

## From Filament to Functional Part: A Comparative Analysis of Structure-Property Relationships in 3D-Printed Coir Fiber Reinforced Polyactic Acid Biocomposites



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### ABSTRACT

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The integration of natural fibers into polylactic acid (PLA) for fused deposition modeling (FDM) holds promise for sustainable additive manufacturing. However, a systematic understanding of how material properties evolve from the composite filament feedstock to the final printed part is lacking. This study bridges this gap by fabricating and comprehensively characterizing PLA biocomposite filaments reinforced with 0, 5, 7, and 9 wt.% sodium bicarbonate-treated coir fibers, followed by a critical comparison with their FDM-printed counterparts. Fourier-Transform Infrared Spectroscopy (FTIR) confirmed the preservation of chemical structures post-processing. X-ray diffraction analysis indicated that coir fibers acted as nucleating agents, enhancing PLA crystallinity with increasing fiber content. Thermogravimetric analysis revealed a marginal decrease in thermal stability attributed to the lignocellulosic nature of the fibers. Tensile testing showed that neat PLA filament exhibited the highest tensile strength (43.26 MPa), whereas the incorporation of coir fibers resulted in lower tensile strengths of 33.96 MPa (5 wt.%), 35.37 MPa (7 wt.%), and 35.22 MPa (9 wt.%). Notably, filaments consistently outperformed their 3D-printed versions across all compositions, primarily due to printing-induced anisotropy and interlayer voids. Scanning electron microscopy corroborated these findings, showing fiber agglomeration at higher contents (9 wt.%) and interfacial gaps that compromised stress transfer. The central conclusion is that for FDM-processed PLA/coir composites, process-induced defects exert a more dominant influence on mechanical performance than fiber reinforcement itself, especially at higher fiber contents. This underscores the necessity for concurrent optimization of material formulation and printing parameters to harness the full potential of these sustainable biocomposites in additive manufacturing.

## 1. INTRODUCTION

In the past decade, additive manufacturing (AM), also referred to as three-dimensional (3D) printing, has undergone remarkable development and has become a key area of interest in both academic research and industrial practice. This manufacturing paradigm enables the direct fabrication of components with complex geometries, high design flexibility, and shortened development cycles using a wide variety of polymer-based feedstocks [1-3]. Among the available AM technologies, fused deposition modelling (FDM) has achieved the most significant level of industrial adoption owing to its straightforward processing principle, relatively low equipment and operating costs, efficient material use, and the widespread commercial availability of thermoplastic filaments [4]. In FDM, a solid thermoplastic filament is continuously fed into a heated extrusion nozzle, where it is melted and deposited sequentially in a layer-by-layer fashion to form three-dimensional objects [5]. Polymers commonly processed via FDM include polylactic acid (PLA), polystyrene (PS), acrylonitrile butadiene styrene (ABS), and polycarbonate

(PC), representing both biodegradable and conventional thermoplastic materials suitable for additive manufacturing applications [6-9].

PLA is among the most intensively studied thermoplastic polymers for FDM owing to its favorable melt processability, balanced mechanical performance, biocompatibility, intrinsic biodegradability, and high optical clarity [6, 10]. Nevertheless, the broader application of PLA in load bearing or thermally demanding environments is restricted by several inherent shortcomings, including limited thermal stability, slow crystallization behavior, low fracture strain, and inadequate flexibility [11, 12]. In response to these limitations, current research efforts have increasingly shifted toward the formulation of PLA-based composite systems by incorporating reinforcing phases. A growing body of literature has demonstrated that integrating natural fibers, nanocellulose, hydroxyapatite, pulp fibers, and various organic or inorganic particulate reinforcements can substantially enhance the mechanical, thermal, and functional properties of PLA, thereby extending its applicability in advanced engineering [13-16].

Natural fibers have gained increasing interest as reinforcements for biopolymer composites due to their renewability, biodegradability, non-toxicity, and low environmental impact. Materials such as jute, sisal, hemp, banana, pineapple leaf fibers, and coir are widely used to reduce carbon footprint while improving composite sustainability. To strengthen fiber matrix interfacial adhesion, various surface modification techniques, including alkali, peroxide, and permanganate treatment, are commonly applied to improve compatibility at the interface [17-19]. Among these natural fiber candidates, coir stands out as a promising reinforcement for PLA-based composites owing to its low density, low thermal conductivity, and cost-effectiveness, making it suitable for lightweight structural applications [20]. Coir also exhibits strong acoustic absorption and favorable resistance to moisture and biological degradation, both of which are attributed to its lignocellulosic composition. Chemically, coir contains moderate cellulose (36-43%), low hemicellulose content, and a relatively high lignin fraction (41-45%), which contributes to its large microfibril angle and results in fibers with lower tensile strength but remarkably high elongation at break [21-23]. These characteristics make coir particularly attractive for applications where toughness and damage tolerance are critical rather than high stiffness alone. Traditionally used in ropes, mats, and household products [24], coir fiber has more recently been incorporated into acoustic panels, thermal insulation materials, construction components, and automotive parts. This expanding range of applications underscores its potential as an environmentally sustainable reinforcement for the development of lightweight and durable PLA biocomposites [25-27].

Extensive research has examined coir-reinforced green composites, demonstrating their effectiveness in enhancing the performance of various biodegradable polymer systems. For PLA-based composites, Dong et al. [28] investigated materials containing 5-30 wt.% coir fibers fabricated via hydraulic pressing. Their findings indicated that increasing fiber content generally reduced tensile and flexural properties. NaOH-treated fibers significantly improved overall mechanical performance, with the highest strength observed at 20 wt.% fiber content. Similarly, Nam et al. [22] developed biodegradable PBS/coir composites and reported that immersing coir fibers in 5% NaOH solution for 72 h increased the interfacial shear strength by 55.6%, resulting in stiffer and stronger composites. Brahmakumar et al. [29] further demonstrated that inadequate fiber surface modification severely degrades fiber-matrix adhesion, leading to pronounced reductions in tensile modulus and strength. While these studies clearly establish the importance of fiber treatment and fiber content, they are mainly limited to bulk-processed or compression-molded composites and do not account for the additional complexities introduced by filament extrusion and layer-by-layer deposition in FDM.

