Multiscale Hybridization of Natural Silk–Nanocellulose Fibrous Composites With Exceptional Mechanical Properties

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Because of the crisis of the petrochemical era, environmentally friendly natural polymers and related material processes are receiving great attention. Cellulose and silk are typical fibrous materials that consist of polysaccharides and proteins, respectively, and have excellent mechanical properties and high physicochemical stabilities due to their unique self-assembly-based hierarchical structures. In this study, highly stable high-strength silk fiber (SF)–cellulose nanofibril (CNF) composites were prepared by the hierarchical fusion of micrometer-scale SFs and nanometer-scale CNFs. This manufacturing process is cost-effective because the raw materials (SFs and CNFs) are used as is. It is an eco-friendly process because it does not require the use of organic solvents or toxic reagents. In addition, it is an energy-efficient process because heat fusion (120°C) takes only 10 min. The results of the Direct Red 80 staining experiments confirm that up to 15 wt.% CNFs were added to the SF nonwoven. With the increase in the CNF amount, the nanometer-scale CNFs form a coating on the micro-scale SFs. At the same time, the CNFs form bonds with the SFs and increase the interfibrillar bonding strength of the CNF-coated SFs. Therefore, the mechanical properties of the SF/CNF composite and its stability in the water environment rapidly increase with increasing CNF concentration. In the case of SF/CNF15, the mechanical and impact strengths increase by 110 and 228%, respectively, compared with SF nonwoven without CNF. In addition, as CNFs are introduced, hydrophobicity of the surface and bulk of the SF/CNF composite can be imparted, thereby maintaining its structural stability in the water environment. This eco-friendly SF/CNF composite can be widely used as reinforcement preform of fiber-reinforced plastics as well as for other applications in the fibrous composite industry.

Keywords: silk fibroin, cellulose nanofiber, fibrous composite, nonwoven, adhesion

INTRODUCTION

Silk is an attractive natural fibrous material with a unique hierarchical composite structure (Shao and Vollrath, 2002; Vollrath et al., 2013; Koh et al., 2015). Its molecular structure consists of repetitive amino acid sequences (Malay et al., 2016), which form a secondary β-sheet-driven protein structure by the surrounding environment to form silk nanofibrils. The self-assembled nanofibrils
agglomerate to produce fibroin microfibers and, finally, a bicomponent nonwoven cocoon fabric in which sericin surrounds two fibroin strands (Keten et al., 2010; Omenetto and Kaplan, 2010; Asakura et al., 2015; Malay et al., 2016; Wang et al., 2018b). Based on its unique hierarchical structure, silk has excellent mechanical properties (Chen et al., 2012; Shah et al., 2014; Sanjay et al., 2018) and is resistant to physical attacks and harsh environmental conditions. In addition, silk cocoons produced by silkworms have the advantage that they occur under environmentally friendly process conditions such as room temperature and atmospheric pressure.

Because of their excellent biocompatibility and biodegradability as well as intrinsic mechanical rigidity, various regenerated silk materials have received increased attention (Han et al., 2017; Magaz et al., 2018; Yazawa et al., 2018; Zhang et al., 2018). However, regenerated silk materials have a significant disadvantage in that they are difficult to scale up (Lee et al., 2018).

During the regeneration process, environmental problems occur because of the use of organic solvents. In addition, molecular weight decomposition occurs in natural silk, leading to the deterioration of its physical properties (Cho et al., 2012; Kim et al., 2014; Lee et al., 2018; Yin et al., 2019). To overcome the above-mentioned disadvantages and effectively utilize silk materials, untreated (no regeneration) natural silk fibers (SFs) should be used (Peng et al., 2019).

Fibrous and fiber preform composites, which are produced by using natural SF as raw material, consist of SFs and additional chemical binders that increase the bonding between the SFs (Song et al., 2009; Yuan et al., 2010; Shubhra et al., 2011; Guang et al., 2015). Their mechanical properties greatly depend on the interaction between the binder and fibers. To improve the interaction between the binder and fiber, chemical modifications (Dányádi et al., 2010; Guo et al., 2016; Lee et al., 2019) (i.e., ionic, electrostatic, polar, and van der Waals forces exerted on the fiber and adhesive) and physical modifications of the fiber surfaces (Song et al., 2010), which increase the roughness and specific surface area of the fibers, have been widely used (Adekunle, 2015). However, these modifications lead to the physicochemical damage of the fibers and thus the mechanical weakness of the product if the modification time is extensive (Baiardo et al., 2004; Carvalho et al., 2010; Favaro et al., 2010; Das et al., 2015; Pandey et al., 2015). Another approach to increasing the interaction between the fibers is to fabricate hairy and fuzzy fibers based on the adhesion of finely sized nanomaterials to the surface of the fibers (Juntaro et al., 2008; Pommet et al., 2008; Lee et al., 2012; Arévalo and Peijs, 2016; Fortea-Verdejo et al., 2016).

