Development of cellulose-based conductive fabrics with electrical conductivity and flexibility

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Abstract

This study aimed to produce cellulose-based conductive fabrics with electrical conductivity and flexibility. Bacterial cellulose (BC) and three chemical cellulose (CC), namely methyl cellulose (MC), hydroxypropyl cellulose (HPMC) and carboxymethyl cellulose (CMC) were in situ polymerized with aniline and the four conductive cellulose fabrics were compared and evaluated. Matrix-assisted laser desorption/ionization time-of-flight mass spectroscopy analysis confirmed that three CC-PANI composites displayed longer and more stable polymerization pattern than BC-PANI because of the different polymerization method: bulk polymerization for BC-PANI and emulsion polymerization for CC-PANI, respectively. The electrical conductivity of BC-PANI and CC-PANI were ranging from $0.962 \times 10^{-2}$ S/cm to $2.840 \times 10^{-2}$ S/cm. MC-PANI showed the highest electrical conductivity among the four conductive cellulose fabrics. The flexibility and crease recovery results showed that MC-PANI had the highest flexibility compared to BC-PANI, HPMC-PANI, and CMC-PANI. These results have confirmed that the electrical conductivity and flexibility were influenced by the type of cellulose, and MC-PANI was found to have the best performance in the electrical conductivity and flexibility.

Introduction

Conductive fabrics are electrically functionalized fabrics with the advantages of flexibility, elasticity, and wearability [1, 2]. These features have enabled conductive fabrics to be applied for textile-integrated batteries [3], wearable sensors [4], and fabric-based energy storage devices [5]. Conductive fabrics can be manufactured by the weaving and knitting of conductive threads [6] or by synthesizing conductive polymers onto the fabric material [7]. The synthesis method is commonly used because of the simple manufacturing process [8]. Conductive
polymers, such as polypyrrole, polyaniline, and polythiophene, are organic-based polymers with extended π-orbital systems along their conjugate backbone through which electrons can move freely [9]. Due to their unique structure, conductive polymers are lighter and better resist corrosion than metals [10].

Among the various conductive polymers, polyaniline (PANI) has been widely studied because of its good chemical stability, good electrical conductivity, and ease of synthesis by the in situ polymerization of aniline [11–13]. In this study, PANI was polymerized onto cellulose to produce the conductive fabric. However, PANI has poor mechanical properties since it is inherently very stiff [14]. To overcome the disadvantage of PANI, a flexible fiber material is required to be synthesized with PANI [15].

Cellulose is one of the most abundant natural fibers. It is composed of glucose chains that contain many hydroxyl (-OH) functional groups [16]. It has excellent physical properties, such as flexibility, and is biodegradable [17]. Because of these properties, cellulose is considered as a good material to produce conductive fabrics. The characteristics of cellulose vary depending on its origin [18]. Presently, bacterial cellulose (BC) and chemical cellulose (CC), especially methyl cellulose (MC), carboxymethyl cellulose (CMC), and hydroxypropyl methyl cellulose (HPMC) were used. BC is produced Acetobacter xylinus bacteria. BC has a three-dimensional nanostructure with pronounced biocompatibility and purity [19, 20]. MC is derived from chemical cellulose without loss of biocompatibility [21]. It is formed by replacing the hydroxyl groups of cellulose with methoxy groups through chemical modification to improve the water solubility of cellulose [22]. The ease of water-based MC modification suggests the possibility of wet spinning and ink material produced by three-dimensional printers. Other advantages of MC include transparency, elasticity, and mechanical strength [22]. Also, HPMC and CMC were used to compare those electrical conductivity to the one of MC. These three CC have similar chemical structures but MC has methyl group instead of cellulose hydroxyl groups whereas HPMC has methyl and hydroxyl propyl groups and CMC has carboxymethyl group. HPMC and CMC also have the properties of reasonable biocompatibility, water solubility, transparency and elasticity [23].

Many studies have explored the production of conductive BC-PANI by polymerizing aniline with BC. However, little is known concerning the polymerization of aniline with CC [24, 25]. Moreover, no study has investigated the production of CC-PANI as a conductive fabric, or compared and evaluated BC-PANI and CC-PANI.