Critically, despite the increasing use of natural fiber-reinforced PLA in additive manufacturing, systematic investigations addressing the fabrication of reinforced filaments and the subsequent evolution of properties during the FDM printing process remain scarce. Most existing studies evaluate only the final printed components or bulk specimens, thereby overlooking the distinct roles of filament quality, extrusion-induced fiber alignment, and printing-induced anisotropy and interlayer void formation. As a result, the transition in material properties from the filament stage to the final printed part remains insufficiently understood for

PLA/coir systems. To address this gap, the present study provides a direct, systematic comparison of extruded PLA/coir biocomposite filaments and their corresponding FDM-printed specimens. Filaments containing 0, 5, 7, and 9 wt.% coir fiber were produced via single-screw extrusion to examine the influence of fiber incorporation on the mechanical, thermal, and structural properties of the composites. Fourier-Transform Infrared (FTIR), X-ray diffraction (XRD), and TGA analyses were conducted to evaluate chemical structure, crystallinity, and thermal behavior. In addition, scanning electron microscopy (SEM) observations were performed on both extruded filaments and 3D-printed specimens to assess fiber dispersion and fiber-matrix interfacial integrity. By explicitly linking filament characteristics to printing-induced microstructural features and mechanical performance, this work aims to clarify the dominant factors governing the performance limitations of PLA/coir biocomposites in FDM applications.

## 2. MATERIALS AND METHODS

### 2.1 Materials

SUNLU PLA + 2.0 served as the polymer matrix in this study. The material was supplied as pellets with a nominal filament diameter of 1.75 mm and a density of 1.24 g/cm<sup>3</sup> and was imported from the United States. Coir fibers obtained from Donggala Regency, Central Sulawesi, Indonesia, functioned as the reinforcing phase. The fibers exhibited a density of 1.15 g/cm<sup>3</sup> and an average diameter of 100-200 μm. Sodium bicarbonate (NaHCO<sub>3</sub>), pro analysis grade, supplied by Merck (Germany), was employed as the chemical reagent.

### 2.2 Fiber treatment

Coir fibers were mechanically extracted from coconut husks and subsequently washed under running water to remove residual impurities. The fibers were then treated with a sodium bicarbonate (NaHCO<sub>3</sub>) solution, a method reported in previous studies to preserve fiber integrity and enhance tensile performance [30]. To leverage this optimal condition, the fibers in the present study were immersed in a 12 wt.% NaHCO<sub>3</sub> solution for 24 h at ambient temperature. After the treatment, the fibers were rinsed with distilled water until the pH was neutral, then dried in an oven at 115°C for 1 h. The dried fibers were subsequently cut to 5-7 mm, ground to a fine powder, and sieved through a 200-mesh screen to obtain a uniform particle size suitable for subsequent compounding. Figure 1 presents an overview of the fiber treatment process.

### 2.3 Filament fabrication and three-dimensional printed samples

Before extrusion, coconut shell fibers after being sieved through a 200-mesh screen were oven-dried at 65°C for 3 h to minimize moisture content and prevent hydrolytic degradation during processing. The dried fibers were subsequently blended with PLA pellets in a milling mixer to prepare composite feedstocks containing 5, 7, and 9 wt.% coir fiber. Neat PLA (0 wt.% fiber) and the prepared composite feedstocks were then processed using a Wellzoom single-screw desktop extruder (China). The extrusion process employed a barrel temperature of 147°C, an extrusion speed of 1200 mm/min, and a target

filament diameter of 1.75 mm. The extrusion process yielded four types of filaments, neat PLA (0 wt.% coir), which served as the reference material, and PLA/5% coir, PLA/7% coir, and PLA/9% coir. All extruded filaments were cooled under ambient conditions, collected, and stored for subsequent mechanical and physical characterization.

Tensile test specimens were fabricated in accordance with ASTM D638-22 [31, 32]. The specimen geometry was designed using computer-aided design (CAD) software and

prepared for printing using OrcaSlicer. For each formulation, seven specimens were manufactured using a Kingroon KP3S FDM 3D printer (China). The printing process employed a nozzle diameter of 0.3 mm, a nozzle temperature of 250°C, a bed temperature of 70°C, a layer height of 0.2 mm, and an infill density of 100%. Figure 2 presents a schematic representation of the filament preparation and specimen fabrication workflow.

