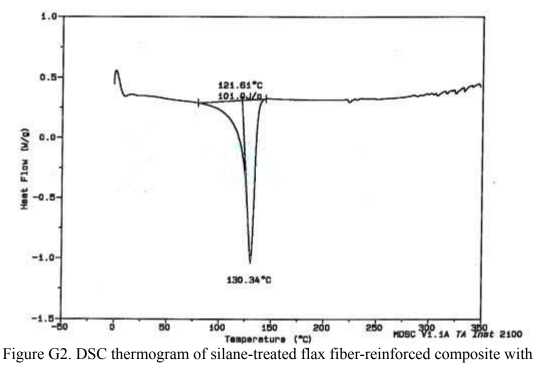
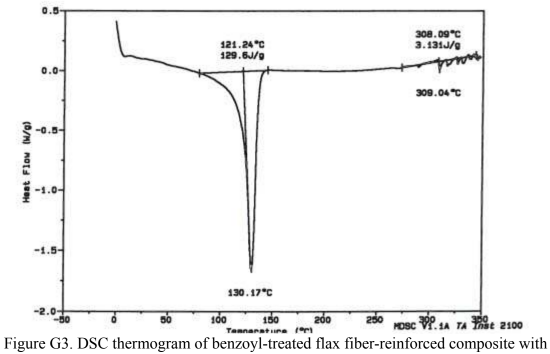


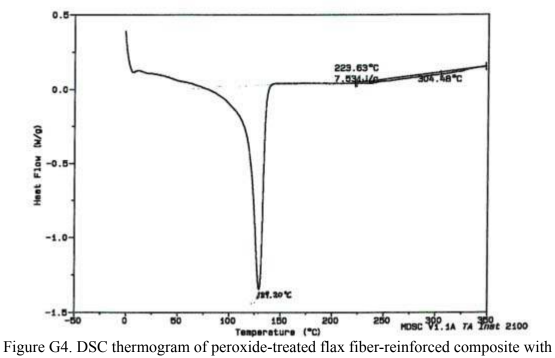
Figure G1. DSC thermogram of untreated flax fiber-reinforced composite with LLDPE.



LLDPE.



LLDPE.



LLDPE.

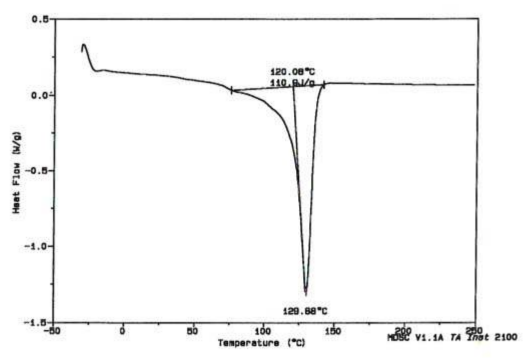


Figure G5. DSC thermogram of pure LLDPE composite.

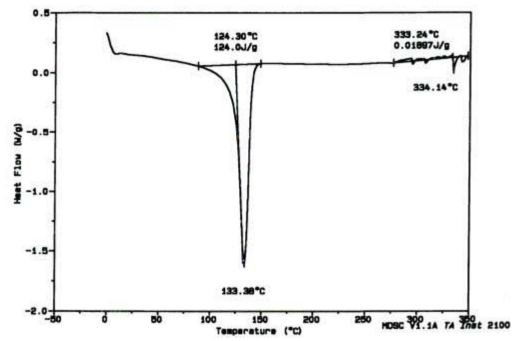


Figure G6. DSC thermogram of untreated flax fiber-reinforced composite with HDPE.

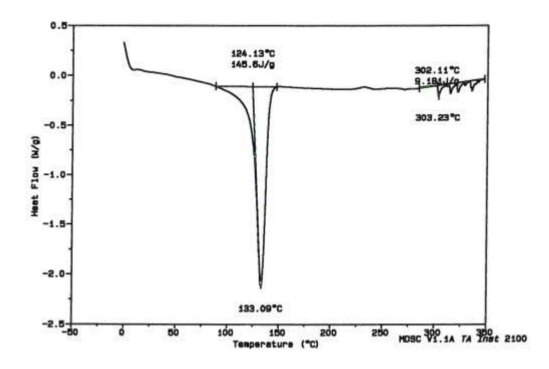


Figure G7. DSC thermogram of silane-treated flax fiber-reinforced composite with HDPE.

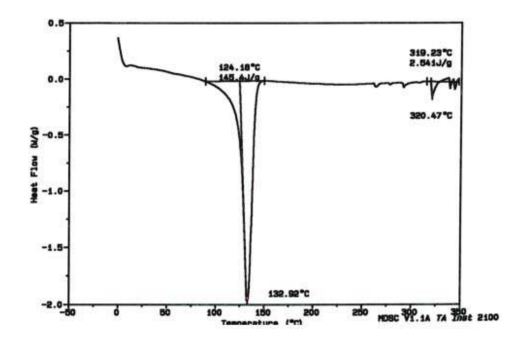


Figure G8. DSC thermogram of benzoyl-treated flax fiber-reinforced composite with HDPE.

