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Splitting and authentication of the newest retrieved cellulose-rich organic fiber from the exterior layer of Bangladeshi palmyra seed sprouts

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The upward trajectory of plant-based cellulosic fiber originating from renewable sources is crucial to visualizing a sustainable future. This article reports a freshly developed and distinctive natural fiber derived from palmyra seed sprout fibers (PSSF) by employing hot water retting. The hygroscopic behavior (moisture regain and content) density, bundle fiber strength, burning behavior, and chemical composition of the fiber are determined using ASTM D1909, ASTM D2654, ASTM D891-18, ASTM D1445, ASTM D1230-22, and TAPPI standards, respectively. Crystallinity, fiber configuration, biological groups, flame behavior, and temperature responsiveness are determined using XRD, SEM, FTIR, burn test, and TGA tests. The reported cellulose-rich textile fiber that contains 64% cellulose with average moisture recovery and content percentages of 14.38% and 12.56%, respectively. The envisioned fiber has an average tensile strength of 11.05 g tex⁻¹, a breaking extension of 1.8 mm, and a crystallinity of 38%. The highest temperature at which the fiber begins to deteriorate is 474 °C. This noble fiber can be utilized to create biological materials, cellulose nanoparticles, composites reinforced with fibers, and more.

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1. Introduction

Sustainable fibers are those that are collected and separated in ways that mitigate their negative impact on the environment and the community.1 These fibers are frequently produced by employing conscientious manufacturing practices that provide acceptable working conditions and social wellbeing.2 When it comes to resource, energy, and pollutant consumption, they typically have little environmental impact.3 Sustainable natural fibers are typically favored over synthetic alternatives because they are biodegradable, have a smaller carbon impact, and may be used in circular economy activities. 1,3 Considering the continual growth in the popularity of natural cellulosic fiber in recent years, both academic and commercial researchers are increasingly studying innovative natural fibers and their possible uses.4,5 They are particularly fascinating because of their unique features, which make them suitable for a wide range of sophisticated fibrous applications, such as reinforcement in composites, textiles, cellulose nanomaterials, activated or conductive carbon or nanotubes, biomaterials, and so on.^{6,7} Natural fibers have specific attributes such as tremendous strength, excellent heat resistance, biodegradability, recyclability, low cost, wide availability, and zero toxicity.8 Flax, hemp, areca, cotton, bagasse-jute, kapok, luffa, maize, coir, luffa,

pineapple, and other materials are known as natural fibers in both traditional and modern contexts. 9,10 Organic cotton, hemp, and linen are among the sustainable natural fibers that are used to make clothing, textiles, and home decor. Since these fibers are grown without the use of harmful pesticides, genetically modified organisms (GMOs), or extended amounts of water consumption, they are more environmentally friendly than traditional fibers. 4,5,11 Their life cycle has less of an impact on the environment because they are also biodegradable. Plasticbased materials in packaging can be replaced by these natural fibers.12,13 Jute, a naturally occurring fiber derived from the jute plant, may be used to make biodegradable sacks, bags, and packaging products.^{1,2} Additionally, these fibers can be used in building construction materials as well as reinforcement materials. For example, bamboo is a naturally occurring fiber that grows swiftly and may be used to make furniture, wall panels, and flooring.12,14

Worldwide, there are over 5000 different types of date palms, demonstrating the date palms' biodiversity. Many more varieties are found in other countries, with botanical descriptions indicating that there are around 1000 cultivars in Algeria, 400 in Iran, 370 in Iraq, 250 in Tunisia, 244 in Morocco, and 400 in Sudan. With fruit measuring up to 100 mm by 40 mm, date palm trees (*Phoenix dactylifera* L.) are the tallest species of *Phoenix* trees, reaching heights of over 30 meters. The fruits are highly palatable and nourishing. Date palms may adapt to a variety of situations thanks to their unique features. They are not erinaceous, although date palm trees may grow well in sand. Their roots have air gaps, which allows them to grow

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successfully in areas where soil water is near the surface. Date palm trees thrive best in higher quality soil and water, yet they can tolerate salinity rather well. Date palms are not xerophyte plants; although their leaves can withstand hot, dry weather, they still need lots of water. The date palm tree produces a large number of offshoots at its base. Something that distinguishes the date palm tree are the numerous offshoots that grow at the base of its trunk. The date palm tree's stalk is covered with dependably dull leaf bottoms. On top, it features a magnificent display of tall, pinnately divided leaves with needle-sharp fronds. There are usually 10 to 20 new leaves produced annually. The date palm's leaves are supported by a thick, fibrous, cylindrical sheath that reticulates around the base of the leaves. Together, they form a tight enclosure around the terminal bud. The same palm tree is near the surface.

The intended objective of this research is to collect fibers from the outer layer of PSSF and determine and analyze their quality in order to present them as a renewable fiber derived from nature.



Fig. 1 Raw palmyra seed with its sprouts.

No researches related to this topic were discovered, proving the research's originality. These features are then compared to those of other lignocellulosic fibers found in nature like jute, cotton, flax, Napier grass, *etc.*²⁰ This comparative study sought to determine whether these materials are suitable for use as reinforcement in thermosetting and thermoplastic polymers for a variety of applications such as food packaging, medical devices, and structural engineering components.