The aim of this study is to produce the conductive cellulose fabric with electrical conductivity and flexibility. BC-PANI and three CC-PANI composites are produced, and then compared to select a conductive cellulose fabric with better performance. The polymerization conditions are optimized by ultraviolet-visible (UV-Vis) spectroscopy, and the degree of polymerization is observed with matrix-assisted laser desorption/ionization time-of-flight mass spectroscopy (MALDI-TOF MS). The electrical conductivity, flexibility and crease recovery of conductive cellulose fabrics are evaluated. Chemical and physical structures of conductive cellulose fabrics are evaluated using Fourier-transform infrared spectroscopy (FT-IR) and X-ray diffraction (XRD). Lastly, surface morphology is observed by field-emission scanning electron microscopy (FE-SEM).

**Materials and methods**

**Materials**

MC powder with a viscosity of 4,000 cP (in 2% solution) and HPMC powder with a viscosity of 4,260 cP (in 2% solution) were obtained from LOTTE Fine Chemical Co., Ltd (Incheon, Korea). CMC sodium salt with a viscosity of 50 to 200 cp (in 4% solution) was obtained from (MSIT) (Grant No. 2018R1C1B5044988), and the 2019 Hongik University Research Fund, Korea.

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Sigma Chemical CO., (St. Louis, MO, USA). Yeast extract and peptone were purchased from BD Biosciences (San Jose, CA, USA). Glucose, hydrogen peroxide (34.5%), sodium hydroxide, ethyl alcohol (99.9%) and acetic acid were supplied by Duksan Pure Chemical Co., Ltd. (Seoul, Korea). Dodecylbenzene sulfonic acid (DBSA) and trifluoroacetic acid (TFA) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Aniline and acetonitrile were purchased from Junsei Chemical Co., Ltd. (Tokyo, Japan). Ammonium peroxydisulfate (APS) was supplied by Kanto Chemical Co., Inc. (Tokyo, Japan). All chemical reagents were used as received.

Production of BC-PANI

BC was produced according to a previously reported method [26] and was pre-treated by following the method of Song et al. [27]. Briefly, *Acetobacter xylinus* inoculum was prepared with Hestrin-Schramm (HS) medium (20 g/L glucose, 5 g/L yeast extract, and 5 g/L peptone) and the medium was boiled at 100˚C for 10 min. After that, BC was statically cultured at 26˚C for 8 days until the thickness of BC became 1 cm. Afterwards, BC was washed with a 3% sodium hydroxide solution at 25˚C for 90 min with shaking at 50 rpm, neutralized with distilled water (DW), and adjusted to pH 3.0 using acetic acid for 30 min. After that, BC was bleached with a 5% hydrogen peroxide solution at 90˚C for 60 min with shaking at 110 rpm, swelled with an 8% sodium hydroxide solution in a liquor ratio of 1:10 (w/v) for 30 min in an ultrasonic bath, and dried at 35˚C.

BC-PANI was produced by the bulk polymerization process by following the method of Kim et al. [28] (Fig 1(A)): First, 2.5 mM aniline and pre-treated BC were added to a 2.5 mM DBSA solution of pH ranging from 2.0 to 4.0. After that, the aniline, BC, and DBSA mixture was magnetically stirred for 1 h. Next, a 2.5 mM APS solution was added to the reaction mixture whereby polymerization was initiated. After several hours at varied temperatures, ranging from 10 to 30˚C, BC-PANI was synthesized. The reaction product was washed with an ethanol solution to remove the residue and dried at 25˚C for 4 h to produce the nonwoven BC-PANI fabric.

