Extraction and Characterization of Fiber Treatment
Inula Viscosa Fibers as Potential Polymer Composite Reinforcement

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Abstract

This research aims to analyze the newly discovered cellulosic fiber from the bark of Dittrichia viscosa, or Inula viscosa (IV), and evaluate the effects of permanganate and alkali chemical treatments on the physical properties to improve the interfacial bonding between Inula viscosa reinforced polymer composites. These permanganate and alkali treatments are both efficacious in helping in reducing hydrophilicity and eliminating impurities from the fiber surface, which has been confirmed by scanning electron microscope (SEM) observation, thermogravimetric analysis (TGA), X-ray diffraction (XRD), and Fourier Transform Infrared Spectrometry (FTIR). Permanganate-treated IVFs have shown higher tensile strength and interfacial stress resistance (IFFS) than alkaline IVFs, and untreated IVFs by tensile and droplet tests. The different characteristics were studied and compared with other bark fibers already available. It is estimated that the fiber treatments will enable high-quality IVF-reinforced polymer composites for use in the industry.

1. Introduction

Societal needs for new environmentally friendly materials make scientists to develop materials from nature itself. The increased numbers of research that working on renewable and biodegradable materials to replace conventional materials. [1–4]. Researchers and scientists are currently focusing on green composite materials from plants fibers with good fiber yields, cost-effective without compromising mechanical properties [5–7]. This is justified because, on the one hand, natural fibers have the status of renewable and durable materials; on the other hand, they are abundant, light, of moderate resistance, high specific modulus compared to synthetic ones [8–10]. Composite materials reinforced by natural fibers have contributed significantly to the human race’s well-being and technological development. The construction industry, vehicle parts, domestic applications, aerospace, sports equipment, and others have been more inclined to use cellulosic fibers because of their benefits [11, 12].

The various sections of a plant, such as bark, root, seed, leaf can be obtained from the cellulosic fiber. The latter is one of the most available and inexpensive polymers on the planet due to its wide variety of industrial and common applications. When assessing natural fibers’ properties, natural fibers properties are also highly influenced by the environment’s growth, such as humidity, temperature, soil, and all affect fiber strength, density, and so on. Also, the fibers’ extraction (chemical, mechanical, or biological technique) leads to a variation of the properties.

Generally, the plants containing natural fibers have different regions and conditions. It depends on the plant’s age, place control and chemical fiber modification [10, 13]. The cellulose’s exceptional properties such as low spiral angle, small diameter, and continuous will enhance the properties of the composite. The outer layer of cellulose fiber is covered by non-cellulosic component (hemicelluloses, lignin, pectin, and wax).
The conditions of the natural fiber are based upon the strength of cellulose fiber. Higher tensile strength is achieved by the chemical composition of the cellulose fiber. Therefore, it is necessary to explore new natural fibers, with the required strength properties of fiber reinforcement. A wide range of diverse natural fibers are available that offer reinforcing effects to polymer composites. The mechanical efficiency of polymer matrix composite is primarily dependent on the interfacial bonding. The fibers may be changed in chemical or physical treatments [12, 14]. Also, natural fibers treated with mercerization, silane treatment, benzoylation, potassium permanganate, etc., have significant advantages, such as lower water absorption, increased surface roughness, and higher thermal stability, less amorphous content, improved crystallinity index, and increased crystallite size [15–17]. Natural fibers are obtained from various plants such as tree bark such as Azadirachta indica, Ceiba pentandra, Acacia Concinna, Grewia tilifolia, Acacia Nilatica L., Acacia leucophloea, ramie, and yam have been a focus of research for scientists as reinforcements of composite because of their specific properties and availability [18–22].

The future research will investigate a new bark fabric Dittrichia viscose L (Inula Viscosa) plant that has not been studied so far, despite its abundance in the Mediterranean region, Southeast Europe, and Asia West. In this study, two different chemical treatments, 3% alkaline and 3% permanganate, were adopted to modify the IV fibers samples of bark were harvested using an ecological technique. The physicochemical, morphological, mechanical and thermal properties of the IV fibers was investigated for the first time in this work. The fiber surface and fiber structure morphology were observed by using a binocular microscope and scanning electron microscopy (SEM).