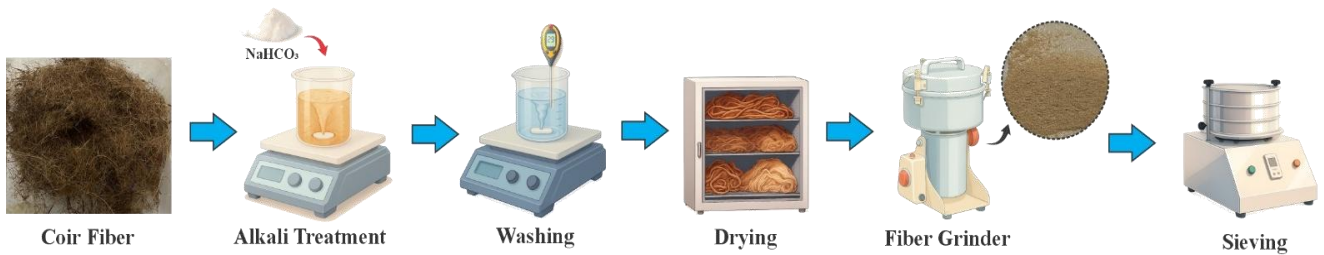


Figure 1. Treatment process of coir fibers

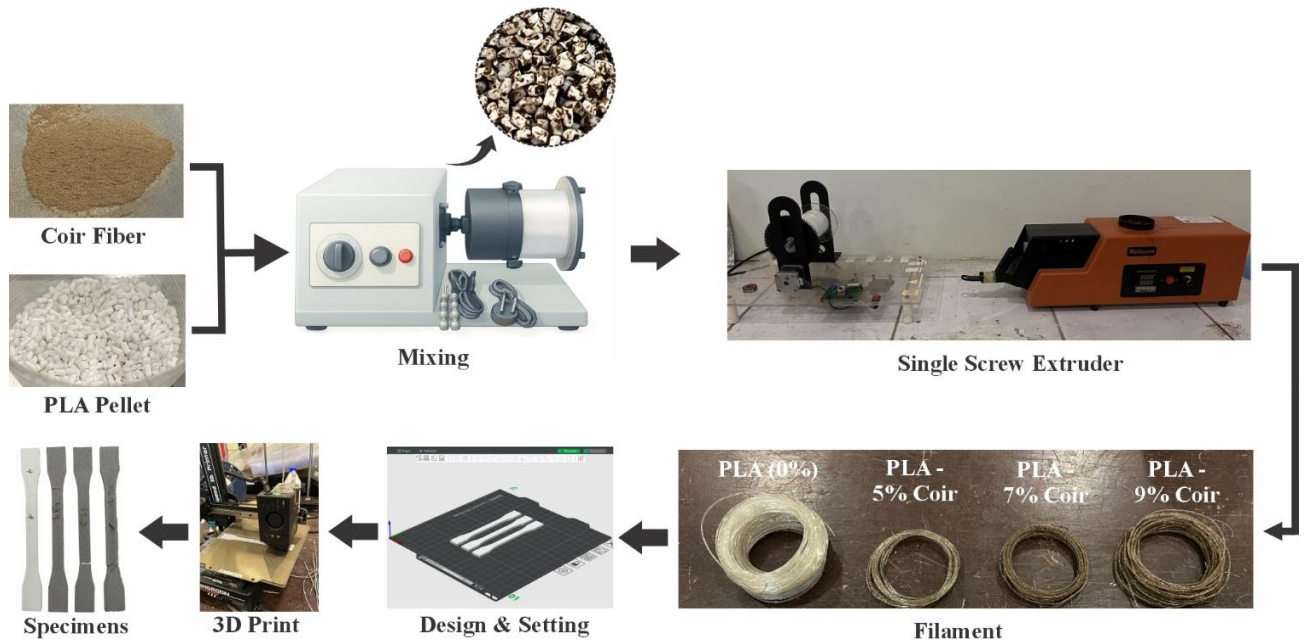


Figure 2. Process schematic for filament manufacturing and three-dimensional (3D) printing of test samples

### 3. CHARACTERIZATION

#### 3.1 Scanning electron microscopy and X-ray spectroscopy

A JSM-IT700HR SEM (JEOL Ltd.), operated at an accelerating voltage of 10 kV, was used to examine the surfaces of the extruded filaments. The instrument was equipped with an energy-dispersive X-ray spectroscopy (EDS) system to perform quantitative elemental analysis. In addition, the fracture surfaces of tensile-tested 3D-printed specimens were analyzed by SEM to characterize the failure mechanisms and to evaluate fiber–matrix interfacial behavior.

#### 3.2 Fourier-transform infrared

A Shimadzu IRAffinity-1S FT-IR spectrometer (Japan) was used to characterize the chemical structures of the coir-reinforced PLA biocomposite filaments over a wavenumber

range of 500-4000 cm<sup>-1</sup>.

#### 3.3 X-ray diffraction

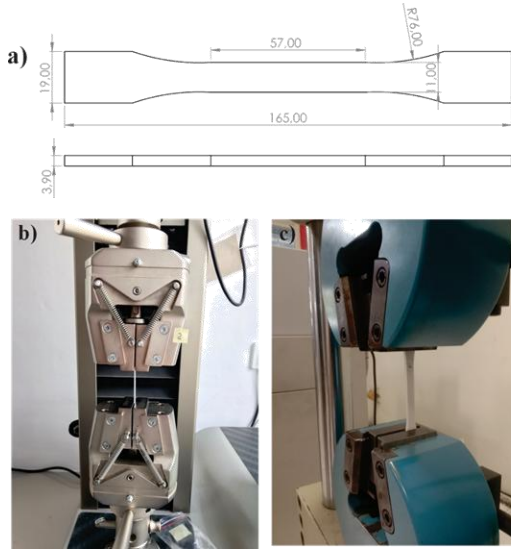
A Rigaku MiniFlex 600 X-ray diffractometer was used to analyze the crystalline structures of neat PLA, coir fibers, and the corresponding composite filaments. The measurements were performed over a 2θ range of 0-50° to characterize the phase composition and crystallinity of the samples.

#### 3.4 Thermogravimetric analysis

A Hitachi STA7300 thermogravimetric analyzer (Japan) was used to evaluate the thermal behavior and stability of the samples. Specimens containing different coir fiber contents were heated from 25°C to 600°C at a heating rate of 10°C/min under a nitrogen flow rate of 50 mL/min to investigate the effect of fiber incorporation on the thermal degradation characteristics.