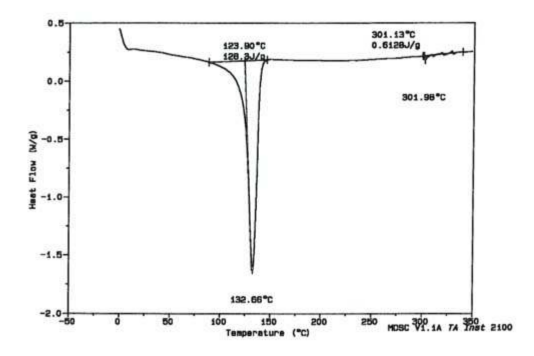


Figure G9. DSC thermogram of peroxide-treated flax fiber-reinforced composite with HDPE.

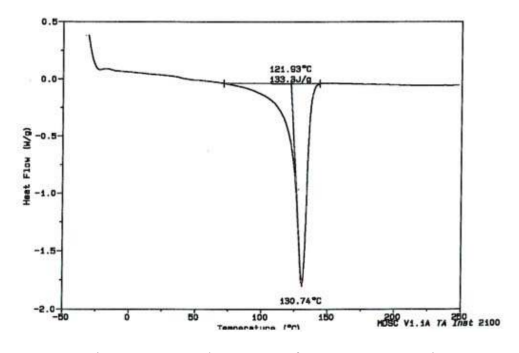


Figure G10. DSC thermogram of pure HDPE composite.

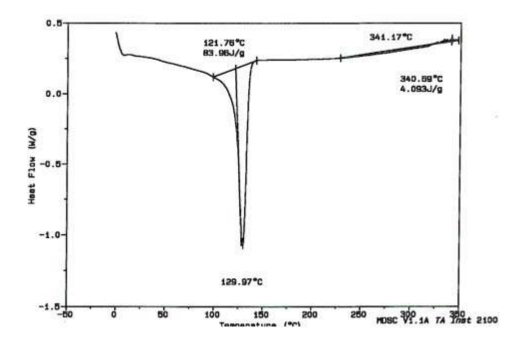


Figure G11. DSC thermogram of untreated flax fiber-reinforced composite with HDPE/LLDPE.

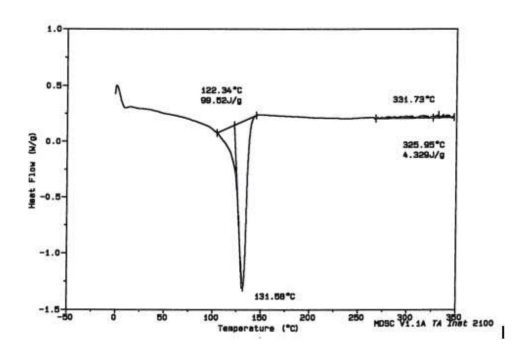


Figure G12. DSC thermogram of silane-treated flax fiber-reinforced composite with HDPE/LLDPE.

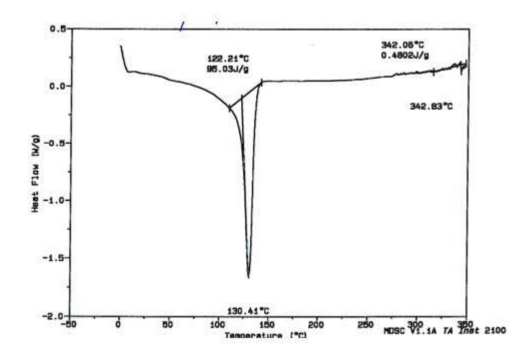


Figure G13. DSC thermogram of benzoyl-treated flax fiber-reinforced composite with HDPE/LLDPE.

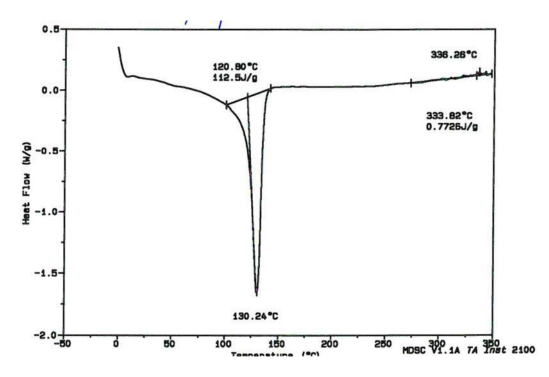
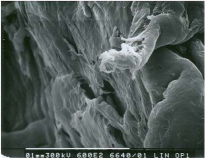


Figure G14. DSC thermogram of peroxide-treated flax fiber-reinforced composite with HDPE/LLDPE.

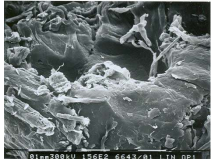
APPENDIX H

The SEM Micrographs of the Cross-session of HDPE and LLDPE/HDPE Flax

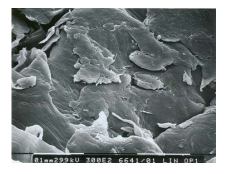
Fiber-reinforced Composites

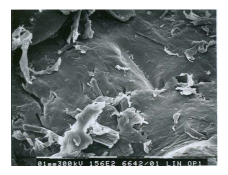


a) Untreated flax in composites



b) Silane treated flax in composites





c) Benzoylation treated flax in composites d) Peroxide treated flax in composites

Figure H1. The SEM micrographs of HDPE with 10% treated flax in composites.





a) Untreated flax in composites

b) Silane treated flax in composites



c) Benzoylation treated flax in composites d) Peroxide treated flax in composites

Figure H2. The SEM micrographs of HDPE/LLDPE with 10% treated flax in composites.