The surface morphology of the new natural fiber was investigated using a binocular microscope and scanning electron microscopy (SEM). Infrared spectroscopy (FTIR) and X-ray diffraction (XRD) were employed to analyze the chemical and molecular structure of untreated and treated IV fibers. In the TGA analysis, thermal stability was conducted to analyze both the untreated and treated IV fibers. This new cellulosic fiber’s mechanical property was investigated using the single filament test according to the ASTMD3379 standard method. The interfacial shear strength (IFSS), post-debonding strength (τpost-debonding), and debonding behaviors were studied using a micromechanical droplet approach.

2. Materials And Method

2.1. Materials

Dittrichia viscose (L.) or Inula viscosa (IV) owes its name to the Greek “inæin” which means purged. This plant is known in Algeria as "Magraman". It is an undemanding plant in the Mediterranean region and widespread in northern Algeria. It grows in grassy places, in fields, and is also found along roadsides and paths. It belongs to the family Asteraceae, Class: Magnoliopsida; Genus: Inula Species: Inula viscosa. It is a wild, perennial, tap rooted, reasonably large (up to 1.50 m tall) plant. Its stems are fairly branched and have dense foliage. The leaves, alternate, elongate to lanceolate, are inserted directly on the stem, without
petioles. The whole plant is coated with glandular hairs that release an odoriferous and sticky resin, with a smell of camphor, Figure 1.

2.2. Fiber extraction from the plant

Fresh Inula viscosa plant was collected from Hodna region (Algeria) in April 2019. The IVs’ barks were submerged in a small water tap for five weeks to detach outer skin from the stalk by using biological retting technique, Figure 2 (a), [23, 24]. Next, the innermost layer was manually removed from the outer layer. The inner layer of the bark yielded fine fibers. The extracted fibers were washed in running water to get rid of unwanted contaminants that were left behind the fiber surface, then dried in an oven at 70 °C for 6 hours, Figure 2 (b).

2.3. Chemical treatments

Two different chemical treatments were used in this analysis. The 3% alkaline and 3% permanganate were used to alter the Inula viscosa fibers' surface and to foster a stronger bonding between these fibers and the resin.

2.3.1. Alkaline treatment

IVFs were immersed in distilled water with a 3% sodium hydroxide solution, for 3 hours, at RT. The fibers were immersed in water prior to having neutral pH reached. After rinsing, the fibers were dried for 48 hours before being characterized.

2.3.2. Permanganate treatment

For the permanganate treatment, the pretreated with alkaline fibers as in Section 2.3.1 were soaked in 3 % of Potassium permanganate (KMnO₄) solution for 3 minutes [25]. These fibers were washed and dried in the air.

2.4. Characterization

2.4.1. Scanning electron microscopy

The surface morphology of untreated raw fibers, 3% alkaline, 3% permanganate -treated Inula viscosa fibers were analyzed using a scanning electron microscope (SEM) machine Model HITACHI S-3400N. The SEM instrument was set up at an emission current of 58 μA. The working distance was set to 6.2 mm, and the acceleration voltage of 5.0 kV. The specimens were coated with a thin gold layer to avoid electrical charges during the examination.

2.4.2. Energy-dispersive X-ray spectroscopy analysis (EDX)

The EDX involves an analytical method was employed for the element analysis of a sample attached with SEM images. This approach allows elements from the whole element of periodic table to be detected
of untreated, 3% alkaline, 3% permanganate treated IVFs, except for H, He, and Li. Therefore, the Hydrogen can't be observed by this technique which represents the major constituents of natural fiber.

2.4.3. Fourier Transform Infrared Spectrometry (FTIR)

The FTIR analysis was carried out for untreated raw fiber, 3% alkaline, 3% permanganate treated Inula viscosa fibers by using an FTIR machine (SHIMADZU81001, Japan). The purpose of using FTIR analysis to distinguish the changes in functional groups on the fiber surfaces of IVs. All spectra were recorded in the range from 4000 cm\(^{-1}\) to 500 cm\(^{-1}\). The sample preparation for this testing, the fibers were chopped and were ground into powder form.