2. Materials and methods

2.1 Palmyra seed sprout collection

Palmyra sprouts are collected from a village near Kapasia, Gazipur, Dhaka. The seeds are kept on average for 4 to 6 months depending on the sprout's fiber maturity. The tested samples were kept for more than 5 months after they were collected. Fig. 1 shows the raw seed as well as the placement of the seed sprouts.

2.2 Fiber extraction process

After collecting the palmyra seeds the sprouts were cut and separated. The PSSF were boiled in water at 100 °C for 1 hour and then the samples were cooled until the room temperature was reached. Subsequently the boiled sprouts were pounded with a medium weight object; as a consequence, the peels became soft and cleaned with normal water 4 to 5 times. Lastly the cleaned samples were kept at room temperature for 24 hours and they dried. The extraction flowchart is shown below in Fig. 2.

2.3 Methods of fiber characterization

2.3.1 FTIR analysis. To identify the active chemical component of the fiber, a Fourier Transform Infrared (FTIR)



Fig. 2 Flowchart of PSSF extraction.

spectrophotometer (FT-IR 8400S, Shimadzu Corporation, Japan) is utilized. To measure the infrared of the fibers, they were ground and mixed with potassium bromide (KBr), which is transparent. The FTIR spectrometer was operating in absorbance mode at a room temperature of 30 °C and relative humidity of 65%. Its scan rate was 32 and its resolution was 2 percent in the 400–4000 cm⁻¹ wavenumber area.

2.3.2 Moisture regain and content analysis. To ascertain the moisture recovery and content, respectively, the ASTM D1909 (ref. 21) and ASTM D 2654 (ref. 21) approach was employed. Under standard fiber testing parameters of 20 °C and 65% relative humidity, a 5 g sample was collected. The weighed fiber samples were placed for a predetermined amount of time at 105 °C in an air oven. The proportion of fiber with moisture content was calculated using the following formula.

$$\frac{W}{D} \times 100\% = MR\%$$

$$\frac{W}{W+D} \times 100\% = \text{MC}\%$$

In those equations, W = weight of water in the samples, D = oven dry weight of the samples, MR = value of moisture regain, and MC = value of the content of moisture.

- 2.3.3 Fiber strength analysis. A stelometer is used to evaluate the strength of the bundle fiber. The test technique belongs to ASTM D1445.²² The well-combed fiber samples from the ten fibers have been mounted on a pair of small clamps with gauge lengths of zero. After placing the fiber clamp at the top of the pendulum, tension is applied to rupture the fibers. The approach collects information on fiber strength when broken and extensibility.
- 2.3.4 XRD analysis. The fiber's crystallinity was evaluated using an X-ray diffractometer (SmartLab SE, Rigaku, Japan) set to 50 mA of current and 40 kV of voltage. The fiber pellet sample was subjected to an X-ray diffraction study. The diffraction intensity of the CuKa (1.54 Å) radiation was measured with a step width of 0.01° and a scanning speed of 15° min $^{-1}$ between 10° and 40° (2 β angle range). The peak height method for determining the crystallinity index (CI) in the cellulosic materials was developed using the preceding Segal empirical equation. 23

$$CI = \frac{I_{200} - I_{am}}{I_{200}}$$

2.3.5 TGA along with DSC analysis. A thermal analyzer was used to perform DSC analysis on the fibers (New Castle, DE, USA; TA Instruments Trios V5.1.1.46572). Fiber samples weighing 10 mg were heated at a rate of 10 °C min⁻¹ between 10 °C and 450 °C. The nitrogen environment used for the experiment was being purged at a rate of 100 mL min⁻¹. The thermal stability of the fiber was assessed using a thermogravimetric analyzer (TA Instruments Trios V5.1.1.46572, New Castle, DE, USA) and an aluminum pan at a heating rate of 10 °C min⁻¹

and a flow rate of 100 mL min $^{-1}$. In this experiment, 10.499 mg of dry material was used. The test chamber temperature of the thermal analyzer was gradually increased at a rate of 10 $^{\circ}$ C min $^{-1}$, from 33 $^{\circ}$ C to 477 $^{\circ}$ C.

- 2.3.6 SEM analysis. In order to depict the morphological properties of fiber interactions 500× magnification, 20 mm working distance, 10 kV propelling voltage, and secondary electron diagnosis mode were the parameters used to capture the SEM picture. The surface morphologies of fiber samples were investigated under the same microscope both before and after chemical treatments. The machine was identified by the model number HITACHI SU1510.
- 2.3.7 Chemical composition analysis. The cellulose, hemicellulose, lignin, extractive, and ash contents of the fiber that was removed from the PSSF are measured using TAPPI standard test methods.²⁴ The extractive percentage was determined using the Soxhlet extraction technique with an ethanol and toluene solvent solution (OM-88 TAPPI T204). Extractive free fibers were treated with NaOH solution to assess the cellulose content of the holo-cellulose that was created using NaClO₂ solution (TAPPI T203 om-93). The Klason method (TAPPI T211 om-83) was utilized to determine the amount of lignin applied. The evaluation of the ash content was conducted using the TAPPI (T211 os-76) test method.²⁴ Two replications were made. The average is taken into account throughout each compositional analysis.
- **2.3.8 Fiber density.** The bulk density of the strand was evaluated using a pycnometer and the water submersion method according to the test standard ASTM D891-18.²⁵ The density was acquired employing the equation indicated below.

