Isolation and characterization of micro cellulose obtained from waste mango

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Abstract

Cellulose fibers are used in polymeric matrices due to their thermal and mechanical capabilities. These biopolymers can be isolated from different natural sources. In this study, micro cellulose was obtained from mango fibrous endocarp (Mangifera caesia Jack ex Wall) waste. Isolation was performed using sulfuric acid and sodium hydroxide as removal agents of lignin and hemicellulose. A comparative analysis between native fiber (NF) and micro cellulose (MC) was performed, using FTIR, DSC and SEM techniques to assess their composition, thermal, structural and crystallinity capacities. The structures of the fibers were not damaged due to the chemical treatments received and their sizes ranged between 40 and 400 μm in length. The extraction of cellulose from mango waste represents an important start in obtaining biopolymers that can be used in the food, pharmaceutical, and other industries.

Keywords: biopolymers, mango waste, micro cellulose.

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1. Introduction

In recent years, different ways have been sought to control and reduce pollution caused by petroleum-derived materials¹-³, and one of the most used alternatives in the world is recycling⁴. However, and even though this conventional method reduces the ecological impact of plastic has not been possible to reduce the large volumes of those kinds of materials that are produced each year⁵. This is mainly due to the fact that a great many products are manufactured in a way that make the plastic content difficult to separate and, therefore very little plastic is currently recycled. An estimated 8.3 billion metric tons of plastic are produced worldwide, and only 9% goes through recycling process⁶. Another alternative for reducing environmental impact of plastics that has been proposed is the use of polymeric materials combined with pro-oxidizing substances⁷-⁸. However, the wastes the materials generate after they are degraded consist of heavy metals and petroleum products⁹-¹⁰, which tend to accumulate in the environment.

To reduce the impact caused by petroleum products, there is need to look for materials that have a polymer matrix formed by biopolymers (natural polymers)¹¹. Since the polymer matrix is biodegradable, the composite which is its integral part is also expected to be biodegradable and environmentally friendly¹². It has an advantage over synthetic polymers because it can be totally biodegraded in products such as CO₂, water and also organic fertilizer¹³. Bioplastics and biomaterials are composed of biopolymers, biomass derivatives or blends of natural and synthetic polymers, which will help mitigate environmental issues in the long term.

Agriculture in Mexico is an important productive sector. The role played by the agricultural sector in the other sectors of Mexico such as economic, social and environmental determines its greatest impact on the country’s development. However, post-harvest residues (bagasse, husks and seeds) represent an environmental problem due to the followings: lack of distribution channels, pests, environmental policies and inadequate management of the large volumes of residues that are generated. Therefore, agricultural residues must be considered as a source sustainable and renewable of products such as cellulose fiber used as a reinforcement for bioplastics and biomaterials, due to their high availability and potential¹³,¹⁴.

Waste mango (seeds, shells and cotyledons) currently have little impact on industries. However, these wastes have been studied to obtain active and functional compounds from mango peel¹⁵, antioxiants from cotyledon¹⁶, to use fibrous endocarp in the paper industry¹⁷, and pulp to extract reducing sugars, organic acids and obtain alcohols¹⁸.

Unlike other agricultural residues, mango (Mangifera caesia Jack ex Wall) is composed of more than one biopolymer, with starch and cellulose being the highest proportion¹⁹. In addition to being the most abundant natural polymer in the world, cellulose is renewable, biodegradable and is a classic example of a matrix reinforcement biopolymer²⁰,²¹ and is expected to be able to reduce and replace many non-renewable materials²². The fibrils have two regions...
known as crystalline (ordered regions) and amorphous (disorder regions). These regions can be hydrolysed with acid, which makes the crystalline region less reactive. This results in obtaining microfibrils, nanofibrils and cellulose nanocrystals.[23]

Cellulose fibers have an inherent structural hierarchy that originates from their different biological sources. Obtaining cellulose from different lignocellulosic fibers has attracted the interest of the scientific and industrial community in recent years.[24,25] The native fibers are longitudinal, rigid and difficult to modify due to the components that form them. These compounds work as a natural glue that gives firmness and shape. The fibers are hydrophobic. This is why, it is necessary to perform a chemical and thermal treatment, that can help to remove glue from the fibers without damaging the structure of the micro-cellulose fibers. The particle size reduction in the cellulose fibers through thermal treatment, generates major areas of contact with chemical substances such as acids and alkalis. Likewise, with the ultrasonic treatments crystallinity increase and morphology of cellulose fibers can be improved; in the same way individual fibrils could be obtained from waste mango and which serve in developing a bio-based material.