### 3.5 Tensile properties

A universal testing machine (Universal Testing Machine (UTM)), TN-MD equipped with a 200 kN load cell was used to determine the tensile properties of the specimens. The tests were conducted at a crosshead speed of 1 mm/min, and seven specimens were evaluated for each formulation to ensure the statistical reliability and reproducibility of the results, as shown in Figure 3.



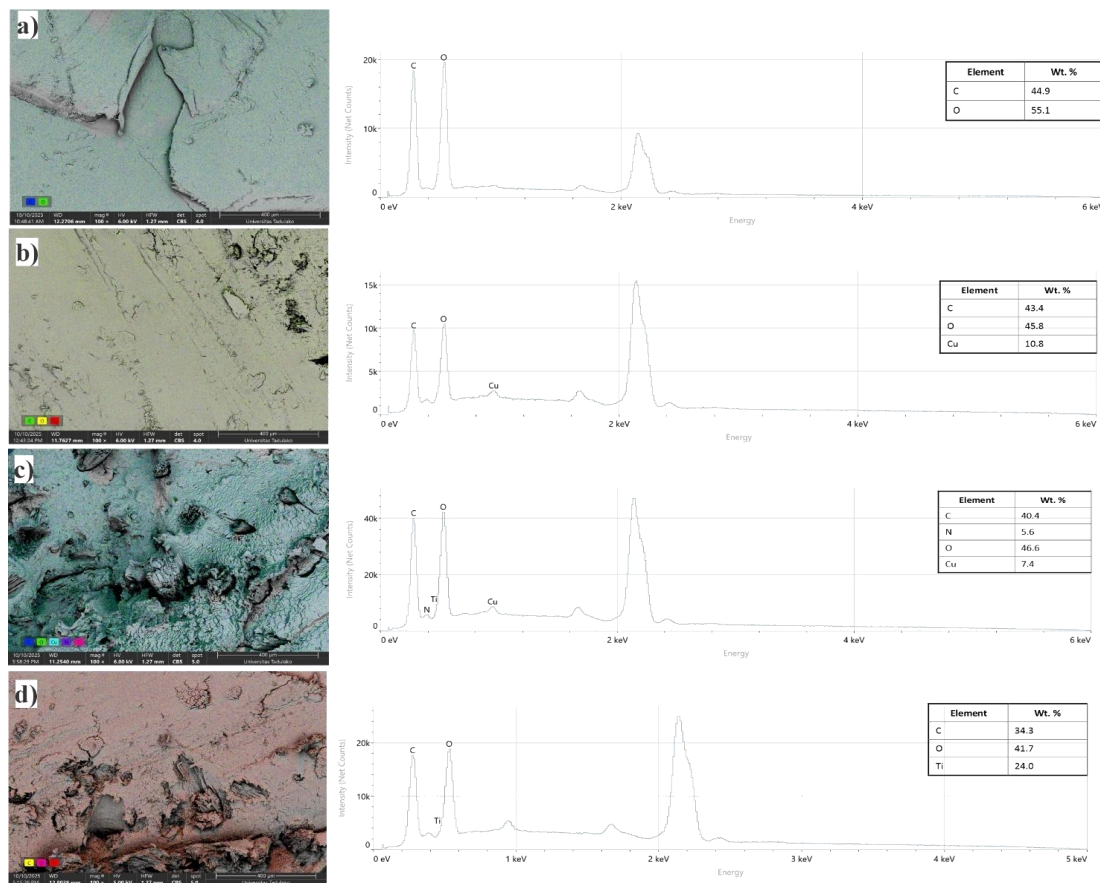
**Figure 3.** Sample design and mechanical testing setup: a) specimen design for tensile test, b) tensile test of filament, c) tensile test of three-dimensional (3D)-printed

## 4. RESULTS AND DISCUSSION

### 4.1 Scanning electron microscopy and X-ray spectroscopy analysis

Figure 4(a) illustrates the smooth and homogeneous surface morphology of neat PLA. At the same time, EDS analysis reveals carbon and oxygen contents of 44.9 wt.% and 55.1 wt.%, respectively, which are consistent with the expected chemical composition of PLA. The incorporation of 5 wt.% coir fibers (Figure 4(b)) increases surface roughness and introduces light–dark contrasts indicative of cellulose-rich regions at a coir fiber content of 7 wt.% (Figure 4(c)), the fibers exhibit a more uniform dispersion within the PLA matrix. In contrast, at a coir fiber content of 9 wt.% (Figure 4(d)), pronounced fiber agglomeration is observed, which is associated with an increased O/C ratio of 1.22, indicative of the higher cellulose content.

The limited fiber dispersion observed at higher fiber contents supports the findings of dos Santos et al. [33], who reported that elevated natural fiber fractions reduce composite homogeneity and promote the formation of interfacial voids or weak interfacial regions. These microstructural defects act as stress concentrators, reducing the composite’s effective load-bearing cross-sectional area and contributing to the decrease in elastic modulus and tensile strength observed at higher fiber contents (Figure 4-11). In addition, SEM observations of the fracture surfaces reveal voids in all filaments formed during the extrusion process. These voids originate from the expansion of the molten material as it exits the extrusion nozzle under pressure gradients, consistent with previous reports [34-36].



**Figure 4.** Surface morphology and X-ray spectroscopy (EDS) compositional mapping of a) neat polylactic acid (PLA), b) PLA/5% coir, c) PLA/7% coir, and d) PLA/9% coir

## 4.2 Diameter measurements of filament

Table 1 summarizes the measured filament diameters. Neat PLA filaments exhibited a diameter of 1.75 mm. In contrast, the coir-reinforced filaments ranged from 1.72 to 1.83 mm, indicating that the incorporation of coir fibers induced only minor changes in filament dimensions. This behavior can be attributed to the physical properties of the coir powder, which affect melt flow and filler–matrix interactions [37], consistent with previous observations in PLA/PCL filaments reinforced with hydroxyapatite [38]. All filament diameters remained close to the nominal 1.75 mm nozzle size, with slight deviations arising from natural shrinkage during cooling [39] or from extrusion parameters such as applied force, melt viscosity, temperature, and screw speed [40, 41]. Overall, the observed diameter variations (0-4.57%) fall within acceptable tolerances for FDM printing.