Production of CC-PANI

CC-PANI was produced by the emulsion polymerization process (Fig 1(B)): A solution of 25 mM DBSA was added dropwise into 9 mL of cold DW without mixing to avoid self-micelle formation. After adding 25 mM aniline, the solution was strongly vortexed for 90 sec until its appearance changed to that of a milky white emulsion. The resulting emulsion was incubated for 30 min in an ice bath, and then, polymerization was triggered by adding 1 mL of a 25 mM APS solution. Following a 4-h incubation period, the solution color turned to dark green owing to the formation of the polyaniline-emeraldine salt. At that point, 0.5 g of each CC powder was completely dissolved in 10 mL of the PANI solution. The MC and PANI mixture was transferred to a petri dish and vacuum was applied to remove extra air bubbles, followed by drying at 4˚C for 24 h to form the CC-PANI fabric without wrinkling. All chemical components and conical tubes were cooled down before use.

The production of CC-PANI by mixing DBSA, aniline, APS, and CC powder, was optimized with respect to the order of mixing as well as the amount of each component (Fig 1(B)): At first, we confirmed the gelation of the CC powder, necessary to generate sheet-type CC. When we mixed CC powder with DW and heated the mixture for 10 min, solution was observed. After that, we cooled down the sample to room temperature, and confirmed the sol-gel transition. As the hydrogen bonds between MC polymer chains and water molecules were broken at high temperature, the methoxy groups became exposed [23]. Thus, hydrophobic interactions could occur, resulting in the formation of a gel network. Bearing this in mind, we
immediately poured the mixture after heating from a glass vial onto a plate, and cooled it down to 4°C, to produce sheet-type CC. Based on this result, we adopted the following procedure: First, we added aniline to DBSA in water and then mixed the solution with APS. After the color of the solution turned to dark green, we added CC. Finally, we poured the aqueous CC-PANI solution onto a plate and dried it for further characterization.

**UV-Vis spectroscopy**

UV spectra of BC- and CC-PANI polymer solutions were acquired with a 96-quartzo microplate reader (SynergyMx, Shimadzu, Japan) for BC-PANI or a cuvette type reader (Thermo Labsystems Multiskan Spectrum, Vantaa, Finland) for CC-PANI, in the absorbance spectrum range from 300 to 800 nm with a resolution of 2 nm. The solutions were diluted 50 times with Milli-Q water before spectrum acquisition.

**MALDI-TOF MS analysis**

The polymerization degrees of BC-PANI and CC-PANI were analyzed using a MALDI-TOF/TOF MS 5800 System (AB SCIEX, USA) equipped with a 1 kHz Opti-Beam™ on-axis laser, and a MALDI-TOF Voyager DE-STR (Applied Biosystems, USA) system equipped with a 337-nm nitrogen laser in the linear positive mode, respectively. All samples were dissolved in a...
TA50 (50% acetonitrile/50% water with 0.1% TFA) solution and mixed with α-cyano-4-hydroxycinnamic acid (CHCA) or 2, 5-dihydroxybenzoic acid (DHB) as a matrix. The samples (0.5 μL) were placed in a 384 Opti-TOF stainless steel plate (AB SCIEX, USA).

**Electrical conductivity**

Dried BC- and CC-PANI were split to 1×1 cm pieces, and the thickness of the samples was calculated by using a micrometer (SYNTEK, China). The electrical conductivity of BC- and CC-PANI was measured with the four-point probe method (CMT-SR1000N, Chang Min Tech Co., Ltd., Korea). The electrical resistivity was measured 5 times on different sides and the average was calculated to obtain the surface resistance value. The electrical conductivity was calculated according to the following Eq (1):

$$\sigma = \frac{1}{(R \times d)}$$

where σ stands for the electrical conductivity (S/cm), R stands for the electrical resistivity (Ω·cm), and d denotes the thickness of the sample (cm).