Multiscale hybridized materials have received great attention by the scientific and industrial communities during the last decade due to their synergistic effects originating from different morphological scales (Duan et al., 2017; Lin et al., 2017; Toprakcioglu et al., 2017; Hadadi et al., 2018). In the composite materials approach, multiscale hybridization is an important key to solve inherent weaknesses such as the deterioration of the mechanical properties due to the poor reinforcement and structural dismantling in wet or aqueous environments (Fortea-Verdejo et al., 2016). In the biomedical fields, the production of multiscale materials can be used to create an effective drug delivery system (Numata et al., 2012; Gao et al., 2019), enabling sustainable and ultrafast drug delivery. At the same time, based on the nano-topologies of the microsurfaces (Mozunder et al., 2011), superhydrophobic materials can be produced. In addition, injectable scaffolds with reinforced mechanical strengths can be prepared by incorporating nano and microparticles into three-dimensional hydrogels (Xiao et al., 2017). Therefore, the fabrication of multiscale hybridizing fibrous composites using natural micro- and nanoscale fibrous materials is a simple and environmentally friendly fabrication method with a synergistic structure–property relationship.

Wood is the most abundant natural resource on Earth and a cellular hierarchical biocomposite (Eichhorn et al., 2010). The self-assembly of hydrogen bond-based repetitive glucose units results in the formation of elemental fibrils, which develop into microcrystalline cellulose (MCC) (Tayeb et al., 2018). The MCC forms a macroscopic biocomposite structure that reinforces the matrix consisting of lignin, hemicellulose, and wax, eventually forming a whole tree with a scale of tens of meters. Various natural raw materials can be obtained from such a complex hierarchical structure of woody biomass (Klemm et al., 2005, 2011; Cervin et al., 2013). Such raw materials have been used to manufacture paper, (Song et al., 2009) pulp, (Theng et al., 2015) fiber, which greatly contributed to human development. Recently, the development of advanced materials using nanocellulose has been actively attempted because of the robust physicochemical properties of nanocellulose, which are based on its unique morphology and crystalline structure (Theng et al., 2015; Amini et al., 2017; Saba et al., 2017b,a; Tayeb et al., 2018; Feng et al., 2019; Kwak et al., 2019b).

To fabricate such a natural multiscale hybridizing polymer fibrous composite, SFs produced by silkworms and cellulose nanofibrils (CNFs) obtained from the woody biomass were used as natural micro and nanoscale fibrous raw materials, respectively. In this study, natural polymeric hierarchical fibrous composites were fabricated by using filtration and thermal fusion methods typically used in conventional paper manufacturing. The successful fabrication of hierarchical nano and microscale hybridization structures was verified using dye stains and the interactions between the nano and microscale fibers were confirmed using disintegration observations. Finally, the mechanical properties and water stability depending on the CNF concentrations of the natural multiscale hybridized fibrous composites were determined.

MATERIALS AND METHODS

Materials

Silk cocoons of the Bombyx mori silkworm were kindly provided by the National Academy of Agricultural Science (NAAS, Korea). Bamboo fibers were obtained from Damyang (2–3 years old, Phyllostachys, Korea) and used as native cellulose fibers. Direct Red 80 was obtained from Tokyo Chemical Industry Co., Ltd., Japan. Na₂CO₃ was purchased from Sigma-Aldrich (Korea).
Degumming of Silk Fibers (SF)
The silkworm cocoons were boiled for 30 min in an aqueous solution containing 0.02 M sodium carbonate (Na₂CO₃, > 99.0%, Sigma-Aldrich) to remove sericin and other impurities. After washing the samples several times with deionized water (DI) water, the resulting protein SFs were dried for 3 days at 25°C. Next, the degummed SF fibers were wound onto a small mandrel to yield 6 cm long aligned fibers.