**Table 1.** Filament diameters of biocomposites with varying fiber contents

Sample	Coir Content (wt.%)	Diameter (mm)
Neat poly(lactic acid) (PLA)	0	1.75 ± 0.007
PLA/5% Coir	5	1.72 ± 0.009
PLA/7% Coir	7	1.82 ± 0.041
PLA/9% Coir	9	1.83 ± 0.014

## 4.3 Fourier-transform infrared analysis

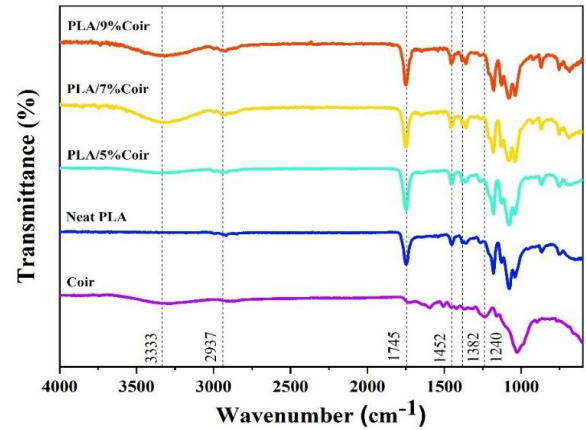
Figure 5 presents the FTIR spectra of neat PLA, coir fibers, and PLA/coir composite filaments containing 0, 5, 7, and 9 wt.% fiber. The coir fibers exhibit characteristic lignocellulosic absorption bands, including a broad hydroxyl (–OH) stretching vibration at 3333  $\text{cm}^{-1}$  and C–H stretching vibrations at approximately 2937  $\text{cm}^{-1}$ . Additional absorption peaks at 1452, 1382, and 1240  $\text{cm}^{-1}$  correspond to  $\text{CH}_2$  bending, C–H deformation, and asymmetric C–O–C stretching vibrations, respectively, which are associated with cellulose and hemicellulose components [39-42]. In contrast, the FTIR spectrum of neat PLA shows distinct absorption bands at 1745, 1452, 1180, 1080, and 869  $\text{cm}^{-1}$ , which are assigned to C=O stretching, C–O stretching, CH bending, and C–C skeletal vibrations, thereby confirming the molecular structure of the polymer [41, 42]. The FTIR spectra of the PLA/coir composite filaments closely resemble that of neat PLA, indicating that the incorporation of coir fibers does not induce chemical modification of the PLA matrix.

However, peak intensity increases with increasing fiber content. The –OH band at 3333  $\text{cm}^{-1}$  broadens and intensifies, reflecting the hydrophilic hydroxyl groups originating from cellulose and hemicellulose [40, 41]. CH-related vibrations at 2998 and 2945  $\text{cm}^{-1}$  and the C=O stretching at 1745  $\text{cm}^{-1}$  remain clearly discernible [40-42]. A minor peak near 1622  $\text{cm}^{-1}$ , attributed to moisture absorption [41, 42], becomes more prominent in the composites, consistent with the hygroscopic nature of lignocellulosic fibers. Peaks at 1452 and 1382  $\text{cm}^{-1}$ , corresponding to  $\text{CH}_3$  bending, increase with fiber content, confirming the contribution of the cellulose-rich filler [40].

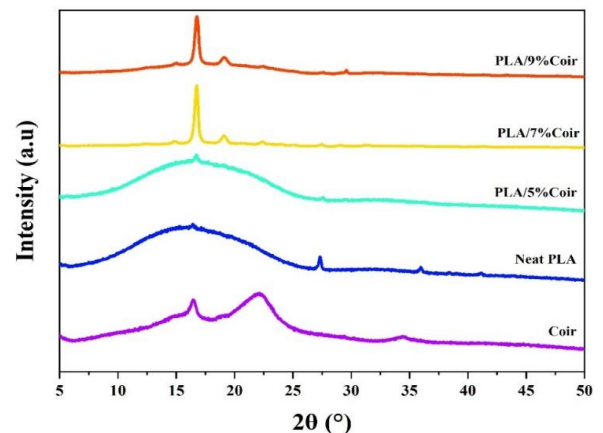
## 4.4 X-ray diffraction analysis

Figure 6 presents the XRD patterns of biocomposite filaments containing 0, 5, 7, and 9 wt.% coir fiber. The coir fibers exhibit characteristic cellulose I diffraction peaks at  $2\theta$

$\approx 15\text{-}6^\circ$ ,  $22\text{-}23^\circ$ , and  $34^\circ$ , corresponding to the 110, 200, and 004 crystallographic planes and confirming their semicrystalline structure [42-44]. Neat PLA shows distinct diffraction peaks at  $16.5^\circ$  and  $19.5^\circ$ , indicating its inherent semicrystalline nature [45, 46]. The PLA/coir composite filaments retain the primary crystalline features of PLA. In contrast, the increasing intensity of the  $16.5^\circ$  peak with higher coir fiber content suggests that the coir fibers act as nucleating agents and promote enhanced crystallinity within the PLA matrix [34].