2.4.4. X-ray diffraction analysis (XRD)

The degree of crystallinity and crystalline size of cellulosic fibers IVs were tested by using an X-ray diffraction test [1]. The XRD testing is non-destructive and rapid testing. The tests were carried out for the untreated raw fiber, 3% alkaline and 3% permanganate-treated of IV fibers. In this testing, the Copper was used as the anode material with monochromatic Cu K\(\alpha\) radiation with a wavelength of 0.154 nm and a current of 30 mA and voltage of 30 kV. The continuous scanning mode, 2\(\theta\) range varying from 5º to 60º with a step size of 0.02º at 25°C. The Segal empirical technique and Scherer’s equation were used to determine, the degree of crystallinity and crystallite size of the untreated raw fiber, 3% alkaline and 3% permanganate treated- Inula Viscosa fiber as displayed in Equation (1) and (2).

\[
CI(\%) = (I_c - I_{am})/I_c \quad \text{Eq. (1).
}
\]

Where, \(I_c\) represents the maximum intensity of the crystalline phase peak and \(I_{am}\) represents the amorphous phase's intensity in the cellulose present in the fiber [26].

\[
CS = K\lambda/\beta \cos \theta \quad \text{Eq. (2).
}
\]

Where \(K = 0.89\) is Scherer's constant, \(\beta\) is the peak's full-width at half-maximum, \(\lambda\) is the wavelength of the radiation, and \(\theta\) is the corresponding Bragg angle [27].

2.4.5. Thermogravimetric analysis (TGA)

The thermal stability and thermal decomposition of untreated raw fiber, 3% alkaline and 3% permanganate-treated IV fibers were obtained by using Thermogravimetric analysis by using GA machine Model (TGA Q 500 TA Instrument, USA). The specimens were measured under room temperature ranging from 30 °C to 600 °C.

2.4.6. Density measurement
The density of the untreated raw ber, 3% alkaline, 3% permanganate-treated IVFs was calculated by a liquid pycnometer for solids with immersion liquid known as the methanol (methanol, \( \rho = 791 \text{ kg/m}^3 \) at 21 °C). The liquid pycnometer was based on direct measurement of specimen's volume. The accuracy of electronic weighing machine was checked at 0.00001 g before the weight measurement of the fiber. The sample of IVFs were cut with the length of 1 mm to suit in the pycnometer [28] and were placed in an oven at 60 °C until the moisture content was reduced to below 5 % before testing [29]. The densities of raw and treated IVFs (\( \rho_{\text{IVFs}} \)) were calculated through the following equation (3) [28]

\[
\rho_{\text{IVFs}} = \frac{m_2 - m_1}{(m_2 - m_1)(m_4 - m_2)} \rho_\text{E} \quad \ldots \quad \text{Eq. (3)}.
\]

Where, \( \rho_{\text{IVF}} \) is the density of IVFs (kg/m\(^3\)), \( \rho_\text{E} \) is the density of methanol (kg/m\(^3\)), \( m_1 \) is the mass of empty pycnometer (kg), \( m_2 \) is the mass of pycnometer with fibers (kg), \( m_3 \) is the mass of pycnometer with methanol (kg), and \( m_4 \) is the mass of pycnometer with fibers and methanol (kg).

### 2.4.7. Diameter measurement

A diameter test was carried out on a single fiber of untreated raw, 3% alkaline at 3% permanganate-treated at room temperature by using Olympus, BX51 optical microscope (Japan) with a magnification of 5X, three replicates were measured for each untreated and treated IV fiber by placing the fiber over a glass plate and taping down its end to obtain a clear image [30].

### 2.4.8. Tensile test

Tensile testing of untreated raw fiber, 3% alkaline and 3% permanganate-treated IVFs were evaluated by Instron universal testing machine (UTM) at a speed of 1mm/min according to ASTM D 30393. The fibers were then fixed and glued to the tab shape, which was performed with the gauge length of 30 mm fibers then inspected to remove the crack, Figure 3. The tensile strength of fibers was calculated from the following equation (4) [31].

\[
\tau = \frac{F}{A} \quad \ldots \quad \text{Eq. (4)}.
\]

Where, \( T \) tensile strength in Pa, \( F \) force to failure in N, \( A \) average fiber area in m\(^2\).