Preparation of Cellulose Nanofibril (CNF)
To remove the waxy material from the bamboo, 2 g of the bamboo fiber was dewaxed on a Soxhlet equipment (SE-Series, Vinci Technologies, France) with a benzene/ethanol [2: 1 (v/v) mixture of] for 6 h. The lignocellulosic material in the sample was then removed using a sodium chlorite (NaClO₂) solution at 75°C for 1 h and repeated same process five times until the color of the sample became white. The residual hemicellulose, starch and pectin were further removed at 90°C for 2 h with 2 wt% potassium hydroxide (KOH). Thereafter, bamboo cellulose dispersion (0.05 wt%) was prepared by adjusting the concentration of the dispersion with distilled water. About 200 mL of the bamboo cellulose dispersion was placed in an ultrasonic generator (Kyungil Ultrasonic Co., Korea), and subsequent sonication (800 W, 4 s on/off pulse) was performed for 1 h to obtain disintegrated cellulose nanofibrils. In order to avoid temperature elevation during high intensity ultrasonic processing, sonication was performed in an ice/water cooling bath system. In order to confirm nanofibril preparation, morphological image of bamboo CNF was obtained using field emission scanning electron microscopy (FE-SEM, AURIGA, Carl Zeiss, Germany), and determination of nanofibers diameters were performed with “Image J” software.

Fabrication of SF/CNF Fibrous Composites
To obtain a homogeneous CNF dispersion, 1 wt.% CNF solution was sonicated at 300 W, 4 s on/off pulse under the ice/water bath condition. Then, 0–1 wt.% of CNF solutions were diluted with the distilled water. SF fibers (1.6 g) were added to the desired CNF solution and incubated for 10 min to fully wet the CNF dispersed solution within the SF fibers. The SF/CNF suspension was poured into a metal mold and homogenously distributed. Thereafter, a paper towel was used to remove excessive water from the SF/CNF mixture in the mold (six times). The moisture-free SF/CNF mixture was placed onto a hot press machine (HLP-12H, Han tech, Korea) and the metal mold was removed. Subsequently, the SF/CNF mixture was pressed at 120°C for 10 min to fabricate a SF/CNF fibrous composite. A polyester nonwoven was placed on the top and bottom of the SF/CNF mixture to avoid the adhesion of the SF/CNF to the plate of the hot press. The overall process of manufacturing SF/CNF composites is shown in Scheme 1. As a control, untreated nonwoven silk was prepared using the above-mentioned pressing method at room temperature. The resultant SF–CNF composite was labeled SF/CNF-n, where “n” corresponds to the initial CNF content (n = 5, 10, and 15 wt.% of SF, respectively).

Characterization
To determine the CNF distribution on the SF surface, the prepared SF/CNF fibrous composite was immersed in 60 mL of Direct Red 80 dye solution (0.0025 wt.%) and incubated for 30 min. Thereafter, distilled water was used to remove the unreacted dye molecules from the sample until no red color was observed on the paper towel, which was pressed onto the SF/CNF fibrous composite sample. The washed fibrous composite was incubated in the oven at 60°C for 1 day and images of the dried sample were obtained. To obtain the Direct Red 80-dyed CNF sample, 1 wt.% of CNF solution was cast on a Petri dish and lyophilized using a freeze dryer. Subsequently, sponge-type CNFs were obtained by pressing at 120°C for 10 min. The above-mentioned dyeing process was also used for this sample and an image was obtained.

Optical images of the prepared SF/CNF composites were obtained using a digital camera (D7200, Nikon, Japan). The morphologies of the SF/CNF composites were analyzed using field emission scanning electron microscopy (FE-SEM, AURIGA, Carl Zeiss, Germany). To obtain cross-sectional images of the SF/CNF composites, the samples were placed in a liquid nitrogen chamber for 6 h and then cut using a cryo-blade. To find out coating and interfacial adhesion behavior of nanocellulose onto the SF fibers, 1.6 g of SF fibers were immersed onto the 40 ml of cellulose nanocrystal (CNC) and CNF solution (0.5 wt%) and it dried 60°C for 24 h. The prepared SF/CNC and SF/CNF composites were immersed the 50 ml of distilled water and it sonicated 200 W for 20 min to detach CNC and CNF from the SF fibers. Then morphology of the SF/CNC and SF/CNF before and after ultrasonication treatment were analyzed using FE-SEM.