$$ho_{ ext{Palmyra}} = rac{m_2 - m_1}{(m_3 - m_1) - m_4 - m_2}
ho_{ ext{w}}$$

In this instance, $\rho_{\rm w}$ denotes the density of water (0.997 g cm⁻³ at 25), $\rho_{\rm Palmyra}$ represents the density of fiber in g cm⁻³, and m_1 represents the mass of the basic pycnometer, m_2 the mass of the pycnometer along with fiber, m_3 the mass of the pycnometer filled with water, and m_4 the mass of the pycnometer filled with both fiber and water.

2.3.9 Burn test. A crucial resource of data on fiber verification, chemical formula, thermal attributes, safety, inspection of quality, and environmental effect is the textile fiber burn test, a procedure endorsed by scientific studies. ASTM D 1230-22 standard was adapted for the burn test of the extracted fiber. The fibers were held at 45° angle in front of a standard flame and the fibers ignite within a second.

3. Results and discussion

3.1 FTIR analysis

An infrared (IR) spectrum acquired by Fourier Transform Infrared (FTIR) spectroscopy is shown in Fig. 3, which shows transmittance (%) as a function of wavenumber (cm⁻¹). There are many significant absorption bands in the spectrum, which correspond to distinct molecular vibrations. Among the notable peaks are those located at 3390 cm⁻¹, 2848 cm⁻¹, 2506 cm⁻¹,

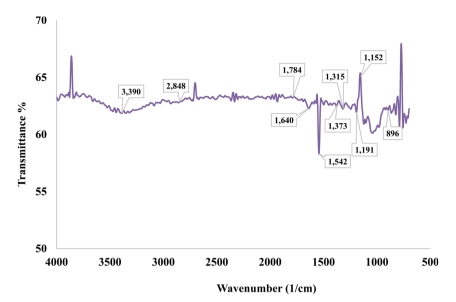


Fig. 3 FTIR spectrum of PSSF.

 $1934~\rm cm^{-1}$, $1640~\rm cm^{-1}$, $1315~\rm cm^{-1}$, $1487~\rm cm^{-1}$, $1542~\rm cm^{-1}$, $1373~\rm cm^{-1}$, $1191~\rm cm^{-1}$, $1152~\rm cm^{-1}$ and $896~\rm cm^{-1}$. The large peak at around $3390~\rm cm^{-1}$ is usually linked to O–H stretching vibrations, which frequently signify the existence of water or hydroxyl groups.

Fig. 3 demonstrates the FTIR spectrum of PSSF. The fiber's peak regions of the corresponding functional group specifies are shown in Table 1.

In the context of cellulose and hemicellulose, the C-H stretching vibration peak is located at 2848 cm⁻¹. ²⁸ Numerous authors state that the peaks at 1786 cm⁻¹ correlate to the carbonyl groups (C=O) of lignin and hemicellulose. ^{9,31,32,34} At around 1640 cm⁻¹, the water absorption of natural cellulose fiber peaked significantly. ²⁹ A signal with incredibly low intensity at 1542 cm⁻¹ indicated the C=C groups of lignin. A correlation was found between the peak at 1373 cm⁻¹ and the C-H bending vibration. The peak occurs at 1315 cm⁻¹ due to O-H bending. ³¹ The C-O stretching vibration of the acetyl group in lignin was measured at 1152 cm⁻¹. Peaks in the concentration of cellulose and hemicellulose in the fingerprint area,

ranging from 1090 to 1200 cm $^{-1}$, were attributed to C–O–C vibration. It was believed that C–OH vibration was responsible for the peak at 1038 cm $^{-1}$, and β -glycoside linkage was responsible for the peak at 896 cm $^{-1}$.28,31,34 This fiber's high cellulose content was verified by the FTIR peak patterns.30 The C–O stretching vibration of the acetyl group in lignin was measured at 1241 cm $^{-1}$. The cause of the cellulose and hemicellulose peaks at around 1100–1200 cm $^{-1}$ is C–O–C vibration.3

3.2 Moisture regain and content analysis

A fiber's composition was greatly influenced by the amount of water it contained. A textile's ability to hold onto body heat under various climate conditions has a significant impact on how comfortable it may be. Moisture control is an essential component of performance in this regard. Textile properties like elasticity, friction, fiber diameter, and tensile strength are all affected by changes in moisture content. A textile may become weaker, more brittle, and fragile when its equilibrium relative humidity decreases. This moisture loss to the