### 2.4.9. Droplet test

In the droplet test, a thin metal rod was chosen to place microdroplets of epoxy resin on single fibers that were fixed to a paper frame and left to solidify. Before testing, the paper frame was cut away when a sample was connected to the tensile testing machine equipped of 5kN (INSTRON UTM). The interfacial shear strength (IFSS) which evaluate the adhesion in specific fiber matrix system according to the following equation (5) [32]

\[
\tau = \frac{F}{\pi DL} \quad \ldots \quad \text{Eq. (5)}.
\]
Where τ is the interfacial shear strength (MPa), $F_{\text{max}}$ is the maximum pull-out force; D is the fiber diameter and L is the embedded length. Both parameters were measured using an optical microscope equipped with a high resolution before testing, Figure 4 (a). The schematics of the microdroplet test are shown in Figure 4 (b).

The ten samples for each untreated raw fiber, 3% alkaline and 3% permanganate-treated, were tested. The fiber diameter measurement was obtained from the point nearest to the droplet to both sides’ fiber contacts.

3. Results And Discussion

3.1. Density measurement

Density measurement of the natural fibers is essential to evaluate the potential density of composite materials that use certain fibers. Various factors affect the density of cellulosic fibers such as the soil plant conditions, humidity present, fiber’s age, the fiber extraction process, etc.

The density value for 3 % alkali and 3% permanganate treated was IVFs (1102 ± 33.42 kg/m$^3$), (1154 ± 13.22 kg/m$^3$) which is a slight density increment compared to untreated IVFs (1040 ± 10.54 kg/m$^3$). This is possibly attributed to the pores and voids in the fiber surface consisted grafted molecules during chemical treatments [33-35]. From this testing, it was noted that the density of novel fiber IV was lower than other natural fiber such as jute (14800 kg/m$^3$)[36], sisal (1500 kg/m$^3$)[37], Alfa and Sabra fibers (1.40 g·cm$^{-3}$ ± 0.02)[34], banana (1350 kg/m$^3$) [38], Cyperus pangorei (1102 kg/m$^3$)[39]. Thus, IVs could be the candidate fiber reinforcement for composite lightweight.

3.2. Diameter measurement

The average diameters of the untreated raw fiber, 3% alkaline and 3% permanganate treated IVs fiber as shown in Figure 5. The diameter of untreated raw fiber (93.50 ± 2.75 µm), 3% alkaline (80.41 ± 1.63 µm), and 3% permanganate treated (78.60 ± 1.82 µm), respectively.

3.3. Scanning electron microscopy (SEM)

The surface morphology of fibers can be determined using the Scanning Electron Microscopy (SEM) method to examine the surface morphology of fibers. Scanning electron micrographs of untreated, alkaline, and permanganate-treated fibers are presented in Figure 6. Figure 6 (a) shows the SEM micrograph of untreated fibers, the surface of IVs showed absence of impurities such as wax and grease, and internal fibrils [40]. At higher magnification, Figure 6 (b) shows that the removed waxes and oils from the fibers' surface were removed by 3% alkaline treatment, thus enable surface roughness on the fiber surface [41, 42]. The alkaline treatment showed the differences compared to raw fiber. In this case, the fiber's surface was smoother than the raw fiber due to removing surface impurities. For permanganate-treated fibers, Figure 6 (c), it can be observed that the fiber became cleaner, with a rougher surface, as
impurities were removed from the surface of the fiber [35]. This rough surface may improve interfacial bonding when IV fibers are used as reinforcing materials.

3.4. Energy-dispersive X-ray spectroscopy analysis (EDX)

The EDX technique relies on the sample's major interaction and the X-ray excitation source. The qualitative findings on the quantity of major elements (carbon, oxygen, calcium, manganese, etc.) provided by the fiber surface of untreated and treated IVFs are shown, Figure 7. In addition, the presence of C and O elements tends to be the most prominent in the EDX continuum since they are the critical components of the architectures of natural fibers [40].

The EDX study of both untreated and handled IVFs in terms of atomic percentage and weight is provided in Table II. It has been found that untreated IVF comprises almost 97.56 percent carbon weight, however, the carbon proportion is decreased to 97.32 percent and 75.08 percent for alkaline and potassium permanganate treatments, respectively, since chemical treatments could have eliminated the outer layer of the treated fiber [42]. This would be due to the IVF’s more non-cellulosic components.