**Figure 5.** Fourier-Transform Infrared Spectroscopy (FTIR) spectra of coir, neat poly(lactic acid) (PLA) and PLA/Coir fiber filaments with different coir contents



**Figure 6.** X-ray diffraction (XRD) patterns of coir, neat poly(lactic acid) (PLA) and PLA/Coir fiber filaments with different coir contents

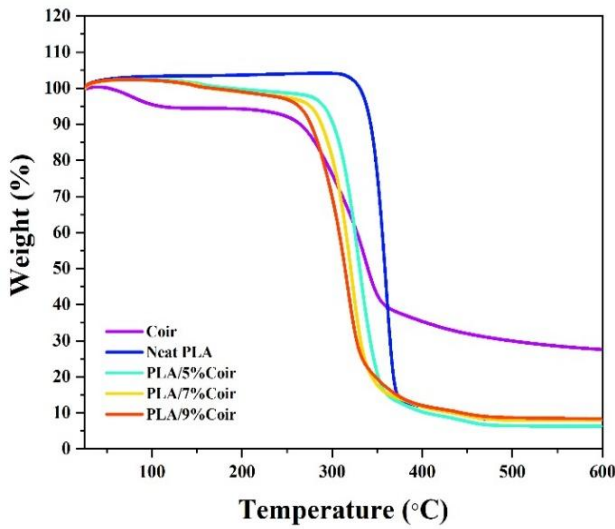
The enhancement in crystallinity arises from the restricted mobility of PLA chains caused by interfacial interactions with cellulose in the coir fibers. Hydrogen bonding between the hydroxyl (–OH) groups of the coir fibers and the polar functional group of PLA forms a stiffer interfacial region, which promotes the nucleation of numerous small crystalline domains that are commonly associated with transcrystalline structures in natural fiber–reinforced composites [47]. This mechanism is consistent with the findings of Tao et al. [48], who reported that lignocellulosic fillers increase PLA crystallinity. At higher fiber contents (7-9 wt.%), the diffraction peak intensities further increase, indicating the formation of additional nucleation sites on the fiber surfaces. The large surface area of the coir fibers facilitates extensive nucleation, leading to the development of overlapping small

crystalline domains, restricting spherulite growth, and resulting in a denser, more fragmented, and thermally stable crystalline structure [49].

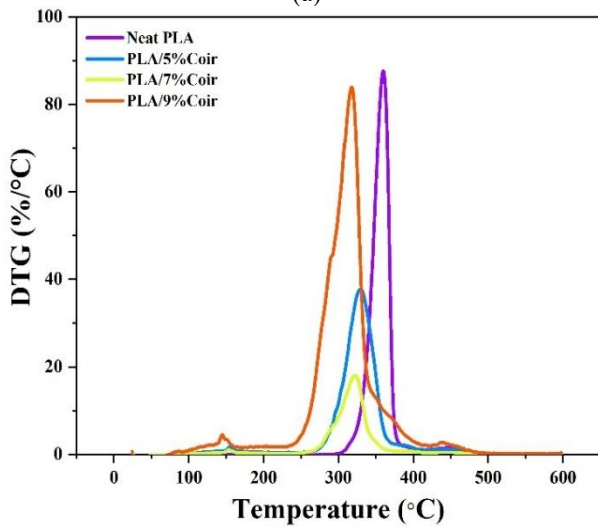
#### 4.5 Thermal analysis results (TGA)

Figure 7(a) illustrates the thermogravimetric behavior of coir fibers, which undergo three principal degradation stages typical of lignocellulosic materials: moisture loss (25-150°C), hemicellulose decomposition (190-290°C), and cellulose depolymerization (290-360°C), while lignin decomposes over a broader temperature range (280-500°C) due to its aromatic structure [50, 51]. Neat PLA exhibits a single, sharp degradation peak with a maximum degradation temperature ( $T_{max}$ ) of 360.67°C (Figure 7(b)), indicative of rapid chain scission within its semi-crystalline structure.

In PLA/coir biocomposites,  $T_{max}$  progressively decreases with increasing coir content (5-9 wt.%), reflecting reduced thermal stability attributable to the lower thermal resistance of lignocellulosic fibers. Similar trends have been observed in PLA composites reinforced with sisal and phormium fibers, where early degradation of the natural fibers accelerates overall mass loss [52, 53]. For composites with low fiber contents, TGA patterns remain dominated by PLA, consistent with reports on other low-fiber-content natural fiber composites [54, 55]. The derivative thermogravimetric (DTG) curves (Figure 7(b)) further confirm that  $T_{max}$  decreases with increasing coir content. Lignocellulosic fibers degrade at lower temperatures than PLA, leading to earlier mass loss, and the hydroxyl-rich groups in cellulose and hemicellulose contribute to this reduction in degradation temperature. Additionally, interfacial interactions between PLA and coir may restrict polymer chain mobility, influencing the thermal behavior of the composites. At higher fiber contents (7-9 wt.%), DTG peaks broaden, reflecting the overlapping degradation of PLA and coir components. Despite the reduction in thermal stability, the composites exhibit slightly higher residual mass at 600°C, consistent with other natural fiber systems [56].