**Flexibility and crease recovery test**

The flexibility of samples was tested according to KS K 0538 (the heart-loop method) with some modifications. The size of samples was modified to 5 x 10 cm and the experiment time was changed to 1 minute. Samples were made to a loop and after 1 minute, the change in physical properties of samples were compared and evaluated. The crease recovery angles of samples were measured according to KS K 0550 (the recovery angle method). The tests were performed five times for each sample and the results were averaged. The crease recovery was calculated according to the following Eq (2):

$$\text{Crease recovery} (\%) = \frac{\text{crease recovery angle (°)}}{180°} \times 100$$

**FT-IR spectroscopy**

FT-IR spectra of BC- and CC-PANI were obtained using a Nicolet IS 50 FT/IR spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA) and TENSOR27 spectrometer (Bruker, Germany), respectively. Scans were acquired between 4,000 cm\(^{-1}\) and 650 cm\(^{-1}\). The peak resolution was 0.4 cm\(^{-1}\) and the number of scans were 32. Baseline correction for each sample spectrum was performed using the Origin 2019 software.

**XRD**

The crystalline structure of BC- and CC-PANI were studied with a D8 Advance diffractometer (Bruker AXS Inc., Karlsruhe, Germany) and a X-ray diffractometer (Rigaku, Japan), respectively, using nickel-filtered Cu Kα radiation (λ = 1.54 nm) operating at 40 kV and a filament emission of 40 mA. The scanning rate and step size was 0.01° min\(^{-1}\) and 0.1° 2θ for BC-PANI, and 1° min\(^{-1}\) and 0.01° 2θ for CC-PANI, respectively, and the diffraction patterns were recorded over a range of 2θ angles from 5 to 40°. The baseline for CC-PANI was subtracted using the Origin software with a smoothing factor of 7 and threshold of 0.01.
All samples were platinum-coated in an ion sputter prior to the observation. BC-PANI and CC-PANI was imaged using a JSM-7600F microscope (JEOL Ltd., Tokyo, Japan) at 2.0 kV. The PANI sample without BC or MC was imaged at 3.0 kV.

Results

Polymerization conditions

The polymerization conditions of BC-PANI and CC-PANI were optimized using UV-Vis spectroscopy. As the PANI polymer solution displays three deterrent colors depending on the oxidation state of PANI, especially green color for PANI emeraldine salt with high electrical conductivity [29], we examined the color change of polymerization solutions of BC-PANI and CC-PANI. UV spectra was observed to determine the optimal point at which PANI was in the emeraldine state. Among three CC-PANI samples, MC-PANI was observed as a representative.

First, the polymerization time was evaluated. As shown in Fig 2, until 0 to 1 h of polymerization, no peak indicative of PANI formation was observed in the UV spectra of both BC-PANI and MC-PANI samples. After 2 h of polymerization for MC-PANI and after 3 h for BC-PANI, the color of the polymer solution changed from white to yellowish-brown, and finally to dark green. Also, the characteristic peaks of PANI were observed in each UV spectrum. In particular, the peaks of PANI corresponded to the π-π* electron transition at 350~360 nm, the polaron-π transition at 420~430 nm, and the π-polaron transition at 760 nm [30]. This result confirmed that the PANI that had polymerized in BC and CC was in the emeraldine state with good electrical conductivity [29]. Thus, the optimized polymerization time was confirmed as 2 h for CC-PANI and 3 h for BC-PANI. The polymerization time for CC-PANI was shorter than for BC-PANI due to the difference in the polymerization method (Fig 1(C)). Bulk polymerization was used for BC-PANI and emulsion polymerization was used for CC-PANI. As emulsion polymerization is conducted in a controlled micro-scale environment, it has the advantage of being a robust polymerization condition, which reduces the polymerization time. Although we attempted to make an emulsion polymerized BC-PANI, PANI was hardly penetrated to the interior of the BC. As the SEM image showed that BC formed dense fabrics with nanofibers of 100 nm or less in diameter, we thought that the emulsion polymerized PANI particles (10–30 nm in diameter) could not be inserted into the BC. Thus, we performed a bulk polymerization for BC-PANI, which allows aniline monomers to penetrate into tightly woven BC fabrics.

Other polymerization conditions, such as the molar ratio of DBSA or APS to aniline, the pH level of the DBSA solution, and the polymerization temperature, were evaluated with the same method. The results are summarized in Table 1.

