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Synthesis, Optimization and Characterization of Bioplastic from Cornstarch Reinforced with Sugarcane Bagasse Cellulose Fiber

Gadissa Mosisa Gobana

Department of Chemical Engineering, Wollega University Shambu Campus, Oromia, Ethiopia Email: mgadisa25@gmail.com

Abstract

Plastics are made into an array of important products offering significant consumer benefits, they can also litter the environment and harm ecosystems; notably, plastic waste constitutes the majority of ocean debris. There is an increasing need for biodegradable plastics because they are environmentally friendly and can replace petroleum-based non-degradable plastics which pollute the environment. Petroleum based polymers are not sustainable as they depend on depleting fossils. This work investigated the effect of bagasse (sugarcane byproduct) fiber weight fraction on starch based composite. This study also explored the potentials of sugar cane cellulose fiber and starch for bioplastic productions. The effect of oven drying and starch-cellulose fiber ratio was also studied. The experimental design was employed using design expert 7.0 with two factor three level central composite designs (CCD) including five replicates at the center point of optimization study requiring 13 experiments on the 3 responses were analyzed namely tensile strength, water absorption and elongation at break. Results have shown that the addition of cellulose fiber (5 wt%, 10 wt% and 15 wt%) to selected gram of corn starch(5g), the value obtained were for tensile strength (13.27MPa, 24.45MPa and 22.18MPa, respectively), water absorption (36.90, 23.6 and 24.01% respectively) and elongation at break (20.25, 5.268 and 9.58% respectively), these values are the averages of replication treatment. Starch-derived bioplastic reinforced with cellulose fiber at the optimal point of the responses namely tensile strength, water absorption and elongation at break, which are biodegradable, have been prepared and characterized for FTIR, compound microscope, Transparency, solubility, and density. From the analysis of experimental results the maximum and minimum value of tensile strength (26.81 and 11.55MPa), water absorption (39.02% and 20.45%) and elongation at break (25.99% and 4.32%) was obtained respectively. Results obtained under optimal condition were found that Transparency of reinforced bioplastic reduced by 5% with respect to the control. Also its solubility decreased from (14.78% to 10.17%) while density was increased from (1.059g/ml to1.069g/ml). Evidence of the existence of strong interactions between the starch matrix and the cellulose fibers was revealed from detailed Fourier Transform Infra-red and compound microscopic evaluation.

Key words: 1. Cellulose fiber 2. Starch 3. Bioplastic 4. Glycerol 5. Strength

1 Introduction

1.1 Background

The word polymer is derived from the Greek, polyand merosmeaning manyand parts respectively. Plastics are polymers or macromolecules which have been modified with additives and mechanically strong plastic structure that adopt a dimensionally stable form. Plastics have become indispensable to modern living and no longer represent only luxury and novelty as they did in the nineteenth century. Almost 35 million tonnes of synthetic polymers are produced annually in the USA and growth is expected as long as petroleum and other feedstocks last and consumer demand continues (Mo,X.Z. 2010).

The raw material used in the production of renewable biopolymers are continually replenished which becomes helpful over traditional plastics for sustainability. Typical commercial biopolymers are polylactic acid (PLA), polyhydroxyalkanoates (PHA), starches, and bagasse. Some applications for PHAs range from stiff packaging to flexible coatings. PLA can be used for flexible packaging or drinking cups as well as similar applications to polypropylene (PP) and polystyrene (PS) (Cao et al., 2006). Applications for starches reside mainly with food packaging. The degradation period of synthetic plastics may last up to 450 years in a landfill whereas some biodegradable polymers, starch and bagasse composites, may degrade within a few months (Janssen et al., 2009).

The solvent casting process of thermoplastic solution is the most utilized technologies at laboratory scale for the plastic manufacturing. Some researchers have reported that these processes can be utilized individually or together for the development of new thermoplastic products which can be used like biodegradable materials for the production of diverse degradable materials with similar or better characteristics than those of commercial plastics, although, with the advantage of decreasing the adverse effect of waste disposal on the global environment (Cha, D. S. et al.,2012). In this work, eco-friendly plastic was produced using the addition of cellulose fiber from sugarcane bagasse as a reinforcement material in corn starch matrix and glycerol as a plasticizer in the process of solvent cast method.

2 Materials and methods

The experimental process for the development of bioplastic is depicted in the Figure 2.1 below. Detailed description of the process is given in the text that follows.