(a)

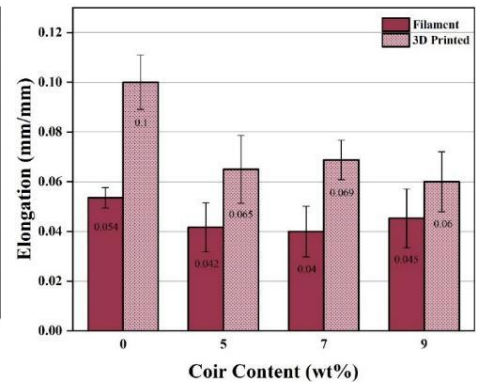
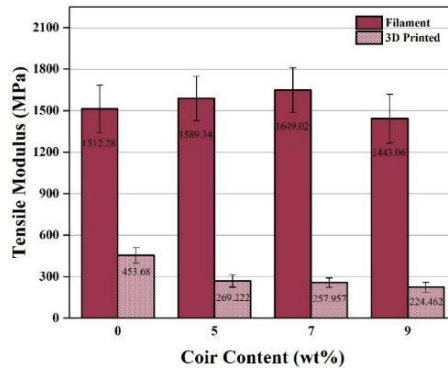
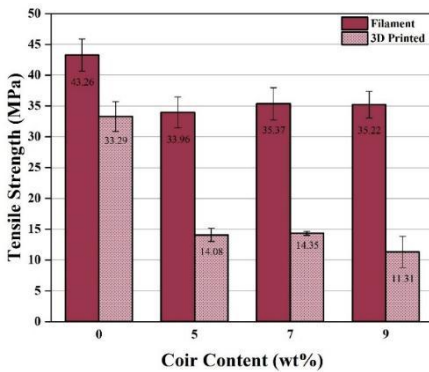


(b)

**Figure 7.** a) Thermogravimetric analysis (TGA) and b) derivative thermogravimetric (DTG) curves of PLA/coir biocomposite filaments at varying coir contents

**Table 2.** Thermal behaviour of coir fiber, neat polylactic acid (PLA), and PLA/coir biocomposites

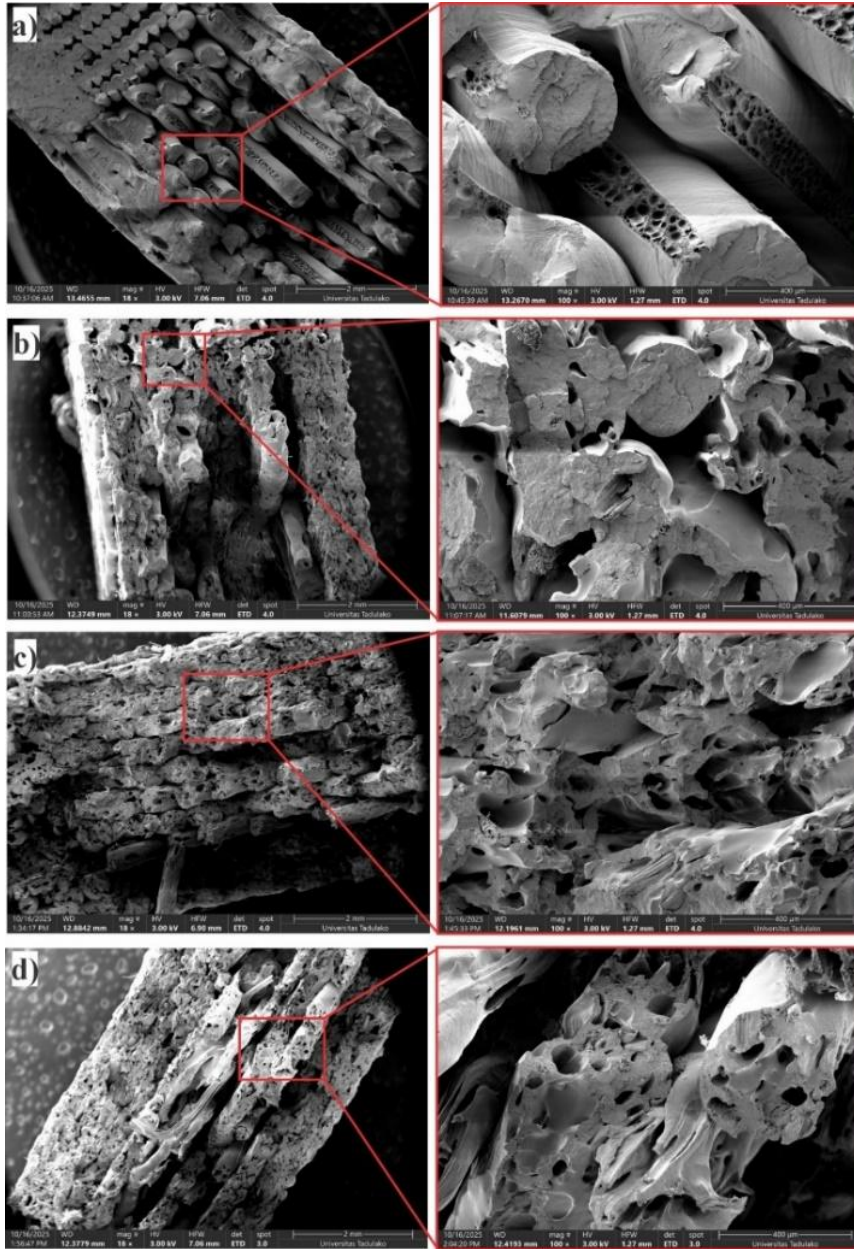
Sample	$T_{onset}$ (°C)	$T_{max}$ (°C)	$W_{residue}$ (%)
Coir	294.30	338.16	8.77
Neat PLA	342.10	360.67	10.83
PLA/5% Coir	302.60	330.59	6.21
PLA/7% Coir	293.80	321.93	4.46
PLA/9% Coir	280.80	316.45	8.16



**Figure 8.** Tensile strength of filaments and three-dimensional (3D)-printed

**Figure 9.** Tensile modulus of filaments and three-dimensional (3D)-printed

**Figure 10.** Elongation at break of filaments and three-dimensional (3D)-printed



**Figure 11.** Scanning electron microscopy (SEM) micrographs of the fracture surfaces of (a) neat PLA, (b) PLA/5 wt.% coir, (c) PLA/7 wt.% coir, and (d) PLA/9 wt.% coir biocomposites at magnifications of 18 $\times$  and 100 $\times$