3.5. Thermogravimetric analysis

The thermal properties of untreated and treated IVF fibers are analyzed using TGA. Figure 8 demonstrate the TGA and DTG curves of these lignocellulosic fibers, an essential feature in biocomposite based on these fibers [30]. The first stage of decomposition was similar to untreated raw and chemically treated (IV) fibers, indicating the weight loss process. At the range between 30 °C and 125°C, a small weight loss (6.15%) was demonstrated, which is agreed by several authors [43, 44]. It was shown that the first curve trend in DTG curves were decline in DTG curves, Figure 8 (b), it is proven that the water evaporation after 3% alkaline, 3% permanganate treatment. The same observation was found in Figure 8 (a). This is due to the reduction of the cellulose fiber's hydrophilic nature when the chemical treatment was employed on the IV fiber as the acquired for fiber reinforced polymer composites. Thus, the reduction moisture loss percentage in the both treated fibers could be the higher of crystallinity properties of the IV fiber [14]. When the temperature rises up to °C, no significant peak is observed in the DTG curve and the similar was agreement with others work [45]. Beyond this temperature, thermal stability is decreasing and the fiber decomposition is happened, Figure 8 (a).

The second stage decomposition at 190°C until 290°C corresponds to hemicellulose decomposition and the third decomposition at 290°C-400°C correspond to cellulose and lignin decomposition. It was reported by another study [46] the least thermally stable was hemicellulose, the intermediate was cellulose and the lignin was the most resistant.

The untreated IV fibers started to degrade at around 200°C as shown in the degradation profile. The first degradation peat at 285°C correspond to the depolymerization of hemicellulose, pectin and glycosidic linkages of cellulose by 18.71% of weight loss. The 3% alkaline and 3% permanganate treated IVs. the peak was not visible, proving the complete removal of hemicellulose from the fiber. The major second
peak was observed at 365.51°C due to degradation α-cellulose by 80.87% weight loss for untreated raw fiber [47], whereas 3% alkaline treatment and 3% permanganate-treated at 363°C and 350 with 67.7% and 62% weight loss respectively [48]. It can be noted that surface modification by both treatments reduced the thermal stability properties of the IV fibers, our findings are consistent with previous works [49, 50]. From ambient to higher temperatures at 600°C, the lignin degradation whose structure is a complex composition of aromatic rings with different branches may occur at a very low weight loss [51].

3.6. Fourier transforms infrared spectroscopy (FTIR)

The comparison of FTIR spectra of untreated raw fiber and chemically treated with 3% alkaline and 3% permanganate of Inula Viscosa (IV) fibers presented in Figure 9, shows absorption bands of chemical groups characteristic of lignocellulosic fiber compounds. The main characteristics of the spectrum of the untreated IVs at the peaks 3329, 2919, 2851, 1731, 1638, 1592, 1423, 1325, 1239, and 1028 cm\(^{-1}\) are α-cellulose, hemicelluloses, lignin, pectin, and water molecules contents. From the large absorption band observed around 3329 cm\(^{-1}\) is linked to OH and CH stretching of cellulose [52]. The strong adsorption peaks depicted at 2919 cm\(^{-1}\) and 2851 cm\(^{-1}\) are related to C-H stretching vibrations from CH and CH2 in cellulose and hemicellulose, respectively [53]. An observable peak around 1731 cm\(^{-1}\) corresponds to the C=O stretching of hemicelluloses [21, 54]. The band around 1638 cm\(^{-1}\) was related to the O–H bending of water absorbed into cellulose fiber structure The peak around 1592 cm\(^{-1}\) corresponds to the aromatic ring C=C of the phenyl propane group in lignin [55]. Also, a small peak near 1423 cm\(^{-1}\) belongs to the aromatic skeletal vibrations and ring breathing with C–O stretching in lignin [56]. The peak observed at 1325 cm\(^{-1}\) is attributed to C-H and C-O groups' bending vibration of the aromatic ring in polysaccharides [57]. Additionally, A small peak around 1239 cm\(^{-1}\) belongs to the acetyl group's C–O stretching in hemicelluloses [54]. A visible peak at 1028 cm\(^{-1}\) is related to the stretching of C–O groups of cellulose [43]. The fibers' FTIR spectra confirm the compositional changes in permanganate and alkali-treated fibers, Figure 9. The two peaks at 1239 cm\(^{-1}\) and 1731 cm\(^{-1}\) were observed in FTIR of untreated fibers, which correspond to hemicelluloses, disappeared in the spectrum of the permanganate and alkali-treated fibers. This result could be explained by the elimination of the residual hemicellulosic materials after the treatment. The removal of an important amount of lignin by the chemical treatment can be noticed through the disappearance of the peaks located at about 1325 cm\(^{-1}\) and 1423 cm\(^{-1}\). It is to demonstrate that the removal of hemicelluloses and lignin from the treated (IV) supports the results of the chemical analysis.