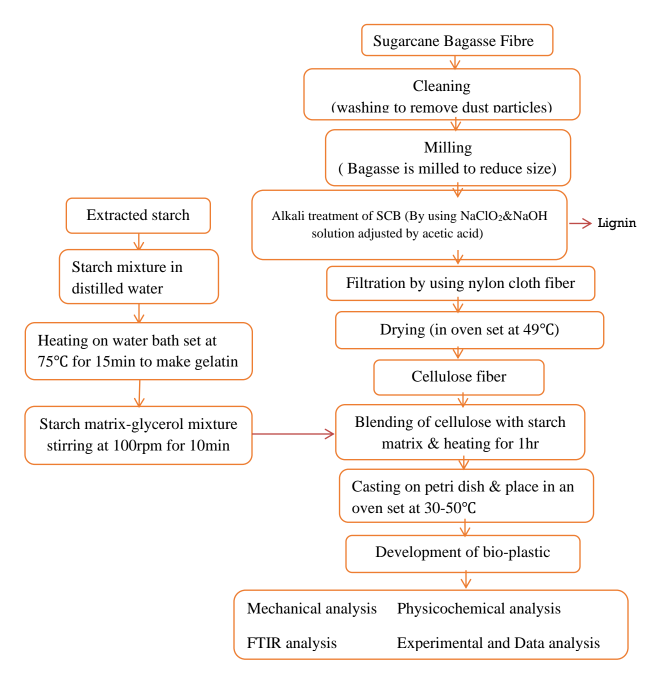


Figure 2.1 The experimental procedure for production of bioplastic from corn starch and cellulose fiber

2.1 Methods

2.1.1 Development of bioplastic from corn starch and cellulose fiber

Chemicals: chemicals used in the production of bioplasticwere glycerol, sodium chlorite (NaClO₂), sodium hydroxide (NaOH), glacial acetic acid and distilled water was the chemicals those used in development process.

Equipments: The equipments used during the experimentations includes Wiley mill, sieve, strip road, Silica crucible, desiccators, digital weighing balance, weight bottle, stopper, tong, Erlenmeyer flask, water bath, vacuum suction, weighed sintered glass crucible, what-man filter paper, measuring cylinder, round bottom flask, cellulose extraction thimble, beaker, scissor, dryer, glove, mask, thermometer, stirring rod, air circulated oven.

Location: Development of the bioplastic and analysis of its physicochemical and mechanical properties are conducted at Leather Industry Development Institute and Addis Ababa University college of natural science (chemistry laboratory).

2.1.1.1 Delignification and isolation of cellulose from sugarcane bagasse fibre

In this work the particle size was selected as determined from the literature, the dried, milled and screened by 80µm sieve size (100g of SCB) was first bleached with 0.7%(w/v) sodium chlorite (NaClO₂) solution (fibre to liquor ratio of 1: 50) at pH4, adjusted by % acetic acid that was used to acidify NaClO₂ solution. The fibre was boiled in the solution for 2hours on water bath set at 75°C to remove lignin completely and hemicellulose partially. The bleaching process was repeated for four to five times until fibre become white and then filtered. The holocellulose thus obtained was boiled with 250 ml 17.5% (w/v) sodium hydroxide solution for 5h to remove the hemicelluloses. After being filtered, the residue was washed for several times with distilled water.

At the end of extraction the insoluble residue (cellulose) was collected by filtration and it was oven dried.

2.1.1.2 Extraction of starch from corn (maize)

Extraction of maize starch was done as described by White et al., (2005) modified by Kaushik, A. S. (2010). First, we clean the shelled corn shipments to ensure that they are free from dust and foreign bodies. Once clean, the corn is soaked in water, called steep water, at 50°C for between 24 hours, during which time it doubles in size. As the corn swells and softens, the mildly acidic steep water starts to loosen the gluten bonds with the corn, and to release the starch. The corn is coarsely milled in attrition mills to separate the germ from the rest of the components. After the fine grinding, which releases the starch and gluten from the fibre, the slurry flows over fixed concave screens which catch the fibre but allow the starch and gluten to pass through. The starch, gluten suspension is sent to the starch separators. The starch, gluten suspension passes through a centrifuge where the gluten, which is less dense than starch, is easily spun out.

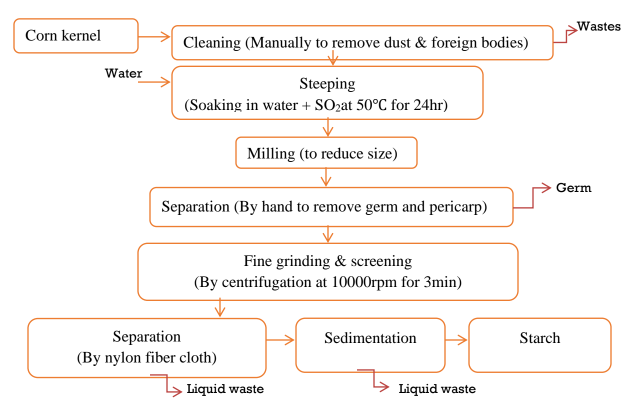


Figure 2.2Extraction of starch process flow diagram

2.1.1.3 Development of bioplastic from corn starch by reinforcing cellulose fiber

The sample was prepared by the procedure adapted from the method described by (Eraricar S.et al., 2009). A film forming dispersion was prepared by mixing the starch (10g) and distilled water (200ml). The dispersion was stirred manually on magnetic strier set at 70-80°C for 15 min while stirring at the same rate until it become gelatinized. Then glycerol(3.6ml) was added and stirred for 10 min. Cellulose fibre was then added at 0.00, 0.50, 1.00 and 1.5g (dry basis) based on a starch weight basis. Each mixture was stirred for homogeneity and to make the gelatin very strong and then allowed to cool to 75 °C before being cast on a non-stick tray. Then the solution, while still hot, the mixture was transferred into petri dishes using weight balance to keep its uniformity (Lafargue et al. 2007). Then dishes will place in an oven set at 50°C, 40°C and 30°C until the film was dry. Subsequently the dishes are removed from the oven and the films was peeled off, and then stored at room temperature in polyethylene bag for further analysis. The analysis variables are drying temperature and fiber content in blends with starch.