The X-ray diffraction (XRD, Rigaku, DMAX 2500) analyses were conducted over a 2θ range of 5°–60° at 40 kV and 100 mA using Cu Kα radiation (λ = 0.154 nm). An attenuated total reflectance Fourier transform infrared (ATR-FTIR) spectrometer was used to examine the chemical structures of the SF/CNF composites. The scan range, scan number, and resolution of the measurements were 4000–700 cm⁻¹, 128, and 2 cm⁻¹, respectively. The absolute density (ρm) of the manufactured SF/CNF composites was measured with a He pycnometer (AccuPyc 1330, Micromeritics, United States). The samples were weighed and then loaded in the measurement chamber. The envelope density (ρe) of the fibrous composites was calculated from the mass and envelope volume of the nonwoven samples. The porosity of the fibrous composite was calculated using Eq. (1):

\[
\text{Porosity (%) = } \left(1 - \frac{\text{Envelope density}}{\text{Absolute Density}}\right) \times 100
\]

To determine the mechanical properties of the SF/CNF composite, tensile tests were performed using a universal testing machine (CS225, Lloyd Instruments, Ltd., United Kingdom). The experiment was conducted by applying a tensile force in accordance with the main arrangement direction of the SF fibers. The gauge length and width of the samples were fixed to 50 and 25 mm, respectively, and tensile tests were performed using a 1000 N load cell at an extension rate of 0.05%. The results were analyzed and the averaged values were presented.
10 mm/min. All samples were preconditioned at 20°C and a relative humidity of 65%. The Izod impact strength of the unnotched fibrous composite specimens was measured with a (QM700A, Qmesys, Korea) machine at a strike velocity of 2.5 m/s according to the ASTMD256 test method. The experiment was conducted by applying an impact force to vertical direction of SF fiber's alignment. At least five samples were tested under each condition and the averages and standard deviations of the tensile strength, elongation at break, and initial Young's modulus were evaluated. To study the physical stability of the SF/CNF composites in a water environment, water droplet images were obtained 1 s after placing a water droplet on the fibrous composite surface. To investigate the stability under hydrolytic conditions, the SF/CNF composites were immersed in 60 mL distilled water for 21 days. Subsequently, optical images of the fibrous composites were obtained.

To determine the moisture absorption behavior of the SF/CNF composites, a water uptake test was performed (de Campos et al., 2013). The sample size was fixed to 60 mm (length) and 25 mm (width). All samples were preconditioned at 25°C and a relative humidity of 51%. Thereafter, the samples were immersed in distilled water. Their weights were determined before and after immersion. At least five samples were tested under each condition and the averages and standard deviations were obtained. The water uptake was calculated using Eq. (2). The parameters $W_0$ and $W_t$ are the initial weight and the weight after immersion for 24 h, respectively.

$$\text{Water uptake (\%) = } \frac{(W_t - W_0)}{W_0} \times 100 \quad (2)$$

**RESULTS AND DISCUSSION**

Using the chemical purification and high intensity ultrasonication process, bamboo-derived cellulose nanofibril (CNF) was successfully prepared. As a result of morphology confirming through the FE-SEM and FE-TEM image, the formation of high aspect ratio CNFs having 15–35 nm of diameter (average diameter: 26.1 nm) was observed (Figure 1).

In order to examine the effect of nonwoven fabrication processing on the silk fibroin (SF) fiber, SF nonwoven fabric was prepared by simply pressing without treatment. Figure 2A shows the image of the degummed SF (sericin removed) and pressed SF nonwoven without heat treatment. The degummed SF shows a three-dimensional cotton candy shape, and it can be confirmed that such SF can be manufactured as a two-dimensional like nonwoven fabric through a simple pressing process. It was found that the pressing process did not change the color of the SF fibers themselves. Meanwhile, images of SF nonwoven and CNF added SF/CNF composite nonwoven prepared by hot pressing method are shown in Figure 2B. In the case of CNF, the hot-pressed CNF film remains white color even though manufactured by the hot-pressing method. However, in the case of SF, prepared hot-pressed silk non-woven is converted to pale yellow. Generally, a yellowing phenomenon can mainly be observed when protein crosslinking (both self-crosslinking and crosslinking with a chemical crosslinker) occurs. During the fabrication of nonwoven SFs, protein self-crosslinking occurs, resulting in a yellowing phenomenon (Lee et al., 2018). Urreaga and de la Orden (2006) prepared chitosan treated cotton cellulose fibers to examine the interaction between amino group of chitosan and cellulose carbonyl groups and found that the higher the amount of added chitosan, the more conjugated imine groups produced, resulting in yellowing behavior of cellulose. However, it is difficult to observe a significant color difference in the SF/CNF composite compared to the SF heat-treated nonwoven. Therefore, the yellowing phenomenon occurring in the SF/CNF composite is largely affected by the crosslinking of the SF fiber itself.