 Table 1
 Chemical stretching distributions along with peak locations for PSSF

Peak locations		
wavenumber (cm ⁻¹)	Allocations	Reference
3390	Wide peak for the -OH bond's H-bonded stretching vibration	27
2848	Vibration of C-H stretching for cellulose and hemicellulose	28
1784	Carbonyl groups (C=O) present in lignin and hemicellulose	9
1640	Significant peak for natural cellulose's water absorption	29
1542	Low peak intensity for lignin's C=C groups	30
1446	Bending vibration of CH ₂ groups in cellulosic matter	9
1373	C-H oscillation	31
1315	As a result of the O-H bending	32 and 33
1152	C-O stretching and acetyl group vibration in lignin	34
1038	C-OH oscillation	3 and 32
896	β-Glycoside bonding	9

Table 2 Assessing the moisture regain and content% for PSSF

Sample no.	Amount of sample $(W + D)$ (g)	Amount of oven dry sample (g) (D)		Moisture regain (MR%)	Average moisture regain (%)	Moisture content (MC%)	Average moisture content (%)
Sample 1	1	0.8693	0.1307	15.03	14.38%	13.07	12.56%
Sample 2	1	0.8865	0.1135	12.80		11.35	
Sample 3	1	0.8742	0.1258	14.39		12.58	
Sample 4	1	0.8712	0.1288	14.78		12.88	
Sample 5	1	0.8703	0.1297	14.90		12.97	
Standard de	viation			0.9150		0.7061	
Coefficient o	of variation			6.37%		5.63%	

environment is minimized by preserving air humidity while processing the fibers.³⁵ The moisture content% and moisture recovery chart for the samples are shown in Table 2.

The study reveals that the average moisture content and recovery are 12.56% and 14.38%, respectively. Five distinct samples weighing one gram apiece from the isolated fibers are used to get the results. The two findings' standard deviations, 0.7061 and 0.915, respectively, indicated that the fiber samples' moisture content and moisture regain values are quite similar. The error values for the five tested samples' moisture content% and moisture recovery% are shown in the bar diagram in Fig. 4.

3.3 Fiber strength analysis

Fig. 5 below depicts the tensile strength of the extracted fiber. The graph's *y*-axis is labeled "strength (g tex⁻¹)" and the *x*-axis is labeled "extension (mm)." The orange curve demonstrates the tensile behavior of palmyra seed sprout fiber. This fiber's strength increases steadily until it reaches its peak, after which it gradually declines. The average strength of PSSF is 11.5 g tex⁻¹ along with 1.8 mm extension. This value is near to that of jute fiber and lower than that of the other natural fiber extracted from *Mikania micrantha*. ^{3,36} *Mikania micrantha* fiber seems to be the strongest fiber in terms of tensile strength since it has the

highest maximum strength among the three fibers. Although the maximum strength of PSSF is lower, their fiber peaks at a shorter extension, indicating a quicker response to tensile stress. As demonstrated by the available data, *Mikania micrantha* fiber appears to be the best choice for applications requiring high tensile strength and extensive extension before breaking. Jute fiber's moderate strength and extension make it an excellent choice for applications needing a balance of flexibility and strength. The fiber from PSSF has the lowest peak strength and extension, making it suitable for less demanding applications.

3.4 XRD analysis

 I_{α} and I_{β} are the two kinds of natural cellulose. Some plant fibers that have a higher availability of I_{β} than others are cotton, jute, flax, and hemp.³⁷ By removing water and creating oxygen bridges between C-1 and C-4, beta cellulose is created. In this case, the stacking of parallel hydrogen-bonded sheets is maintained in part via a van der Waals contact.³⁸ The graph in Fig. 6 depicts the X-ray diffraction (XRD) pattern of natural fiber retrieved from palmyra seed sprouts. The x-axis is labeled "2 theta (degree)" and encompasses 0 to 70 degrees, while the y-axis is labeled "intensity (counts)". The graph displays peaks at multiple vantage points (2θ) representing the sample's

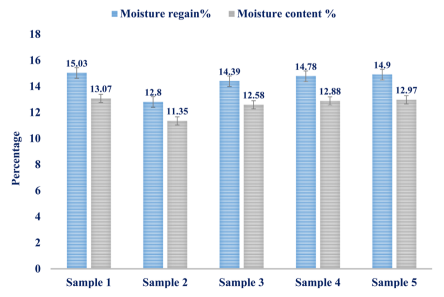


Fig. 4 Moisture regain and content represented in a bar diagram for PSSF.

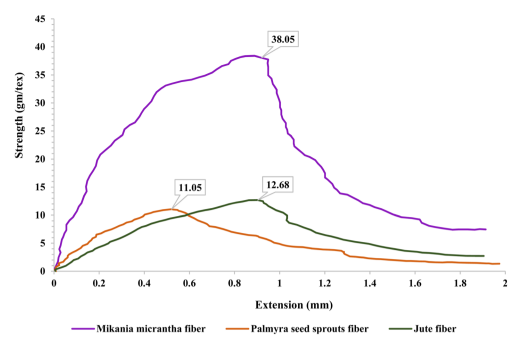


Fig. 5 Strength vs. extension curve for three different natural fibers

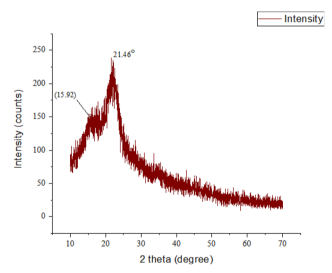


Fig. 6 XRD diagram for PSSF.