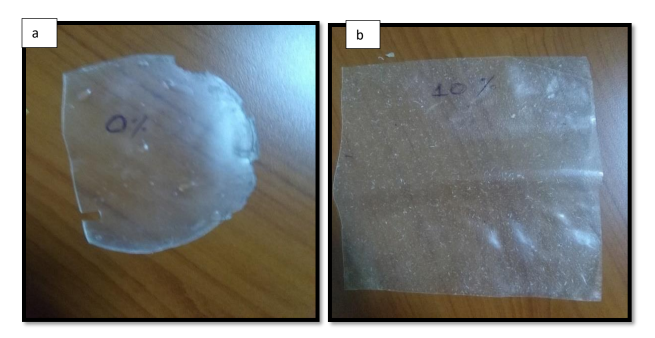


Figure 2.3pictures of developed bioplastic with a) 0% starch matrix b) optimal cellulose fibre (10%wt)

2.2Evaluation of mechanical properties of developed bioplastic

Tensile strength and elongation at break

Tensile strength and elongation at break are the most important mechanical properties of the packaging bio-plastic. Both Tensile strength and percentage elongation at break were calculated as follow

Tensile strength at yield (MPa) =
$$\frac{Force(N)}{Area(width in mm x thickness in mm)}$$
 (2.1)

$$\%Elongation = \frac{Increase in length}{Original length} x 100$$
 (2.2)

2.3 Characterization of physicochemical properties of produced bioplastic

For every physical testing, before starting any test the sample was conditioned for 48hr in a specified temperature and humidity. For this purpose we use the standard test method(ISO 2418:2005 and ISO2419:2005, Sampling and Conditioning). Water absorption was estimated by using the following formula

Water Absorption =
$$\frac{W_{f-}W_{o}}{W_{o}} \times 100\%$$
 (2.3)

Where, W_f = represents the final weight of the film after 5 minutes of absorption W_o = is the initial weight of the film.

The amount of water solubilty was estimated using the following formula:

Water Solubility =
$$\frac{M_f - M_i}{W_f} \times 100\%$$
 (2.4)

Where M_i = is the initial mass and

 M_f = is the final mass of the sample.

The transparencies of the films are determined using spectrophotometer (UV 7804C). The transmittance of films will be determined at 600nm as described by Bourtoom and Chinnah (2009). The film samples are cut into rectangles and placed in the internal side of the spectrophotometer cell.

Transparency(%T) =
$$\frac{-\log T600 nm}{X}$$
 (2.5)

Where T600 = is the transmittance at 600nm and X is the film thickness (mm)...

2.4 Experimental design and data analysis

Response surface methodology (RSM) will be adopted in the design of experimental combinations.A three variable (three levels of each) central composite experimental design was employed (Patrycja, Wojciechowska. 2012). The ratio of cellulose fiber concentrations has strong impact on the final performance of prepared bioplastic. Cellulose extracted from sugarcane bagasse is used to improve the mechanical properties of the starch based bioplastic for different application.

The two main factors selected in this study were oven-drying temperature, concentration cellulose fiber. The level of the selected factors is determined from the literature research and is presented in table 2.1the experiment performed as a completely randomized design with two main factors at three levels and three response variables. The processing variables (concentration of cellulose fiber and drying temperature (°C) were optimized using RSM to study their effect on the functional properties of prepared plastics. The responses that were considered during the optimization of the Processing variables are solubility, water absorption, tensile strength and elongation at break. The prepared bioplastics physical (solubility and water solubility) and mechanical (TS and EB) properties, and the effect of independent variables on the prepared bio-films were checked.

Table 2.1Levels of independent variables for the development of bioplastic based on central composite design

Independent Variable	Units Coded symbol		Coded Levels			
		Syllibor	-1	0	+1	
Drying Temperature	°C	A	30	40	50	
Cellulose fiber to Starch	w/w%	В	5	10	15	

Data analysis has performed by DESIGN EXPERT® 6.0.8 software using Response surface methodology design method and randomizes the runs. Randomization ensures that the conditions in one run neither depend on the conditions of the previous runs nor predict the conditions in the subsequent runs. Randomization is essential for drawing conclusions from the experiment, in correct, unambiguous and defensible manner.