In the present study, cellulose fibers were obtained from waste mango fibrous endocarp. The shape, size and structure of the fibers were characterized as well as the decrease in lignin and hemicellulose subjected to a modified acid-alkaline treatment was analysed by wet chemistry methods. Taking advantage of waste residues facilitates the use of cellulose as a biopolymer and reduces the impact generated by inappropriate disposal of agricultural waste and are able to generate profits.

2. Materials and Methods

2.1 Cellulose insulation

Fibrous endocarp of Ataulfo mango fruits (Mangifera caesia Jack ex Weill) was obtained in the local market in El Arenal, Guerrero, Mexico, and used as the raw material for this study. Reagents used to obtain and extract the native cellulose fibre (NF) and chemically modified cellulose fibre (designated micro-cellulose, MC) were: Sodium hydroxide (CAS 1310-73-2), sulfuric acid (CAS 7664-93-9), hydrogen peroxide (CAS 7722-84-1) and distilled water (CAS 7732-18-5).

2.2 Pre-treatment of mango fibre

The mango endocarp was cut into squares of approximately 1 cm². 50 g was weighed and hydrated using distilled water. It was stirred at room temperature for two hours and subsequently dried in a continuous flow oven at 40 ºC for 24 hours.[26]

2.3 Alkaline treatment

The dried fibrous endocarp was treated in a solution of 2% NaOH (w/v) in a 1:20 ratio (fibre: solution). The solution was continuously stirred at 80 ºC for two hours. After the alkaline treatment was done, the fibres were washed with distilled water and dried at 40 ºC for 24 h. The dried fibres were grinded using an E3303.00 mini cutting mill (Eberbach Corp.), equipped with a stainless steel blades to separate the bonded fibres formed after the drying process. After the milling was completed, the fibres were stored in an airtight bag.[27]

2.4 Whitening

The fibres were bleached in a 1:20 ratio (fibre: solution) in a solution composed of H₂O₂ (v/v) and 4% NaOH (w/v), and stirred for two hours at 50 ºC. After the treatment, the fibres were washed with distilled water and dried at 40 ºC for 24 h.[28]

2.5 Acid Hydrolysis

The fibres were subjected to 52% sulfuric acid (w/w) in a 1:20 ratio (fibre: solution), and stirred for two hours at 45 ºC. The reaction was stopped by thermal shock using a vessel with distilled water at a temperature of 10 ºC.[27]

2.6 Sonication

Sonication process was applied after acid hydrolysis in order to disintegrate the fibres into microfibres in water dispersion to reduce the viscosity and allow efficient propagation of the vibration through the dispersion. Szymańska-Chargot et al.[29] and Kasuga et al.[30] techniques were modified and the dispersion was then subjected to sonication for 5 min at a power of 99 W using a Branson 5510MT ultrasonic cleaner. Subsequently, the samples were centrifuged at 6300 x g at 4 ºC for 15 min and dried at 40 ºC for 24 h.[27]

2.7 Chemical analyses

The lignin content was determined according to Klassen method[31]. The dried fibres was extracted by using Soxhlet method[32] with ethanol-benzene solvent for 5 h. The residue extract was oven dried at 103 ºC for 1 h, and later treated with 72% H₂SO₄ for 2 h while stirring at 37 ºC. The material was then diluted to 3% H₂SO₄ and then refluxed at 80 ºC for 4 h; the sample was oven dried at 105 ºC for 1 h and cooled in a desiccator to obtain constant weight. The lignin content was calculate using the following equation[31]:

\[
\text{Lignin} \% = \frac{A_{100}}{W}
\]

Where \( A \) is weight of lignin and \( W \) is oven-dry weight of test specimen.