To confirm the effect of CNF concentration on the residual CNF contents in the SF/CNF composites, the weight gain ratio of SF/CNF were calculated based on the dry weight and the obtained result was shown in Figure 2C. As shown in the figure, as the CNF content increases to 15% of the SF weight, the weight gain ratio of the SF/CNF composite also increases proportionally. However, it can be seen that there is a limit to the amount of CNF that can penetrate into the SF nonwoven.
fabric by reduced weight gain ratio of SF/CNF20 and SF/CNF25 compared to SF/CNF15.

In the degumming process, which is the removal of silk sericin from cocoon silk, a Direct Red 80 staining test is widely used to confirm the removal of the gummy sericin protein (Nisal et al., 2014; Kwak et al., 2017). Depend on the hydrophilicity of silk proteins (fibroin and sericin), Direct Red 80 easily stains the gummy and hydrophilic sericin protein, while degummed hydrophobic fibroin fiber cannot be dyed (Kwak and Lee, 2015). In addition, (Saleem et al., 2007) confirmed that the Direct Red 80 dye is easily dyed to cellulose cotton fibers. By using the different dyability between cellulose and silk fibroin, the distribution of CNF in SF/CNF composite was visualized and the optimal CNF content was found. The Direct Red 80 dyed fibrous SF/CNF composite samples are shown in Figure 2D. As shown in the figure, Direct Red 80 does not dye nonwoven SF, probably because the SFs are hydrophobic and the dye molecules do not penetrate the β-sheet-dominated SF structure. On the other hand, in the case of the nonwoven CNF fabric, the Direct Red 80 dye penetrates the fabric and a vivid pink color can be observed. This is the result of dyeing through hydrogen bond with cellulose chain after the Direct Red 80 dye dissociates under neutral aqueous conditions. The CNF concentration of the SF/CNF composite can be simply determined based on the dyeing performance observed during the Direct Red 80 staining test. As the CNF content of the SF/CNF composite increases from 5 to 15 wt.%, the color of the fibrous composite surfaces becomes more vivid (pink). In the case of the SF/CNF5 composite, a white part can be observed because the amount of CNF between the SFs is not high enough. In the case of SF/CNF10 and SF/CNF15, the surfaces of the SF fibrous composites are fully covered with CNF molecules, which results in uniform pink layers. In addition, as the CNF weight increases, the color becomes more vivid, indicating that the CNF concentration of the SF nonwoven is higher. However, SF/CNF20 and SF/CNF25 are partially pale pink similar with SF/CNF5 despite the increase in the CNF
Between the SFs. CNF concentration is due to the fact that the CNF does in the porosity of the SF/CNF composite with increasing the void filling effect of the sericin proteins. The decrease of the porous structure of the jute fiber composites due to adhesion of jute fibers and observed a similar offset phenomenon preform composites using sericin as bio-adhesive for the

Supplementary Table S1

$\pm$ two density measurements. The CNF content increases; porosity The calculated porosity was obtained based on the results of

Figures 3A–C and absolute densities, as shown in Figures 3A–C and Supplementary Figure S1.

To investigate the macroscopic changes of the nonwoven SF structure after the combination of the CNFs with SFs, the envelope and absolute densities of the SF/CNF composites were measured. The porosities of the SF/CNF composites were calculated based on the relationship between the envelope and absolute densities, as shown in Figures 3A–C and Supplementary Table S1.