Table 3.2Two factors, three levels face centered cube design with five center point formulation

Standard order	A:Temperature °C	B:Cellulose fiber (%wt)
1	30.00	5.00
2	50.00	5.00
3	30.00	15.00
4	50.00	15.00
5	30.00	10.00
6	50.00	10.00
7	40.00	5.00
8	40.00	15.00
9	40.00	10.00
10	40.00	10.00
11	40.00	10.00
12	40.00	10.00
13	40.00	10.00

3 Result and discussion

The determination of mechanical properties involves not only scientific, but also technological and practical aspects(Cosimo, C. E. 2013). Thus, the mechanical properties of the films were studied to determine the values of important parameters such as tensile strength (TS) and percent of elongation at break (E). The value of tensile strength, water absorption and Elongation at break for Bio-plastic at given level of two factors is depicted in table 3.1 below

Table 4.1Experimental design and responses

Run order	Factor	Factor	Response 1	Response 2	Response 3
	A(°C)	B(wt%)	Tensile strength(MPa)	Water absorption(%)	Elongation atbreak(%)
1	30.00	5.00	10.79	44.54	27.3
2	50.00	5.00	15.47	37.35	24.41
3	30.00	15.00	15.58	30.24	15.89
4	50.00	15.00	25.75	22.23	7.85
5	30.00	10.00	17.87	30.37	14.88
6	50.00	10.00	24.83	22.23	10.42

7	40.00	5.00	16.55	37.02	22.99
8	40.00	15.00	25.23	20.45	9.01
9	40.00	10.00	26.25	20.88	5
10	40.00	10.00	26.01	19.97	7.56
11	40.00	10.00	25.89	20.52	4.32
12	40.00	10.00	26.81	20.25	4.7
13	40.00	10.00	26.47	19.97	4.85

The resulting data, Table 3.1; were analyzed using Design expert® 6.0.8 software to determine the effects of oven-drying temperature and concentration of cellulose fiber. The dependent variables used as a response parameter were the tensile strength, water absorption and elongation at break. All experiments were carried out in a randomized order to minimize the effect of unexpected variability in the observed response due to extraneous factors. And the design summary for the experiment is shown in table 3.2.

Table 3.2 Design summery

Study Type	Central
	composite
Initial Design	Response
	surface
Center Points	0
Design Model	Quadratic
Runs	13
Blocks	No Blocks

Factor	Name	Units	Туре	Low	High Actual	
				Actual		
A	Temperature	°C	Quadrati	30	50	Levels: 3
			С			
В	Cellulose	%	Quadrati	5	15	Levels: 3
	fibre		С			

3.1 Development of regression model equation

Table 3.1 summarizes the result obtained with the experimental design which was aimed at determining the condition that favors maximum tensile strength reduce water absorption and increase elongation at break. A quadratic model equation shown below was fitted to the data model for predicting responses; Tensile strength, Water absorption and elongation at break respectively. F ratio is calculated for 95% of level of confidence.

Tensile strength of produced bioplastic

Equation in Terms of Coded Factors:

Tensile Strength = $+26.16 + 3.63 * A + 3.96 * B - 4.48 * A^2 - 4.94 * B^2 + 1.37 * A * B$

Equation in Terms of Actual Factors:

Tensile strength = $-76.77115 + 3.67341 * A + 3.64608 * B - 44805 * A^2 - 0.19762 * B^2 + 0.027450 * A * B^2 + 0.$

Where: A = Oven-drying temperatureB= Cellulose fiber

Water absorption of bioplastic

Equation in Terms of Coded Factors:

Water Absorption = $+21.78 - 3.72 * A - 6.33 * B + 6.35 * A^2 + 4.29 * B^2 - 0.70 * A * B$

Equation in Terms of Actual Factors:

Water Absorption = $+21.78379 - 3.72333 * A - 6.33167 * B + 6.35172 * A^2$

+4.28672 * B² -0.70500 * A * B

Where: A = Oven-drying temperature B= Cellulose fiber

Elongation at break for bio-plastic

Equation in Terms of Coded Factors:

Elongation at break =+5.91-2.57 * A-7.00 * B+5.20 * A^2 +8.55 * B²-1.28 * A * B

Equation in Terms of Actual Factors:

Elongation at break = $+137.28264-4.15822 * A -7.21422 * B +0.051957 * A^2$ $+0.34183* B^2-0.025550* A*B$

Where: A = Oven-drying temperature B= Cellulose fiber

3.2Effect of process variables

The effect of each independent variable on the physical and mechanical properties of the bio films was investigated by keeping other variables constant. Moreover, from the model equation, the coefficients of the independent variables show the effect of each independent variable on physical as well as mechanical properties of the bioplastic.

3.2.1 Effect of individual process variables

3.2.1.1 Effect of drying temperature on tensile strength of bio plastic

The effect of drying temperature on Tensile Strength of bio-plastic was shown in Figure 3.1 below. From the graph it can be observed that as temperature start to increase from low level coded (-1) up to center point coded (0) tensile strength also increases, but beyond center point the tensile strength shows slight decreasing and become parabolic. This can be caused by the influence of higher temperatures that can cause intermolecular bonds in starch chains becoming weaker. The hydrogen bonds between amylose chains undergo termination of the bond. Then further heating will break

glycosidic bonds (bonds between monomers) in amylose. Based on research Gilfillan WN, N. D. (2012) increasing the heating temperature can cause depolymerization in amylose chain, the straight chain amylose falater and become shorter, thus decreasing amylose content (Halley, 2001).