3.7. XRD analysis

Figure 10 illustrates the XRD pattern of the 3% alkaline, 3 % permanganate treated and raw fibers specimen of IV and the corresponding planes involved. From the Figure 10, it was shown that each specimen showed two peaks, respectively. For 3% permanganate and 3 % alkaline treatment, the first peak represents the amorphous peak demonstrated at 2θ = 15.96°, 16.45° and 16.06° at lattice plane (110). respectively. While for the second-high intensity peak the 3% alkaline, 3% permanganate and
untreated IV fibers represents the crystalline peak observed at $2\theta = 22.48^\circ$, $22.52^\circ$ and $22.08^\circ$ respectively, belongs to the (200) plane of cellulose [40]. The value of crystallinity index (CI) was higher for 3% permanganate at 55.93% followed by 3% alkaline treatment at 54.25% and untreated raw fiber at 51.63%. It was shown that chemically treated with alkaline and treatment were improved compared to untreated.

The increase in CI with permanganate and alkali treatments is related to the loosening of cellulosic chains resulting in the disappearance of excess amorphous constituents, such as lignin, hemicellulose, etc. [58]. This result was agreed with SEM morphology the impurities of fiber was removed with chemical treatment. Furthermore, the crystallite size (CS) of the untreated raw, the 3% alkaline and the 3% permanganate-treated of the IVFs were obtained as 0.8 nm, 1.85 nm and 2.0 nm, respectively. The crystallite size may reduce the chemical activity and the water absorption capacity of the fibers.

3.8. Tensile test

The study of mechanical properties of natural fiber reinforced polymer composites is important to understand their potential for various structural applications. Figure 11 (a), depicts the impact of untreated raw, 3% NaOH and 3% permanganate-treated IVFs on the tensile strength of IVFs. From the graph, the tensile strength trend was increasing for 3% permanganate, followed by 3% alkaline treatment compared to untreated raw fiber with the value of 195.88 MPa, 173.047 MPa and 163.60 MPa, respectively. The increased tensile strength of chemically treated IVs fiber due to the elimination of impurities from the IV fiber surface. Previous research shows the values of tensile strength from plants fiber was approximate with IVs fiber such as sisal (274-526 MPa) [59], date palm (170-275 MPa) [60], Lygeum Spartum (LS) (113 MPa) [43], pineapple leaf fiber (PALF), and Arundo Donax 248 MPa [30]. The strain rate of untreated fibers and alkali and permanganate treated IVFs is 1.172%, 1.439 %, and 1.562% respectively, which directly affects the micro fibrillation angle of the IVFs Figure 11 (b). The Young modulus of raw natural fibers and fibers treated with alkali and permanganate is 11.407 GPa, 11.596 GPa, and 12.25 GPa, respectively, Figure 11 (c). The tensile modulus of IV fibers with 3% alkaline and 3% permanganate was higher than untreated raw fiber. The values are quite approaching to other plant fiber such as artichoke (11.6 GPa), sisal (9.4 - 22 GPa) and bamboo (11-17 GPa) [30].

3.9. Droplet test

This work evaluated the bonding strength between the IV fiber and epoxy resin by a droplet test. As shown in Figure 12, the IFFS interfacial shear strength was obtained from the test results. Unlike other fiber pull-offs, this technique allows the average shear stress to be calculated once the fiber is peeled off by force ($F_d$).