crystalline areas. The pattern shows two conspicuous peaks: $2\theta \approx 15.92^\circ$, which is wide and intense, and $2\theta = 21.46^\circ$, the tallest and sharpest peak, indicating a strong crystalline phase. This peak identifies the predictable diffraction of cellulose-I. The 38% crystallinity index (CI) implies that 62% of the natural fiber extracted from PSSF is amorphous and 38% crystalline. Fibers with higher crystalline structure areas show greater mechanical stability and stiffness. The fiber's mechanical strength and stiffness are modest, with this CI. It also denotes that the fiber is thermally stable and may withstand a large temperature range before disappearing. The fiber has a greater heterogeneous component (62%) than more crystalline polymer chains, making it more prone to swelling and disintegration in certain solvents. The fiber's reduced crystallinity index implies that it

would be considered as an adequate choice for bio-composite materials that require a certain ratio.³⁸ The tested and proven natural fiber CIs are contrasted in Table 3.

3.5 TGA and DSC analysis

Fig. 7 below shows the palmyra seed sprout fiber sample's thermogravimetric behavior along with the values found from differential scanning calorimetry. The PSSF's mass change as a function of temperature or time is displayed in this graph. The weight reduction as a percentage of the sample's starting weight is displayed on the graph's left *y*-axis. The heat flow in watts per gram (W g⁻¹) is displayed on the graph's right *y*-axis. The temperature in degree celsius (°C) is displayed on the *x*-axis. Small circles are displayed to represent the data points on the graph. Connecting the data points are lines as well. The weight of the sample reduces with increasing temperature, as the line graph illustrates. This implies that as it is heated, the sample is disintegrating. A more intricate heat flow curve is shown.

Initially the weight reduction starts significantly at 100 °C, reaching 87.7%; this is probably because moisture or volatiles evaporate at that temperature. With a heat flow of 0.182 W g⁻¹ and a first endothermic peak at 59.5 °C, dehydration is suggested. In the intermediate phase, subsequent material changes, maybe a phase shift, are indicated by a large endothermic event at 132.5 °C (0.361 W g⁻¹). Effective decomposition is indicated by the weight loss of 84.342% at 251.5 °C. An exothermic peak at 343.5 °C (-0.188 W g⁻¹), indicating further breakdown, is the most notable thermal event. The weight reduction is substantial by 368 °C (31.284%), which denotes that the decomposition starts. Finally the material continues to decompose or convert at 474 °C as evidenced by its weight loss of 23.102% and heat flow of -0.2 W g⁻¹.

Table 3 PSSF and its comparative investigation of chemical contents with several natural fibers

Name of the fibers	PSSF	Mikania micrantha	Jute	Cornhusks	Cotton stalk	Napier grass	Sugar cane straw	Sorghum stems	Flax	Hemp
Cellulose%	64	56.42	64.4	80-87	79	45.66	33.5	65	62-71	67-75
Hemicellulose%	8	21.42	12	_	_	33.67		_	16-18	16-18
Lignin%	24	15.78	11.8	_	13.7	20.60	25.8	6.5	3-4.5	3-5
Crystallinity index	38	72	71	48-50	47	_	_	_	80	88
Extraction process	Boiling water, 100 °C	5% NaOH, 90–100 °C	Water	5% NaOH	2N NaOH, boiling temperature	Chemical	_	2% NaOH	_	_
References	Present work	3	12	39	40	41	34	42	43	43

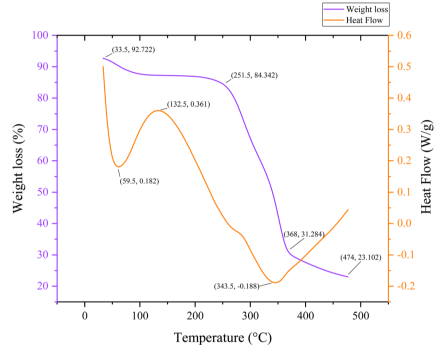


Fig. 7 Diagram of TGA and DSC study for the fiber of PSSF.

The compatibility of PSSF with the other fibers is illustrated in Table 4, which compares the thermal behavior of the multiple fibers listed below.