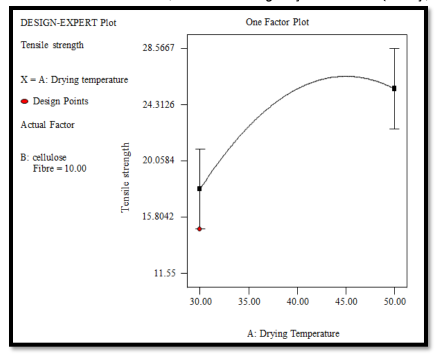


Figure 3.1 Effect oven drying temperature on tensile strength of developed bio-plastic

3.2.1.2 Effect of cellulose fiber concentration on tensile strength

As depictedin figure 4.2 below the effect of Cellulose fiber to starch ratio on tensile strength can be observed that as the ratio of Cellulose fiber increases the response of tensile strength increases up to center point coded (0) and beyond to some extent and become decreases as cellulose fiber increases further up to higher level coded(+1). The fibre content in the matrix of synthetic or biological polymers increased the values of elasticity modulus and tensile strength of bioplastic; and decreased the values of elongation at fracture (Oskman et al., 2003).

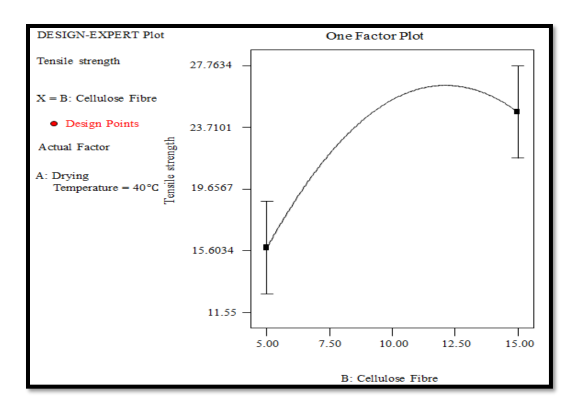


Figure 3.2 Effect of cellulose fiber concentration on tensile strength of developed bioplastic

3.2.1.3 Effect of drying temperature on water absorption of bioplastic.

The effect of drying temperature on water absorption of bio-plastic was shown in figure 4.5. As shown in the figure, the lowest water absorption was observed at center point coded (0) or at 40° C. As we move from lowest coded level(-1) to center point the water absorption decreases, and as drying temperature increases from center point to higher level coded(+1) water absorption start to increases this was in agreement with some literature results.it was found out that in the work of (Ahmedet al., 2014)

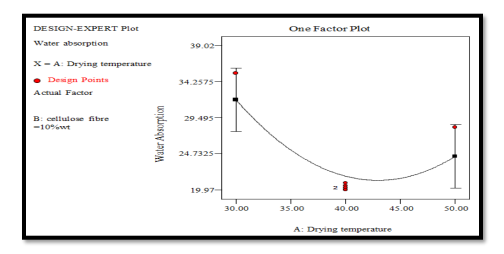


Figure 4.3 Effect of oven drying temperature on water absorption of developed bio-plastic www.journal-innovations.com

3.2.1.4 Effect of cellulose fiber on water absorption of bioplastic

As shown from the figure bellow the water absorption decreased when the percentage of Cellulose fiber increased from lowest level to center point coded (0) or up to 10% of Sugarcane bagasse cellulose in the bio-composites, as we move from center point to higher level water absorption starts to increase slightly and become a show parabolic line. According to Sarifuddinet al. (2012), this moisture absorption reduction in bio-composites can be attributed to stronger hydrogen bonds between the matrix and the reinforced material. Furthermore, Müller et al. (2009) demonstrated the resistance to water absorption in TPS when cellulose fibers are incorporated.

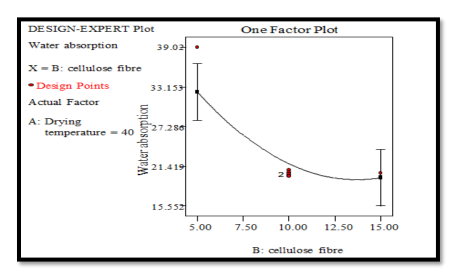


Figure 3.4 Effect of cellulose fiber on water absorption of bioplastic

3.2.1.5 Effect of drying temperature on elongation at break of bio-plastic

From figure 4.5 it illustrated that increasing oven drying temperature of the bio - plastic solution causes the value of elongation at break decreases until it reach the center point. Which was due to increasing the tensile strength of bio-plastic as well as increasing drying temperature from lower level coded(-1) up to center point coded(0). The result of elongation at break decreases with increasing the temperature and as shown from the figure as drying temperature increases to high level coded(+1) elongation at break start to increases smoothly. This is because the heat that is given causes an increase in the kinetic energy of the molecules in which the molecules vibrate and create a free volume to allow larger molecular chains rotation (Patrycja, Wojciechowska, 2012).