The absolute density ($\rho_m$), commonly referred to as bulk density, is the ratio between the mass and the unit volume of a sample. Voids or pores in the material are not considered. The absolute density insignificantly changes, even if CNF is added to the SF structure. On the other hand, the envelope density ($\rho_e$), defined as the ratio between the mass of the material and the sample volume, includes voids or pores in the material. As the CNF content increases, the envelope density of the SF/CNF composite increases from 0.79 $\pm$ 0.03 for nonwoven SF to 1.02 $\pm$ 0.02 for the SF/CNF15 composite. The calculated porosity was obtained based on the results of two density measurements. The CNF content increases; porosity decreases from 40.91 $\pm$ 1.08 (sf nonwoven) to 25.91 $\pm$ 1.36 (SF/CNF15). Kwak et al. (2019a) fabricated jute fiber–sericin preform composites using sericin as bio-adhesive for the adhesion of jute fibers and observed a similar offset phenomenon of the porous structure of the jute fiber composites due to the void filling effect of the sericin proteins. The decrease in the porosity of the SF/CNF composite with increasing CNF concentration is due to the fact that the CNF does not only form a coating on the SF but also fills the space between the SFs.

To identify the effect of the CNF binder on the morphological changes of the SF/CNF composites, surface and cross-sectional FE-SEM images were obtained. The results are shown in Figures 4e–h. As the CNF content increases, the gaps between the SFs are filled and the sizes of the pores decrease. This effect can be clearly observed in the FE-SEM cross-sectional images with higher magnification. In the case of SF nonwoven, the boundary between the SFs can be clearly observed, whereas in the case of the SF/CNF15 composite, CNFs are observed between SFs, which makes the boundary between fibers unclear. Nevertheless, pores between the SFs can be observed, indicating that the CNFs reduce the porosity of the SF structure but do not fill the entire pore structure of the SF matrix.

To investigate the chemical structure and crystalline properties of SF/CNF composites after CNF addition, FTIR spectra and XRD patterns were obtained. The results are shown in Figure 5. The FTIR spectra of the SF/CNF composites are shown in Figure 5A. In the spectrum of neat CNF, an O-H stretching mode at 3300 cm$^{-1}$, C-H stretching mode at 2800 cm$^{-1}$, and C-O stretching peak at 1013 cm$^{-1}$ can be observed, which are typically associated with cellulose structures. In the case of SF/CNF composites, the SF shows a crystalline secondary $\beta$-sheet structure typical for SF proteins. Regardless of the CNF content, the spectra of all SF/CNF composites show amide I (C = O stretching) and amide II (N-H bending and C-N stretching) peaks at 1620 and 1510 cm$^{-1}$, respectively, which are associated with the secondary $\beta$-sheet structure of SF (Jackson and Mantsch, 1995; Lu et al., 2010; Nagarkar et al., 2010; Boulet-Audet et al., 2015). This means that the CNFs do not induce the conformational transition of the $\beta$-sheet structure of SF, even if CNFs are introduced to the SF structure. In general, silk-based materials, including natural and regenerated SFs, have excellent mechanical properties when the silk materials exhibit a $\beta$-sheet structure. Therefore, maintaining the secondary

**FIGURE 3** Absolute density (A), envelope density (B), and calculated porosity (C) of the SF/CNF composites with various amounts of CNFs.
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**FIGURE 4** | FE-SEM images of the SF/CNF composites with various CNF concentrations. Surface (a–d) and cross section (e–h) images; (a,e): SF nonwoven, (b,f): SF/CNF5, (c,g): SF/CNF10, (d,h): SF/CNF15. The insets represent high-magnitude images.

**FIGURE 5** | Characteristics of CNF, nonwoven SF, and SF/CNF composites with various amounts of CNFs: (A) FTIR spectra, (B) XRD pattern.

The β-sheet structure of the SF, even when CNF is added, will have a positive effect on the mechanical properties. On the other hand, as the CNF content increases, the intensity of the C-O stretching absorption peak at 1013 cm\(^{-1}\) increases,(Morán et al., 2008; Han et al., 2013; Feng et al., 2019) implying the settling of CNF on the surface of the SF. Furthermore, in the spectra of the SF/CNF composites, the O-H peak of SF merges with that of CNF. The spectra exhibit a blue-shifted broader peak, indicating the interaction between SFs and CNFs through the hydrogen bonds (Lee et al., 2013).