Holocellulose was measured by means of the Wise chlorite method[33] through dried fibers treatment with an acidified sodium chlorite water solution in acidic medium at 70-80 ºC for 1 h. The process was repeated until the product became white. Determination of cellulose to dry fiber samples was using the Seifert method[34] wherein the cellulose is separated from the plant material in the process of its processing with aqueous solutions of acetylacetone, dioxane, and hydrochloric acid. Hemicellulose content was calculated theoretically from the difference in the contents of holocellulose and cellulose.
2.8 Scanning Electron Microscopy (SEM)

Cellulose samples were sprinkled on a double adhesion carbon conductive tape, which was previously fixed on aluminium support of the scanning electron microscope Carl Zeiss EVO L.S. 10. Subsequently, they were observed at a voltage of 2.5 kV, with a resolution of 3-10 nm; the micrographs were taken at 100, 500 and 1500 magnifications[35].

2.9 Fourier Transform Infrared Spectroscopy (FTIR)

Fourier transform infrared spectroscopy (FTIR) studies were carried out using a Perkin-Elmer-Spectrum 100/100 N FT-IR infrared spectrometer. The infrared region was within a range of 4000-650 cm\(^{-1}\) in the transmittance mode, with a resolution of 16 cm\(^{-1}\) and 8 scans[36].

2.10 Differential Scanning Calorimetry (DSC)

A differential scanning calorimeter model TA 2010 was used. The equipment was calibrated with indium (melting point of 156.4 °C and enthalpy of 6.8 cal/g). Between 5 and 10 mg of the sample was weighed, using a micro balance with an accuracy of ± 0.01 mg, a heating program ranging from 20 to 400 °C and a heating rate of 10 °C/min[37]. This generated an inert atmosphere in the cell by circulating 50 mL/min of nitrogen gas (99.9% purity).

2.11 Crystallinity

The degree of crystallinity was determined based on the DSC enthalpy of fusion data using the following equation[37]:

\[ X_c = \frac{\Delta H_f}{\Delta H_o} \]  \hspace{1cm} (2)

Where \( X_c \) is the crystalline fraction, \( \Delta H_f \) represents the enthalpy of fusion measured by DSC and \( \Delta H_o \) is the enthalpy of fusion for 100% crystalline polymer.

3. Results and Discussions

3.1 Chemical Analyses

Despite the economic importance of agricultural residues there are no studies on the chemical composition of the mango endocarp. In general, the studies of mango chemistry are limited to chemical characteristics based mainly on mango agroindustrial waste, fruit by-products and silage[38-41].

The results of lignin and calculated hemicellulose of native (NF) and alkaline modified (MC) fibers decreased, from 10.32 wt % to 0.64 wt % for NF and 29.75 wt % to 1.11 wt %, respectively. Regarding the lignin content, the data from this investigation (10.32 wt %) is close to those found for Guzmán et al.[38], different values are obtained when the food-processing industry method is employed (10.32 wt % vs. 9.0 wt %). This method is similar to Klason lignin extraction process, except a final step of 1 h of calcination at 550 °C. Nevertheless, a large difference was observed between our lignin results: 5.76 wt % and 6.97 wt % of lignin in mango peel and mango by-products, respectively[39,40]. Concerning the hemicellulose content from this study (29.75 wt %) is lower than the values (31.1 wt % and 31.75 wt %) found for mango peel and mango by-products[38,41]. This can be explained by the fact that it seems that in the food-processing industry method, the neutral detergent solution used for determining the Neutral Detergent Fibre (NDF = Hemicellulose + Cellulose + Lignin) content does not remove completely hemicelluloses and is, therefore, this method is not indicated for this work.

3.2 Structural characterization

The native fibres presented very dense outer layers covering the cellulose sacs. The layers are considered to be composed of lignin (Figure 1a). Lignin wraps the cellulose in the inner part covering it with its stiffness and hardness property. Small globular particles and pores along the fibres and impurities can be seen on the surface.