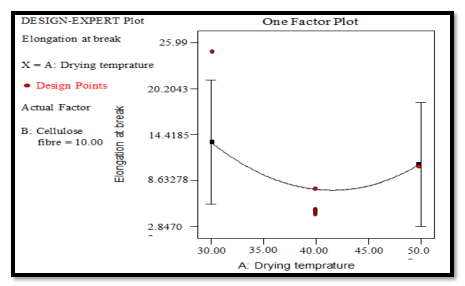


Figure 3.5Effect oven drying temperature on elongation at break of bio-plastic

3.2.1.6 Effect of cellulose fiber on elongation at break of bioplastic

As shown from the figure below increasing fiber content from low level coded(-1)up to center point coded(0) decreases response elongation at break up to cellulose fibre content reaches 10% and elongation at break become increasing as cellulose fibre content further increases beyond 10% or center point. This is in agreement with other literature, the work done by Funkeet al (1998).

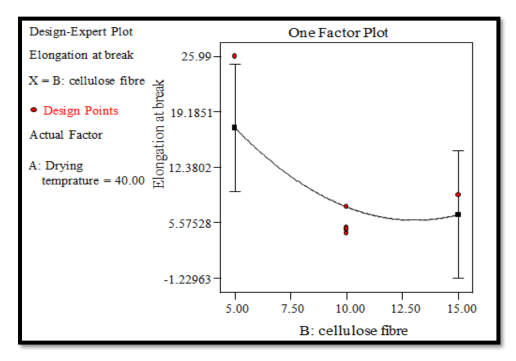


Figure 3.6Effect of cellulose fiber on elongation at break of bioplastic

3.2.1.7 Effects of interactive parameters between process variables on tensile strength, water absorption and elongation at break

Figure 3.7 shows the effect of the interaction of drying temperature and starch-cellulose fiber ratio where the maximum values of tensile strength (26.81 MPa) were at drying temperature of 40°C and cellulose fibre (10 wt%) concentration. However, further increase in oven drying temperature above 40°C and the increase of fibre content above 10 % in the blend smoothly increase the values of water absorption and elongation at break. In general the values of tensile strength were favored at the center values of drying temperature and cellulose fiber. The increment of oven drying temperature values decreased drastically both water absorption and elongation at break (see figure 3.11 and 3.12 below) for water absorption and elongation at break respectively

In a general way the values of Water absorption and elongation at break were increased with an increasing of and cellulose fibre and also at high values of drying temperature and cellulose fibre the value of both responses start to increase. Thus, the values of both water absorption and elongation at break were favored at 40°C and 10% of cellulose fibre (CF) and counter surface plot. The incorporation of cellulose fibre acts as reinforcement in thermoplastic materials (Averouset al., 2001).

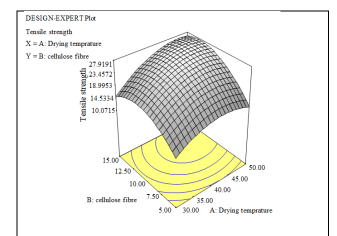


Figure 3.7Effects of drying temperature and fibre interactive on strength, water absorption and elongation at break.

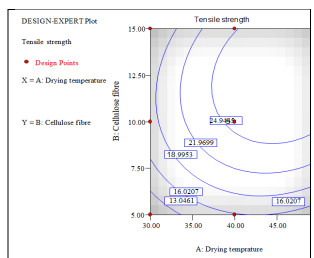


Figure 3.8 Effect of drying temperature and cellulose fibre on tensile strengthcontour plot bioplastic

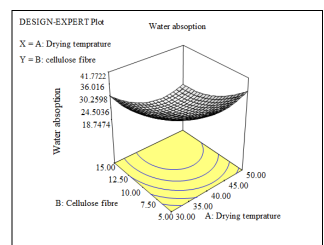


Figure 3.9 Interactive effects of drying temperature and cellulose fibre on water absorption of developed bioplastic

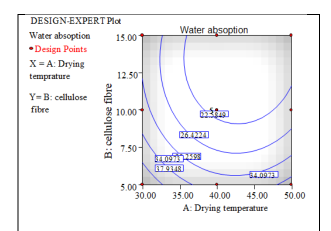


Figure3.10 Contour effect of drying temperature and cellulose fibre on water absorption of bioplastic

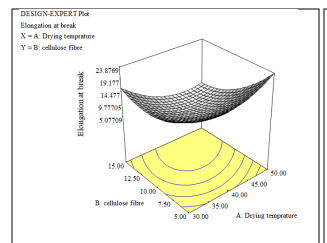


Figure 3.11 Interactive effects of drying temperature cellulose fibre and elongation at break of bioplastc

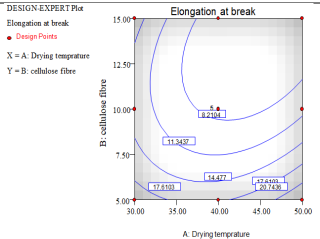


Figure 3.12Effect of drying temperature and cellulose fibre on elongation at break contour figure

3.3 Optimization of process factors

The broad optimization objective is the maximization of quality and minimizing of oven drying costs. RSM (Response Surface Methodology) is a collection of statistical and mathematical techniques used for developing, improving, and optimizing processes in which a response of interest is influenced by several variables and the objective is to optimize the response.