The onset temperatures ( $T_{\text{onset}}$ ) presented in Table 2 justify these observations. Coir fibers exhibit  $T_{\text{onset}}$  of 294.30 $^{\circ}\text{C}$ , neat PLA 342.10 $^{\circ}\text{C}$ , and PLA/coir composites show a progressive decrease with increasing fiber content: PLA/5 % at 302.60 $^{\circ}\text{C}$ , PLA/7 % at 293.80 $^{\circ}\text{C}$ , and PLA/9 % at 280.80 $^{\circ}\text{C}$ . These trends are consistent with those observed in PLA/cellulose nanocrystal composites, where low-molecular-weight components such as hemicellulose and lignin induce early weight loss [14].  $T_{\text{max}}$  follows a similar pattern, decreasing from 360.67 $^{\circ}\text{C}$  for neat PLA to 330.59, 321.93, and 316.45 $^{\circ}\text{C}$  for 5, 7, and 9 wt.% coir, respectively. Residual mass ( $W_{\text{residue}}$ ) in the composites ranges from 4.46–8.16 %, reflecting the interplay between PLA and coir during pyrolysis.

#### 4.6 Tensile properties of filaments and three-dimensional-printed specimens

Figure 8 illustrates that the tensile strength of the filaments decreases with increasing coir fiber content. For each composition and processing condition, tensile properties were

determined from at least seven specimens, and the reported values correspond to the standard deviation. Neat PLA exhibits the highest tensile strength (43.26 MPa), whereas composites containing 5–9 wt.% coir demonstrate reduced strengths, reaching 35.37 MPa at 7 wt.% and 35.22 MPa at 9 wt.%. The reduction in mechanical performance results from the lower intrinsic strength of lignocellulosic fibers and the presence of interfacial defects, such as voids and fiber pull-out, consistent with previous studies on natural fiber-reinforced PLA composites and evidenced by the SEM observations shown in Figure 11. The 3D-printed specimens exhibit further decreases in tensile strength due to anisotropy and interlayer voids that act as stress concentrators, thereby limiting molecular entanglement between adjacent printed layers [57–59].

Figure 9 illustrates that the tensile modulus of the filaments initially increases with the incorporation of coir fibers, reaching a maximum of 1649.02 MPa at 7 wt.% coir, before decreasing to 1443.06 MPa at 9 wt.%. At moderate fiber contents, the PLA matrix is effectively stiffened due to

efficient stress transfer between the matrix and the fibers [28]. In contrast, higher fiber contents promote fiber agglomeration, inadequate wetting, and the formation of voids, which collectively reduce stiffness, consistent with observations in high-fiber-content natural fiber composites [28, 58]. Across all compositions, the 3D-printed specimens exhibit lower tensile modulus than their filament counterparts, attributed to interlayer gaps and internal porosity [60, 61].

Figure 10 demonstrates that neat PLA exhibits the highest ductility, whereas the rigid coir fibers restrict polymer chain mobility. A slight increase in elongation at break for 3D-printed composites containing 7-9 wt.% coir may be attributed to fiber bundles facilitating localized deformation [58]. Nevertheless, the printed specimens remain less ductile than the filaments, reflecting the influence of interlayer defects and voids that promote premature crack initiation, as similarly reported for PLA/cellulose nanocrystal and PLA/recycled-paper composites [62, 63].

#### 4.7 Morphological structures

Figure 11(a) shows that neat PLA exhibits layer lines, interlayer cracks, and small voids resulting from incomplete interlayer fusion, which leads to brittle fractures along the printing direction. These microstructural features are consistent with previous reports indicating that deficiencies in interlayer bonding in FDM-printed PLA act as primary sites for crack initiation and consequently reduce tensile strength [64].

Figures 11(b-d) illustrate the fracture surfaces of the PLA/coir composite specimens. Fiber pull-out generates voids in all composite samples, while the composite containing 5 wt.% coir (Figure 11(b)) exhibits partial PLA-coir interfacial bonding. At a coir fiber content of 7 wt.% (Figure 11(c)), numerous cavities indicate limited matrix wetting and weak fiber-matrix adhesion. In addition, partial fiber breakage and fiber pull-out are observed, reflecting modest improvements in stress transfer and consistent with the findings reported by Dong et al. [28]. At a coir fiber content of 9 wt.% (Figure 11(d)), larger fiber clusters and agglomerates act as stress concentrators, reducing tensile strength and tensile modulus, as shown in Figures 8 and 9. These microstructural features facilitate crack propagation, disrupt interlayer fusion, and produce irregular fracture patterns, which are consistent with prior studies [65].

#### 5. CONCLUSIONS

The key finding of this study is that process-induced constraints more strongly influence the mechanical performance of FDM-printed PLA/coir biocomposites than fiber addition. SEM observations revealed interfacial voids, fiber pull-out, and incomplete interlayer fusion, which act as stress concentrators and limit effective load transfer. Consequently, despite enhanced crystallinity at low fiber contents, all PLA/coir composites exhibited lower tensile strength than neat PLA. At coir fiber contents of 5-7 wt.%, improved fiber dispersion and nucleation-induced crystallinity resulted in moderate increases in tensile modulus, whereas a higher fiber content of 9 wt.% promoted agglomeration and porosity, leading to reductions in both tensile strength and stiffness. These results demonstrate that FDM-related defects at higher fiber contents progressively outweigh the beneficial

effects of fiber reinforcement. The central implication of this work is that effective optimization of natural fiber-reinforced PLA for FDM applications requires the co-design of material formulations and printing parameters, as composite performance cannot rely solely on fiber incorporation but must also address extrusion stability, filament quality, and interlayer fusion to exploit sustainable biocomposites in additive manufacturing fully.

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