Figure 1b shows an accumulation of long fibres, rolled and small traces of waste. This is attributed to the modification of the alkaline treatment using NaOH. Cellulose fibres can be seen more clearly, but there is still a need to remove part of the lignin and hemicellulose that holds them together. Cellulose fibres are more available due to the removal of lignin, and their characteristics (e.g. shape, surface) are similar to those of the commercially available cellulose (Cived™); the SEM images (Figure 1b and c) evidence that separate individual fibres have also similar morphologies with smooth surfaces.

The fibres went through a whitening process with the help of hydrogen peroxide. The peroxide helped in eliminating some waxes, tannins, lignin and alkaline treatment residues.

In Figure 2a, completely separated fibres can be seen; there are no traces of lignin, as well as porous components. In this procedure, the decrease in thickness, shape, size and discolouration of the fibres is more evident. After bleaching, the fibres had sizes ranging from 40 to 400 μm in length with a uniform surface and rectangular shape (Figure 2b). Owi et al.[42] characterized micro celluloses chemically isolated from fruit bunches and sugarcane bagasse as well as commercial cellulose (Sigma™) with characteristics similar to those obtained in this study after extraction, particularly in fibre shape and surface. In a similar manner, Nagalakshmaiah et al.[43] report the shape and diameter of chilli fibres by an acid hydrolysis treatment. These results are consistent with the ones presented in this study. This is consistent with the report of Atiqah et al.[44] that amorphous portions like hemicellulose and lignin were removed after applying alkali and bleaching treatment on raw kenaf fibre.

3.3 FTIR analysis

Figure 3 shows the native and micro cellulose fibre spectra obtained by chemical treatment. A peak at 894 cm\(^{-1}\) can be observed in the fibres with alkaline treatment. This signal is attributed to the anomeric carbon present in cellulose. This signal is usually in the polysaccharide absorbing region of 950-700 cm\(^{-1}\), which also reveals the component structures of β-glucans. In the native fibre, this signal is less intense because the lignin coating prevents specific cellulose signals that are present.

A narrow absorption band in the region between 1170 and 920 cm\(^{-1}\) ascribed to the polysaccharide bands, shows the stretching vibrations of C – O, C – C and C – OH bonds which
Figure 1. Micrograph of (a) native mango fiber, (b) alkaline modified fiber (c) commercial cellulose (Civeq™).

Figure 2. Micrographs of (a) fiber after whitening (b) and thermal sonication.

Figure 3. Infrared spectra for native fibre (NF) and microcellulose (MC).

are in cellulose, lignin and hemicellulose. A strong signal was present in the treated fibre (MC). The increase is based on the breaking of bonds during the acid-base treatment.

The native fibre showed a peak at 1230 cm⁻¹. This is attributed to the stretching of C – O – C bonds of the alkyl-aryl ether, a compound present in the structure of lignin, and which disappears after chemical treatments. This means that the lignin was partially removed in MC as further confirmed by lignin results obtained in this work, 0.64 wt %, determined by Klason method.

In the native fibre, a characteristic peak of the C–C unsaturated bond of lignin was observed at 1623 cm⁻¹, as well as the water band at 1636 cm⁻¹ decreasing its intensity slightly in MC. Likewise, a peak of 1740 cm⁻¹ is observed,
which is associated with the stretching of $C = O$ groups linked to aliphatic carboxylic acid and ketone. It could also be related to the presence of hemicellulose. Both peaks decrease when the sample is subjected to acid treatment, which allows the removal of lignin and hemicellulose, from 10.32 to 0.64 wt % and from 29.75 to 1.11 wt %, respectively.

Between the range of 2940 and 2800 cm$^{-1}$, there are two peaks in the native cellulose, which are assigned to the asymmetrical and symmetrical stretching vibrations of alkyl, aliphatic and aromatic rings ($H – C – H$) groups present in hemicellulose and lignin. These bands were ascribed to the aliphatic materials such as cutin, waxes and cutani. The signal decreases when the sample is subjected to an acid-base treatment, due to the removal of these components. 1740 cm$^{-1}$ band associated with the stretching of $C=O$ in ester groups disappeared after basic hydrolysis, confirming the partial removal of hemicellulose, which presented variation in hemicellulose content (e.g. 1.11 wt % compared to 29.75 wt % for NF sample).