Optimization analyses were conducted on the data from the Central Composite Design(CCD) to relate drying temperature and sugar cane bagasse cellulose fibre concentration to Tensile strength, water absorption and elongation at break of developed specimen. The numerical optimization was generated by design expert 6.0.8 software and elucidated in a table 4.6 below as a function of two factors drying temperature and sugar cane bagasse cellulose fibre. An optimal processing conditions of the bioplastic developed from corn starch and cellulose fibre was selected for further characterization based cellulose fibre and oven drying temperature setting in the range and as well as maximizing the tensile strength and minimizing water absorption and elongation at break of the sample (see table 4.3 below). Therefore, considering those parameters the value obtained from optimum experimental for drying temperature (40.85°C), cellulose fibre (12.20%), tensile strength (27.2175), water absorption (18.3257%) and Elongation at break (4.25203%) were obtained with desirability of 1.00

Table 3.3 Optimization constraint for developed bioplastic

Constraints Name	Goal	Lower Limit	Upper Limit	Lower Weight	Upper Weight	Importance
Drying temperature	Is in the range	30	50	1	1	1
Cellulose fiber	Is in the range	5	15	1	1	1
Tensile strength	Maximize	10.79	26.81	1	1	1
Water absorption	Minimize	19.97	44.54	1	1	1
Elongation at break	Minimize	4.32	27.34	1	1	1

3.4 Physicochemical characterization of developed bioplastic from starchcellulose based fiber

3.4.1 Compound microscopy

In Fig 4.13 the compound micrographs of bio-plastics prepared with particle sizes of 80µm (a) unfilled starch matrix of control and (b) reinforced bioplastic are shown. The biocomposite samples observed in Fig.4.13 correspond to thermoplastic material prepared unfilled matrix without reinforcing cellulose particles, and the samples shown in (b) prepared with a treated sugarcane bagasse fiber (cellulose) amount of 10% contained. As can be observed in reinforced bioplastic, an even distribution of cellulose particles in the thermoplastic starch matrix is clearly evident due to the obtained transparency. In addition, the methodology applied to prepare bioplastic in the present work avoids problems such as fluidity by high viscosity in the processing of the thermoplastic starches by extrusion and injection that is determined in literature (Ma et al. 2005).

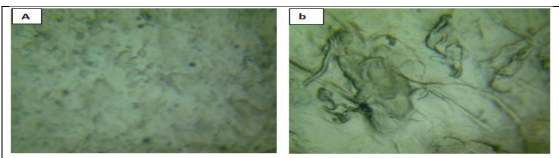


Figure 3.13 Micrographs of bioplastic developed from a) unfilled starch matrix and b) filled with 10% treated fiber.

3.4.2 FTIR spectroscopy analysis

Figure 3.14 shows the FTIR of the initial bio plastic from starch (100%) as a control. The starch spectrum shows the common signals for these kinds of polysaccharides, with glucopyranose rings such as OH bands at 3450 cm⁻¹, C-H stretching vibrations of aliphatic groups at 2926 cm⁻¹, adsorbed water signals at 1646 cm⁻¹, C-C and C-O stretching at 1100 cm⁻¹, and C-O-H bending vibration at 1007 cm⁻¹. It also can be seen that pure TPS bio-film exhibits FTIR main peak positions in the range of 1458 to 1150cm⁻¹, 1410-1230cm⁻¹, and 1200-1000cm⁻¹, representing C-H asymmetric stretching of -CH₂-, -CH₂- deformation, C-O-C stretching, and C-O-H stretching, respectively. The peak position at approximately 1646 cm⁻¹ is due to the bound water present in the starch (Prachayawarakorn *et al.*, 2011).

Table 3.4Signal Assignments of Functional groups identified by Fourier transform- Infrared spectroscopy analysis of bioplastic

Wave number (cm ⁻¹)	Functional groups
1005-897	C-O-H stretching
1150-1005	C-C stretching
1235-1150	C-O-C stretching of Easter
1734-1631	Bounded water
3000-2826	C-H stretches
3600-3450	-OH stretching

The FTIR spectra of bio-composites with reinforced material (sugar cane bagasse cellulose fibre) content amounts (10%) and organized by particle sizes of 80µm which is keeping it at constant is depicted in Figure.4.15. In this case, different signal patterns are observed from bio-composites prepared (Fig. 4.15) as a result of their different chemical composition due to addition of fiber, the following changes in the intensity of some signals, such as those at 1764, 1660, and 1424 cm-1, can be attributed to the addition of cellulose fibre at optimal concentration. The amount of adsorbed water seemed to have a small increase in the bio-composite that had 10% fibre content. However, this may be only an optical effect caused by the decrease of the adjacent carbonyl band (1720 cm⁻¹). In contrast, the signal at 1099 cm⁻¹ for control decreased to 1050cm⁻¹ when 10% of cellulose fibre was added to the starch.