Degree of polymerization

The degree of polymerization (DP) of BC-PANI and CC-PANI was analyzed by MALDI-TOF MS. As shown in Fig 3, consistent peaks of PANI were observed in the mass spectra of BC-PANI and CC-PANI, confirming that PANI was successfully polymerized with BC and CC. In addition, mass intervals of 90 or 91 m/z appeared in the mass spectra of BC-PANI and CC-PANI, corresponding to the molecular weight of PANI monomer [31]. DP was analyzed according to this interval (Table 2). BC-PANI had short PANI chains with a DP_{max} <5, whereas CC-PANI had longer PANI chains with a DP_{max} of 16 to 17. This result was due to the difference in the polymerization method described in Fig 1(C). The reason why the degree...
of polymerization rises in an emulsion polymerization is that the emulsion polymerization is based on a radical polymerization, which is occurred with emulsified monomer. Namely, as the concentration of the emulsified monomer in a micelle is high, the interval between the emulsified monomers for mass transfer is shorter than the bulk polymerization [32]. In this study, we used DBSA as a surfactant that traps hydrophobic monomer molecule of aniline into a micelle. But, the monomer molecules of aniline are being dispersed in case of the bulk polymerization. Therefore, the concentration of aniline from emulsion polymerization becomes relatively higher than the one from water-based bulk reaction in the same batch volumes. When the aniline molecules are polymerized, radical molecules around the aniline molecules, such as initiators and activated oligomers, are also being efficiently polymerized, resulting long and stable PANI chains. However, as the aniline monomers are being distributed to the whole container and the distance between aniline monomers in a bulk polymerization are long, the molecules are difficult to reach each other. Thus, the evaluation confirmed that CC-PANI had a more stable polymerization pattern compared to BC-PANI.

Fig 2. The color change and the UV spectra of (A) BC-PANI solution with varied polymerization time from 0 to 3 h, and (B) CC-PANI solution with varied polymerization time from 0 to 2 h.

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Table 1. Polymerization conditions of BC-PANI and CC-PANI.

| Polymerization conditions | BC-PANI | CC-PANI |
|--------------------------|---------|---------|
| Time                     | 3 hours | 2 hours |
| Molar ratio (DBSA: Aniline) | 1.0     | 1.0     |
| Molar ratio (APS: Aniline)     | 1.0     | 1.0     |
| pH level of DBSA solution   | pH 2.0  | pH 2.0  |
| Temperature               | 20˚C    | 4˚C     |

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Electrical conductivity

The thickness and weight of BC-PANI and CC-PANI produced under optimized polymerization conditions were measured, and their electrical conductivity values were $1.990 \times 10^{-2}$ S/cm, $2.840 \times 10^{-2}$ S/cm, $2.080 \times 10^{-2}$ S/cm, and $0.962 \times 10^{-2}$ S/cm, respectively (Table 3). The electrical conductivity measurement indicated that MC-PANI and HPMC-PANI had higher electrical conductivity than BC-PANI, confirming that the electrical conductivity improved as the DP increased due to the formation of long and stable PANI chains.

However, the DP value of CMC-PANI was higher than that of BC-PANI whereas the electrical conductivity of CMC-PANI was lower than that of BC-PANI. This can be explained by the differences in the viscosity of cellulose. For conductive polymer composites, it is important how well the polymer is dispersed in the cellulose [33]. If the conductive materials are concentrated in one place, the other place is empty, resulting in the electron’s passage being cut off. Therefore, it is important that conductive polymers spread without aggregation to form a uniform network [34]. A cellulose material with high viscosity prevents the movement of conductive nanoparticles inside the cellulosic structure and helps dispersion [35], thus making nanoparticles to form a uniform electrical network [34]. CMC-PANI showed more than 20 times less viscosity than MC-PANI or HPMC-PANI. This could explain the low electrical conductivity of CMC-PANI.