3.6 SEM analysis

The surface morphology of the sample may be seen intuitively in the SEM image, and its surface roughness and texture can be represented statistically in the 3D surface plot. When these

resources are combined, they enable an exhaustive analysis of the sample, which is useful for research applications where surface properties are critical, as well as material characterization and integrity management.³⁴ At a 100-fold magnification, Fig. 8 provides a thorough look of the sample. The view is around 500 micrometers (μ m) broad, according to the scale bar at the bottom right. Similar to the tested fiber's surface ridges, the surface seems to be composed of parallel, elongated

Table 4 Comparison list for the thermal behavior of different natural fibers

Fiber name	PSSF	Mikania micrantha	Prosopis juliflora	Azadirachta indica	Phaseolus vulgaris	Heteropogon contortus	Saccharum bengalense grass	Jack tree fiber
Preliminary decomposition temperature (°C)	100 °C	103 °C	66.2 °C	97 °C	91.2 °C	90 °C	100 °C	104.5 °C
Imminent disintegration temperature (°C)	474 °C	477 °C	351.3 °C	321.2 °C	328.5 °C	337.7 °C	600 °C	565 °C
Reference	This work	3	44	45	46	47	48	27

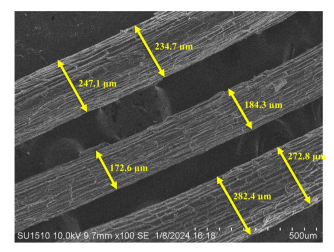


Fig. 8 SEM image of PSSF

structures. These formations have a rough texture that resembles a layered surface. Fiber diameters vary from one another.

The fiber diameter is 232.3 µm on average. Lignin and hemicellulose, which make up the PSSF's outer layer, are surface binding components. Surface roughness enhances the contact surface area, which improves fiber adhesion to the matrix in composite applications.⁴⁸ In Fig. 9, the 3D surface map portrays the sample's surface as visible in the SEM photograph.

The z-axis displays the intensity or height information, which most likely correlates to the grayscale values in the SEM

image, while the x and y axes display the image's pixel dimensions. The surface map depicts height fluctuations over the fiber's surface, with peaks and valleys representing places with higher and lower electron emission, respectively. Differences in electron density, which are connected with intensity values, are related to fiber surface properties such as texture, roughness, and perhaps composition. The end result is that the fiber surface is appropriate for bonding with the matrix for composite reinforcement.

3.7 Chemical composition analysis

Determining the chemical behavior of any kind of substance requires identification. Properties, structure, characteristics, and processing capabilities of fiber are all influenced by its composition. Larger quantities of lignin, hemicellulose, and αcellulose were detected in the fiber obtained from palmyra seed sprouts. It is made up of 24.36% lignin, 08.74% hemicellulose, and 64.74% alpha-cellulose. Elevated cellulose concentration promotes high-value uses and facilitates simple processing for many uses. It can enhance the crystalline features, mechanical properties, biodegradability, hydrolysis resistance, and thermal stability of a material, among other attributes. Small amounts of hemicellulose and lignin are present in any fiber appropriate for high-value composite applications, helping to regulate the fibers' stiffness and bundling. The ash content was 1.58%, moisture content was 12.58%, and extractive percentage was 2.88%, according to the results. The pie chart represents the percentage of the chemical components in Fig. 10 below.

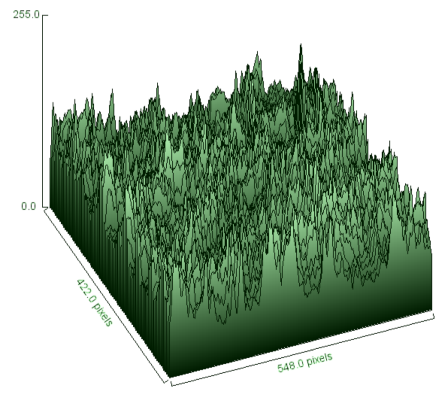
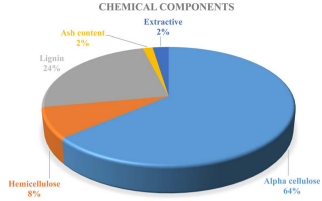


Fig. 9 Surface roughness of PSSF.

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Biological components of PSSF

Table 5 Density measurement of PSSF

Sample no.	Density (g cm ⁻³)	Average (g cm ⁻³)
Sample 1	1.3759	1.3354
Sample 2	1.3194	
Sample 3	1.3567	
Sample 4	1.2985	
Sample 5	1.3267	
Standard deviation		0.0307
Coefficient of variation		2.30%

3.8 Fiber density measurement

Based on the molecular weight and intermolecular tensions inside the fiber, a fiber's mass per unit volume determines its density. The configuration of the fibers and their molecular makeup affect density. Because of their innate cellular structures, natural fibers like wool and cotton typically have a range of densities, while synthetic fibers may be made to have a particular density by regulating the polymerization approach. Fiber tensile strength, elasticity, and durability are mechanical qualities that are influenced by density. Strength and resistance to deterioration are increased in higher density fibers because their molecules are usually packed closer together. A pycnometer examination of the recently extracted fiber from PSSF revealed a density of 1.3354 g cm⁻³. The coefficient of variation is 2.3%. The data are represented in Table 5. Compared to other fibers like cotton, jute, flax, etc., this fiber has a lesser density.49 This suggests that it could be a wise choice for creating fiberreinforced composites.

Responses of the burn test

The PSSF manifested instantaneous combustion, and sparking within seconds upon contact with the flame. A bright yellow flame was encountered, pointing to a quick combustion process. The burning process produced an odd fragrance that was similar to burning wood or leaves, which is a characteristic of an organic cellulose material. 50 After burning, the fibers turned into a light, fluffy gray ash. This ash broke easily in the hands and was quite brittle. The smoke generally appeared gray

or white during the fire. Most importantly, the absence of a fume threat from the smoke indicated that there were no hazardous byproducts. These results have implications for quality control in cellulose-based goods, textile analysis, forensic investigations, and identification.