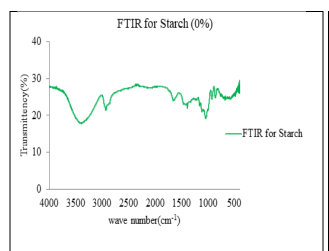


Figure 3.14 FTIR result of the starch-based bio-plastic plot

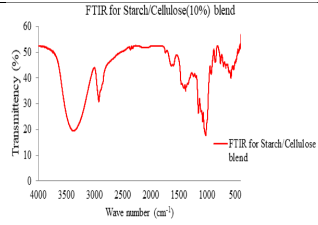


Figure 3.15 FTIR result of starch-cellulose, bio-plastic plot

3.4.3 Transparency of developed bioplastic

Transparency of the films is of importance in some instances, when used as packaging materials (Assefa Z *et al.*, 2013). Addition of cellulose fiber generally causes the films to reduce their transparency, but at less small difference compared to control. Starch composite film without cellulose was rather transparent.

The percentage of transmittance of corn starch film at a wavelength of 600 nm and containing optimum cellulose fiber (CF)(10% concentration) prepared by solvent casting process. The results showed (see Table 3.5) that the films were off white in color but highly translucent. Addition of cellulose increased film cloudiness for starch-based films. This was due to the cellulose being cloudier than the corn starch so the transparency decreases of 118% for starch to 113% for 10% cellulose containing bio-plastic (see Table 3.5).

3.4.4 Solubility and density of developed bioplastic

The film with a 10% of cellulose fibre content had a lower swelling capacity than the films with only containing starch matrix. From the experimental result the solubility of the specimen was 10.17% for starch-cellulose based (10% cellulose fibre) bioplastic and 14.78 % for 100% starch-based bioplastic can recommend for packaging purposes. Density of film developed from 100% starch and 10% CF containing was 1.059 g/ml and 1.068g/ml, respectively (see Table 3.5), there was an increment in density in the case of CF, this is may be due to the addition of 10% of cellulose fibre.

Table 3.5Experimental result data for measuring bioplastic transparency, solubility and density

Product	Transparency (%)	Solubility (%)	Density(g/ml)
Starch (100%) as a control	118	14.78	1.059
10%SCBF containg	113	10.17	1.068

4 Conclusion and recommendation

4.2 Conclusion

In this study, development of bioplastics from corn starch and cellulose fibre from sugar cane was investigated. Sugar cane bagasse fibers with an average diameter of 80µm derived from bleached or treated pulp, slightly decreased the moisture content of starch films, but doubled the tensile strength. These tensile test results are slightly higher than those obtained by wheat grass cellulose fibre composites but similar to those of flax cellulose crystals (Cao et al. 2008). As well as, in this work the strength of the cellulose fibre composite decreased at fibre loadings above 10 wt%. This may be caused by fibre agglomeration and entanglement due to the cellulose fibre having a high aspect ratio.

The output of the experiments conducted has been analyzed by design expert 6.0.8 of three levels and two factor central composite designs and response surface methodology was employed. In the development of bioplastic the three responses were significantly depend on starch-cellulose fibre ratio and oven drying temperature. Generally, as processing condition (cellulose fibre and oven drying temperature) increase from low level to center point tensile strength increases, but, was slightly decreased at some interval as both processing conditions increased from the center to the highest level. From the design software employed the maximum tensile strength of 26.81 MPa, water absorption of 20.52% and elongation at break of 4.32% were obtained at optimal value of the processing condition. Using numerical optimization of design expert 6.0.8 for development of bioplastic from cellulose fibre and cornstarch at operating condition for drying temperature (44.68°C), cellulose fibre (13.99%) determined for developing bioplastic by solvent cast process.

Compound macroscopic and FTIR analyses provided evidence of the strong interactions that existed between starch and bagasse fibers. The Compound micrographs revealed that the cellulose fibers are well wetted by the starch matrix, and showed that the fibers are strongly attached to the starch matrix since it pulls the matrix out of shape where the fibre is attached during tensile testing.

Results showed that the starch composites reinforced with sugar cane bagasse cellulose fibre have a potential application in biodegradable packaging and biocomposite medical science (Alvarezet al., 2004). Regarding transparency, solubility and density the developed bio plastic at optimal point shows lower transparency, solubility and higher density (13%, 10.7% and 1.068q/ml) compared to control (100% starch) made bioplastic(118%, 14.7% and 1.059g/ml restively) this was due to introduction of fibre into a starch matrix which from hydrogen bond.

4.3 Recommendation

Even though bioplastic is produced further investigations of the following points can be recommended.

- Since corn starch is edible food it creates problem of food security of our country, so focus should be given to non-edible based starches
- > Based on the present study developed bioplastic can be used for food packaging so in the future study focus on the feasibility of using bioplastic for food packaging.
- In order to explore more effective bioplastic based on starch and cellulose fibre more wide studies should be carried out by considering other processing factors such as drying time, plasticizer, starch gelatinization temperature, fibre particle size and solvents on the quality, mechanical and physicochemical property should be investigated.
- Since corn starch is edible food it creates problem of food security of our country as well as the world, focus should be given to non-edible based starch source.
- > In addition, extensive researches are still needed on the permeability (water vapor and gas transmission rates), barrier properties, and the effect of hydrophobic plasticizers and effectiveness of these bioplastic on different application.

Conflict of Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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