The XRD patterns of the SF/CNF composites are shown in Figure 5B. The nonwoven SF exhibits a peak at 20.2°, which corresponds to the silk II β-sheet crystallite structure (Lee, 2004; Cheng et al., 2015; Guang et al., 2015; Ming et al., 2015). In the case of CNF, characteristic peaks at 20 = 16.3° and 22.6° are observed, representative for the typical cellulose type I structure (Morán et al., 2008; Han et al., 2013; Kwak et al., 2019b). Based on the XRD patterns of the SF/CNF composites, the CNF binder affects their crystalline structures. In the case of SF/CNF5, because the CNF does not cover the whole SF surface, the crystal structure of SF appears to be sharp. On the other hand, in the case of SF/CNF10, the crystal structure appears to be mixed, but the peak at 20.2°, which represents the crystal structure of silk, is more dominant. In the case of SF/CNF15 with a higher CNF content, the peak at 22.6°, which corresponds to CNF, is more prevalent. Based on the FTIR and XRD results, the introduction of CNF does not lead to the transfer of the β-sheet structure of SF, but the crystal structure of the entire SF/CNF composite changes.

To visually identify the interaction between CNF and SF, a ultrasonication detaching test was applied using nanocellulose coated SF fibers. To find out the effect of nanocellulose morphology on the interaction between SF and nanocellulose, cellulose nanocrystal (CNC) with a low aspect ratio and CNF with a high aspect ratio were used as a SF coating agent. Thereafter, bonded nanocellulose onto SF fibers were detached by ultrasonic treatment, which is a strong physical treatment technique. Figure 6a shows the morphology of a freeze-dried CNC. It can be seen that the aspect ratio of CNC is lower than that of CNF as shown above image (Figure 1a). Also, as shown in Figure 6b, neat silk fibers have a smooth surface. Figure 6d show that the SF has a smooth surface, while the CNC-coated SF has a rough surface. This smooth surface of SF changed to rough with a short aspect ratio of CNC covering (Figure 6c). The rough surface of the CNC-coated SF indicates that the CNC covers the surface of the SF and forms a core–shell-like structure. Although the roughness decreases after sonication, the CNC-coated SF surface is rougher than that of the conventional SFs. This means that more than just physical adsorption occurs between CNC and SF. Compared with CNC, CNF has a longer aspect ratio, so it can be applied not only for simple coating but also for interfacial adhesion between SF fibers. In order to confirm the adhesion behavior of CNF, CNF coating was applied
to SF fibers, and FE-SEM images of SF/CNF fibers subjected to sonication to separate them are shown in **Figures 6e,f**. As can be seen in the figure, the CNF bundle wraps around the SF fibers, so the SF fibers have a rough surface. Not only this, you can clearly observe the presence of CNF bundles between SF fibers. Through this, when the CNF is introduced onto the SF fibers, CNF can play not only the simple coating reagent but also the interfacial adhesive of the SF fibers. On the other hand, similar to the CNC, defibrillation and detachment of CNF from the SF fibers occurred by strong ultrasonication treatment, but nanoscaled CNF bundles exist, indicating that there is a certain binding force between CNF and SF. However, further consideration of the exact mechanism between CNF and SF is needed.

Based on the introduction of CNF, which allows the simultaneous coating and binding of SFs, SF/CNF composites with excellent mechanical properties can be easily fabricated. To confirm this effect, the mechanical properties of SF/CNF composites with various amounts of CNFs were investigated using impact and tensile tests. The results are shown in **Figure 7** and **Supplementary Table S2**. **Figures 7A,B** present the average tensile strength and modulus of SF/CNF composites with the various amounts of CNFs, respectively, and indicate the improved tensile behavior of the SF/CNF composites. The SF/CNF fibrous composite with 5 wt.% CNF (SF/CNF5) exhibits a 60 and 61% increase in the tensile strength and modulus, respectively, compared with nonwoven SF. The mechanical reinforcement further enhances with increasing CNF content. The SF/CNF15 composite shows a 110% and 150% increase in the tensile strength and modulus, respectively. The impact strength test results (**Figure 7C**) confirm the mechanical reinforcement effect of CNF. Based on the impact tests, the SF/CNF5 displays a ~53% increase in the impact strength compared with SF nonwoven, whereas SF/CNF10 and SF/CNF15 show an increase in the impact strength of ~109 and ~228%, respectively. These data prove that CNF can reinforce the low impact strength of nonwoven SF and that higher CNF concentrations can accentuate this trend. The mechanical reinforcement phenomenon of SF/CNF composites can be explained as follows. First, because the introduction of CNF does not affect the secondary β-sheet structure of SF, the mechanical properties of the SFs can be maintained, even after the preparation of the SF/CNF composites. In addition, by coating CNF onto the SF surface, a core–shell-structured SF/CNF fiber forms, which promotes the hydrogen bonding between the microscale SFs and nanoscale CNFs. In addition, as the amount of CNF increases, high-aspect ratio CNFs can form an interfibrillar bridges between the hybrid fibers of the SF/CNF composites.
thereby improving the interfacial adhesion characteristics of the SF/CNF composites.