### 3.4 Thermal characterization

As for the DSC results, both samples studied in nitrogen atmosphere show a similar trend and, all the reactions during the decomposition of the cellulose were endothermic processes in inert atmosphere (Figure 4). The native endocarp fibre presented three steps of thermal degradation in relation to cellulose, lignin and hemicellulose. This is in line with a research done by Stevulova et al.

NF shows an endotherm at 113.64 °C and a plateau at 145.3 °C; while that of MC is at 135.11 °C. This is mainly due to the elimination of moisture by the heat range applied in the DSC, as well as reducing sugars and other volatile components joined to the fibres, that are thermally decomposed. Espinosa-Andrews and Urias-Silvas mention that there is a humidity loss, specifically in fructans, in temperatures under 130 °C. In addition to this, a small endotherm at 232.12 °C can be observed in NF probably due to the degradation of hemicellulose, reducing sugars and those hemicelluloses that are still joined to the cellulose-lignin complex. NF and MC presented endotherm peaks at 325.52 °C and 335.25 °C, respectively. The endothermic peak observed in a 220-370 °C temperature range corresponds to that used for the simultaneous decomposition of hemicelluloses, cellulose and lignin. Based on the report of Miyahara et al., this endothermic peak represents the degradation of hemicellulose. As reported by Xiao et al., cellulose is degraded within the range of 250 and 345 °C.

The enthalpies in the initial endotherm peak of the NF sample (121.92 J/g) are lower compared to the MC sample (183.27 J/g). This could be due to a greater presence of traces of waste and volatile components that were impregnated through acid-alkali hydrolysis. Therefore, a greater amount of heat energy is needed for degradation to occur. Conversely, with respect to final endotherm peak for both samples, the enthalpy of MC (172.92 J/g) is lower compared to that of the native fibre (178.32 J/g), which could be due to the elimination of larger sugar chains through acid-alkali hydrolysis. Besides, under nitrogen atmosphere, cellulose is more resistant to thermal treatment, probably due to its crystalline structure. The range at which lignin is decomposed is between 250 and 600 °C. The analysis was performed up to 400 °C, which indicates the degradation of 40% of the total lignin in the native sample. In the composition of native cellulose, it can be seen that the compound that degrades first is hemicellulose, followed by cellulose and lignin.

### 3.5 Crystallinity

Applying the Equation 1 and with the use of the DSC enthalpy of fusion and taking 230J/g as the value for the fully crystalline Polycaprolactam (PA6), a lower value of crystallinity for native fibre (NF) was found compared to microcellulose (MC) (Table 1).

As reported by Li et al., the percentage of crystallinity in native fibre will always be lower, due to the presence of hemicellulose and lignin, amorphous compounds that encapsulate the fibrils. It is believed that the crystallinity index in micro cellulose increases due to the elimination of the amorphous regions of cellulose, in which hydrolysis drives the breakdown of glycosidic bonds, releasing individual crystals. The crystallinity of mango micro cellulose reported in this study is similar to that reported by Bolio-López et al. in banana peel micro cellulose (72% crystallinity) but greater than that obtained by Naranjo et al. in agave fibre (50.07%). The greater the crystallinity of a fibre, the better the thermal stability and properties it can provide in a polyblend to obtain biodegradable materials.

### 4. Conclusions

Cellulose microfibers were isolated from waste mango fibrous endocarp (Mangifera caesia Jack ex Wall), after which they were subjected to an acid-alkaline treatment and an ultrasonic treatment that helped homogenize their sizes. The acid-alkaline treatment helped decrease the size of the fibres and was also involved in the removal of lignin and cellulose. Ultrasonication process is a very effective way to depolymerize cellulose and that led to reduce the formation of cellulose.
of agglomeration and increase the dispersion of cellulose disintegrating into individual microfibers. We hope that with this simple chemical-ultrasonic treatment applied to cellulose will further stimulate interest in isolating microcellulose fibers from natural sources. The use of agricultural waste as a natural source of biopolymers has become an excellent option for innovation and the development of new products with potentially biodegradable properties by having a polymeric matrix formed by biopolymers and, their utilization can minimize negative environmental effects.

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