Conclusion 4.

Cellulose, a highly prevalent basic constituent in the textile industry, has a variety of applications, including fiberreinforced composite materials, medical textiles, clothing, and others. The goal of cellulose-based research has always been to identify new sources. The research also introduced and characterized palmyra seed sprouts, a novel cellulosic fiber. Following retting with water, the substance was evaluated. The chemical analysis suggests that it has more cellulose (64%). The XRD and FTIR data support the comparison of the fibers to other cellulosic fibers. SEM testing confirmed an excellent morphology. The fiber is thermally stable up to 251 °C, with significant deterioration happening between 251 °C and 368 °C, according to thermogravimetric experiments. This breakdown pattern shows the fiber's cellulose content and crystal structure. This fiber offers a moderate density, strength, and elongation. This fiber's potential as a reinforcement for bio-composite materials, cellulose nanoparticles, and biomaterials is being intensively researched.

Data availability

The datasets used and/or analyzed during the current study available from the corresponding author on reasonable request.

Author contributions

Fahmida-E. Karim: conceived and designed the experiments; wrote and revised the paper. Md Redwanul Islam: analyzed and interpreted the data. Afsar Uddin: performed the experiments. Shahidul Islam: contributed reagents, materials, analysis tools or data.

Conflicts 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.

References

- 1 S. Palamutcu, Sustainable Textile Technologies, 2017, pp. 1-22.
- 2 M. M. Devarajan, G. Kumaraguruparan, K. J. Nagarajan and C. Vignesh, Int. J. Biol. Macromol., 2024, 254, 127848.
- 3 F.-E. Karim, M. R. Islam, R. Ahmed, A. B. Siddique and H. A. Begum, Heliyon, 2023, 9, e19360.
- 4 H. Nawaz, X. Zhang, S. Chen, T. You and F. Xu, Carbohydr. Polym., 2021, 267, 118135.

Paper

5 B. M. Bright, B. J. Selvi, S. Abu Hassan, M. M. Jaafar, S. Suchart, S. Mavinkere Rangappa and B. Kurki Nagaraj, *J. Nat. Fibers*, 2022, **19**, 9373–9383.

- 6 M. T. Selvan, M. Ramesh, A. F. Sahayaraj, H. J. Prabu and K. J. Nagarajan, *Int. J. Biol. Macromol.*, 2024, **270**, 132492.
- 7 S. T. K. Rajan, K. J. Nagarajan, V. Balasubramani, K. Sathickbasha, M. R. Sanjay, S. Siengchin and A. N. Balaji, *Int. J. Adhes. Adhes.*, 2023, **126**, 103492.
- 8 N. Sgriccia, M. C. Hawley and M. Misra, *Composites, Part A*, 2008, **39**, 1632–1637.
- 9 F.-E. Karim, M. R. Islam, M. H. Supto, A. A. M. Rafi, T. R. Tanni and H. A. Begum, *Clean. Eng. Technol.*, 2024, 19, 100736.
- 10 J. Biagiotti, D. Puglia and J. M. Kenny, *J. Nat. Fibers*, 2004, **1**, 37–68.
- 11 A. Karimah, M. R. Ridho, S. S. Munawar, Ismadi, Y. Amin, R. Damayanti, M. A. R. Lubis, A. P. Wulandari, Nurindah, A. H. Iswanto, A. Fudholi, M. Asrofi, E. Saedah, N. H. Sari, B. R. Pratama, W. Fatriasari, D. S. Nawawi, S. M. Rangappa and S. Siengchin, *Polymers*, 2021, 13, 4280.
- 12 E. Awwad, B. Hamad, M. Mabsout and H. Khatib, 2nd Int. Conf. Sustain. Constr. Mater. Technol., 2010, pp. 575–582.
- 13 G. R. Raghav, K. J. Nagarajan, M. Palaninatharaja, M. Karthic, R. A. Kumar and M. A. Ganesh, *Int. J. Biol. Macromol.*, 2023, 249, 126119.
- 14 M. R. Sanjay, G. R. Arpitha, L. L. Naik, K. Gopalakrishna and B. Yogesha, *Nat. Resour.*, 2016, 7, 108–114.
- 15 A. M. A. Osman, Acta Hortic., 1984, 231-238.
- 16 D. A. Demason and W. W. Thomson, *Bot. Gaz.*, 1981, **142**, 320–328.
- 17 A. A. Jaradat and A. Zaid, *Food, Agric. Environ.*, 2004, 2, 208–217.
- 18 M. Chandrasekaran and A. H. Bahkali, Saudi J. Biol. Sci., 2013, 20, 105–120.
- 19 M. I. Hussain, M. Farooq and Q. A. Syed, *Food Biosci.*, 2020, 34, 100509.
- 20 V. Balasubramani, K. J. Nagarajan, M. Karthic and R. Pandiyarajan, *Int. J. Biol. Macromol.*, 2024, **259**, 129273.
- 21 Standard Test Methods for Moisture in Textiles, 2023, https://www.astm.org/d2654-22.html.
- 22 Standard Test Method for Breaking Strength and Elongation of Cotton Fibers (Flat Bundle Method), https://www.astm.org/d1445-05.html.
- 23 L. Segal, J. J. Creely, A. E. Martin and C. M. Conrad, *Text. Res. J.*, 1959, **29**, 786–794.
- 24 A. D. O. Betene, F. E. Betene, F. Martoïa, P. J. J. Dumont, A. Atangana and P. M. A. Noah, *J. Miner. Mater. Charact. Eng.*, 2020, 08, 205–222.
- 25 Standard Test Methods for Specific Gravity, Apparent, of Liquid Industrial Chemicals, https://www.astm.org/d0891-18.html.
- 26 Standard Test Method for Flammability of Apparel Textiles 1, https://cdn.standards.iteh.ai/samples/112408/9887694ba6e545e0b446d3e4b8117f74/ASTM-D1230-22.pdf.