The stability of the fibrous structure including the nonwoven fabric in wet or high-humidity environments is a very important property. Because most of the natural fibers are hydrophilic, composite materials with natural fibers as reinforcing agents often have disadvantages in that they are more vulnerable to moisture environments than materials without a reinforced matrix (Fages et al., 2013; Theng et al., 2015, 2017). Many natural polymer-based CNF-reinforced composites fabricated using common blending methods show hydrophilic properties such as a decrease in the water contact angle (WCA) (Kwak et al., 2018; Wang et al., 2018a), increasing water uptake or absorption capacity, and higher swelling ratio with increasing CNF concentration. On the other hand, when the surface coating and layer-by-layer coating method are used, the CNF-reinforced composite materials have hydrophobic properties because CNF actively affects the surface morphologies or topologies of the composite materials. The WCA is used as an indicator for the degree of surface hydrophobicity or hydrophilicity of film, textile, and membrane materials. Water droplet images representative of the SF/CNF fibrous composites and the average WCA results are shown in Figures 8A,B. Generally, a film with a WCA above 65° is considered as hydrophobic (Vogler, 1998). The WCA of the nonwoven SF without CNF is ~32.5°, which indicates that the nonwoven SF is hydrophilic. In contrast, when CNF is added to the SF structure, the WCA significantly increases. In the case of SF/CNF15, the contact angle is ~69.4°, which means that the CNF coating on the SF surface and CNF bonding with SFs sufficiently hydrophobizes the surfaces of the SF/CNF composites. The hydrophobic effect of CNF is due to the change in the surface topology of the entire SF/CNF composite.

In addition, the hydrostability of SF/CNF composites in wet environments was also examined using a water uptake test. The results are shown in Figure 8C. As the CNF content increases, the water uptake of the SF/CNF composites decreases (Fages et al., 2013; Theng et al., 2015, 2017). This might have the following reasons: (1) the initial diffusion rate of the water molecules decreases because of the aforementioned surface hydrophobicity, and (2) by introducing CNF into SF fibrous systems, the porosity of the SF/CNF composites decreases, thus reducing the space for water impregnation. Finally, the increase in the hydrogen bonding between SF–CNF and CNF–CNF reduces the affinity to water molecules. The hydrophobic effect of CNF greatly contributes to the morphological stability of the SF/CNF composites in wet environments. As shown in Figure 8D, nonwoven SF without CNF shows that each SF disintegrates in a 4-week stirring environment, causing the loss of the dense SF structure. In contrast, the dense nonwoven structure of the SF/CNF15 composite can be well maintained, even after 4 weeks of stirring in an aqueous environment.

**CONCLUSION**

In this study, natural micro- and nano-fiber fused fibrous composites were fabricated using silk and cellulose as green raw materials. Robust SF/CNF composites were fabricated using a simple short-term thermal bonding process. This process is eco-friendly and energy-efficient because it does not require an organic solvent, additional chemical reagents, long processing time, and high energy consumption. The introduction of CNF to SF increases the interfibrillar bonding of SFs as well as the coating on the SF surface. As the CNF content increases, the coating and bonding effects of CNF rapidly increase. Therefore, the mechanical properties and structural stability of the SF/CNF composites increase compared with that of nonwoven SF without CNF. A maximum of 15 wt.% CNF can be added to the SF/CNF composite. The SF/CNF15 composite shows a 110 and 225% increase in the impact and tensile strength, respectively, compared with nonwoven SF. The results also show that the hydrophobicity of the SF/CNF composite
surface and bulk materials can be imparted. Such green and facile SF/CNF composites have immense potential for a wide range of applications, particularly in the nonwoven and fiber-reinforced plastics industries.

**DATA AVAILABILITY STATEMENT**

All datasets generated for this study are included in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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**SUPPLEMENTARY MATERIAL**

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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