It should be noted that the apparent adhesive force measured with micro bonding tests varies greatly. The variability causes this in the IV fiber dimension that have different diameters. The mechanism of droplet test showed that when the increase of load to pull out the IV fiber, the easily IVs fiber to break. From this result, the higher interfacial shear strength was permanganate-treated fiber with 4.50 MPa, followed by alkaline treatment, and untreated of IVs fiber with 3.36 MPa and 2.93 MPa, respectively. The
increased value of t IFFS for permanganate was 53.58% and alkaline treatment by 11.26% compared to untreated IVs fiber. The data obtained is comparable to the IFFS of flax, hemp, and sisal [61]. The adhesion between the epoxy resin and the Inula Viscosa fiber was improved by permanganate treatment and alkalization. According to the results, the adhesion between the treated Inula Viscosa fiber/epoxy was better than the adhesion bonding between the untreated Inula Viscosa fiber/epoxy.

**Conclusions**

In this study, the untreated IVs, 3% alkaline treated IVs, and 3% permanganate treated IVs were evaluated for their physical, morphological, crystalline structure, thermal degradation, mechanical and micro-mechanical properties. The conclusions concluded from the experiments.

1. The density of untreated raw IVF and alkali and permanganate treatment IVF is lower than that other natural fibers have.
2. The decrease in diameter for alkaline, permanganate-treated fiber for IVs fiber due to impurities surface removal compared to untreated raw IVs fiber.
3. The XRD analysis confirms that the treated IVFs was higher strength and more resistant to humidity than untreated raw ones obtained from CI and CS values.
4. The reduce hemicellulose, lignin, and wax content of IVs fiber by chemical treatment was confirmed by FTIR and were agreed with surface morphology in SEM analysis.
5. The thermal stability of untreated and chemically treated with alkaline and permanganate of IV fiber can be withstanding up to 190°C.

The tensile properties of IVs fiber such as tensile strength, Young’s modulus, and strain to break depicted the improved strength by chemically treated IVFs due to the higher cellulose content and CI values. The highest IFSS properties were permanganate, followed by alkaline treatment and untreated raw fiber.

This finding showed that the importance of chemical fiber treatment leads to the natural fiber reinforced polymer composites’ enhanced properties. Overall, the chemical modification of IVs fiber exhibited better mechanical properties than untreated. The is due to that the improved adhesion and polar interaction between the fiber and polymer interfaces.

**Declarations**

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**Tables**

**Table I. Abbreviations of samples names.**

| Names of samples                          | Abbreviations |
|-------------------------------------------|---------------|
| Raw Inula Viscosa fiber                   | R-IVF         |
| Alkali treated Inula Viscosa fiber        | A-IVF         |
| Permanganate treated Inula Viscosa fiber  | P-IVF         |

**Table II. (a) The elements composition found on the surface of R-IVF.**
| Element | Weight % | Atom % | Formula | Compnd % |
|---------|----------|--------|---------|----------|
| C       | 97.56    | 98.78  | C       | 97.56    |
| O       | 0.98     | 0.75   | —       |          |
| S       | 0.36     | 0.14   | SO₃     | 0.91     |
| Ca      | 1.10     | 0.33   | CaO     | 1.54     |
| **Total** | 100.00  | 100.00 | 100.00  |          |

**Table II. (b)** The elements composition found on the surface of A-IVF

| Element | Weight % | Atom % | Formula | Compnd % |
|---------|----------|--------|---------|----------|
| C       | 97.32    | 98.84  | C       | 97.32    |
| O       | 0.76     | 0.58   | —       |          |
| Ca      | 1.91     | 0.58   | CaO     | 2.68     |
| **Total** | 100.00  | 100.00 | 100.00  |          |

**Table II. (c)** The elements composition found on the surface of P-IVF

| Element | Weight % | Atom % | Formula  | Compnd % |
|---------|----------|--------|----------|----------|
| C       | 75.08    | 91.39  | C        | 75.08    |
| O       | 2.22     | 2.03   | —        |          |
| Mg      | 0.60     | 0.36   | MgO      | 1.00     |
| Ca      | 3.44     | 1.26   | CaO      | 4.82     |
| Mn      | 17.61    | 4.69   | Mn       | 17.61    |
| Fe      | 1.04     | 0.27   | Fe₂O₃    | 1.49     |
| **Total** | 100.00  | 100.00 | 100.00  |          |