- 27 S. Hossain, M. A. Jalil, T. Islam and M. M. Rahman, *Heliyon*, 2022, **8**, e11667.
- 28 A. Azanaw, A. Haile and R. K. Gideon, *Cellulose*, 2019, 26, 795–804.
- 29 P. Senthamaraikannan, M. R. Sanjay, K. S. Bhat, N. H. Padmaraj and M. Jawaid, J. Nat. Fibers, 2019, 16, 1124–1131.
- 30 G. Rajeshkumar, G. L. Devnani, J. P. Maran, M. R. Sanjay, S. Siengchin, N. A. Al-Dhabi and K. Ponmurugan, *Cellulose*, 2021, 28, 5373–5385.
- 31 S. Indran and R. E. Raj, *Carbohydr. Polym.*, 2015, **117**, 392–399.
- 32 P. Manimaran, P. Senthamaraikannan, M. R. Sanjay, M. K. Marichelvam and M. Jawaid, *Carbohydr. Polym.*, 2018, 181, 650-658.
- 33 R. Dalmis, S. Köktaş, Y. Seki and A. Ç. Kılınç, *Cellulose*, 2020, 27, 127–139.
- 34 S. M. Costa, P. G. Mazzola, J. C. A. R. Silva, R. Pahl, A. Pessoa and S. A. Costa, *Ind. Crops Prod.*, 2013, 42, 189–194.
- 35 A. Célino, S. Fréour, F. Jacquemin and P. Casari, *J. Appl. Polym. Sci.*, 2013, **130**, 297–306.
- 36 J. Gassan and A. K. Bledzki, *J. Appl. Polym. Sci.*, 1999, **71**, 623–629.
- 37 H. Zhao, J. Kwak, Z. Conradzhang, H. Brown, B. Arey and J. Holladay, *Carbohydr. Polym.*, 2007, **68**, 235–241.
- 38 B. A. N. and N. K. J., Carbohydr. Polym., 2017, 174, 200-208.
- 39 N. Reddy and Y. Yang, Green Chem., 2005, 7, 190.
- 40 N. Reddy and Y. Yang, *Bioresour. Technol.*, 2009, **100**, 3563–3569.
- 41 K. O. Reddy, C. U. Maheswari, M. Shukla and A. V. Rajulu, *Mater. Lett.*, 2012, **67**, 35–38.
- 42 N. Reddy and Y. Yang, J. Agric. Food Chem., 2007, 55, 5569-5574.
- 43 G. Gedik, Cellulose, 2021, 28, 6899-6915.
- 44 S. S. Saravanakumar, A. Kumaravel, T. Nagarajan, P. Sudhakar and R. Baskaran, *Carbohydr. Polym.*, 2013, **92**, 1928–1933.
- 45 P. Manimaran, P. Senthamaraikannan, K. Murugananthan and M. R. Sanjay, *J. Nat. Fibers*, 2018, 15, 29–38.
- 46 B. Gurukarthik Babu, D. Prince Winston,
 P. SenthamaraiKannan, S. S. Saravanakumar and
 M. R. Sanjay, J. Nat. Fibers, 2019, 16, 1035–1042.
- 47 N. R. J. Hyness, N. J. Vignesh, P. Senthamaraikannan, S. S. Saravanakumar and M. R. Sanjay, *J. Nat. Fibers*, 2018, 15, 146–153.
- 48 R. Vijay, D. L. Singaravelu, A. Vinod, I. D. F. Paul Raj, M. R. Sanjay and S. Siengchin, J. Nat. Fibers, 2020, 17, 1739–1747.
- 49 I. Elfaleh, F. Abbassi, M. Habibi, F. Ahmad, M. Guedri, M. Nasri and C. Garnier, *Results Eng.*, 2023, **19**, 101271.
- 50 M. N. Azman Mohammad Taib, T. S. Hamidon, Z. N. Garba, D. Trache, H. Uyama and M. H. Hussin, *Polymer*, 2022, **244**, 124677.