Flexibility and crease recovery

Fig 4 indicates the results of the flexibility of BC-PANI and CC-PANI based on the modified heart-loop method. A sample has good flexibility and dimensional stability when it has circular shaped loop and is easily returned to its original shape [36]. As shown in Fig 4, a difference in the flexibility was observed in each sample. First, in BC and BC-PANI samples, angular shaped loops were formed rather than circular shaped ones that were formed in chemical cellulosics such as MC, HPMC, and CMC. Moreover, after 1 minute of making loops, BC and BC-PANI maintained the folded shape and did not return to their original state. Based on these results, it

Table 2. The degree of polymerization, molecular weight and polydispersity index from mass spectra of PANI, BC-PANI and CC-PANI (polymerization conditions for BC-PANI: pH level of DBSA solution = 2.0, DBS:aniline:APS = 1:1:1, 20°C, 3 h, polymerization conditions for CC-PANI: pH level of DBSA solution = 2.0, DBS:aniline:APS = 1:1:1, 4°C, 2 h).

|             | DP$_{\text{max}}^*$ | Mw$^{**}$ | PDI$^{***}$ |
|-------------|---------------------|-----------|-------------|
| PANI (BC)   | 5                   | 725.81    | 1.18        |
| PANI (CC)   | 12                  | 530.07    | 1.28        |
| BC-PANI     | 5                   | 688.30    | 1.17        |
| MC-PANI     | 16                  | 948.97    | 1.18        |
| HPMC-PANI   | 17                  | 859.08    | 1.27        |
| CMC-PANI    | 17                  | 927.58    | 1.27        |

*: Maximum degree of polymerization  
**: Molecular weight  
***: Polydispersity index.
was confirmed that the flexibility of BC and BC-PANI was relatively lower than chemical cellulosic. MC and MC-PANI samples formed circular loops and returned to their original state without causing a deformation. CMC, HPMC, and HPMC-PANI samples also formed round shaped loops and displayed a tendency to retain their original states. However, fabric deformation was observed in those samples and CMC-PANI was fractured due to its low flexibility.

In order to verify the flexibility test results with quantitative measurements, crease recovery test was conducted (Table 4). The highest crease recovery value was obtained at MC-PANI with 100% of crease resistance. High crease recovery angle stands for good crease recovery of the sample. Thus, the crease recovery test result indicated that MC had relatively good crease recovery after the polymerization of PANI. From these results, it was observed that MC-PANI had the highest flexibility compared to other samples such as BC-PANI, HPMC-PANI, and CMC-PANI.

Chemical structures

The chemical structures of BC-PANI and CC-PANI were evaluated by FT-IR spectroscopy (Fig 5). The IR spectra of BC-PANI and CC-PANI showed peaks indicating the presence of cellulose (Table 5). The peaks ranging from 3340 cm\(^{-1}\) to 3450 cm\(^{-1}\) were due to the stretching vibration of hydroxyl groups [37]. The peaks near 2900 cm\(^{-1}\) were due to the asymmetric stretching vibration of the C—H bonds in cellulose [38–40]. Particularly, in the spectrum of CMC-PANI, a peak that appeared at 1589 cm\(^{-1}\) was ascribed to the carboxyl groups in CMC [41]. These peaks confirmed that BC and CC maintained cellulose structures after PANI was polymerized.

In addition, characteristic peaks of PANI were observed in the IR spectra of BC-PANI and CC-PANI (Table 6). The original O—H stretching vibration peaks were shifted to 3240 cm\(^{-1}\) to 3230 cm\(^{-1}\) because of the N—H stretching of PANI [42]. The absorption peaks at 1612 cm\(^{-1}\) to 1606 cm\(^{-1}\) were ascribed to the C〓N stretching of quinoid ring [42–44]. The peaks at 1436 cm\(^{-1}\) to 1411 cm\(^{-1}\) were assigned to the C〓C stretching of quinoid ring [43, 45–48]. The peaks at 1315 cm\(^{-1}\) to 1307 cm\(^{-1}\) confirmed the protonated state of PANI with C〓N stretching [46–50]. The results confirmed that PANI was successfully polymerized while maintaining the cellulose structures of BC and CC.

Crystalline structures

XRD was used to characterize the crystalline structure of BC-PANI and CC-PANI. Both BC-PANI and CC-PANI showed reflection peaks characteristic of a cellulose structure (Fig 6). The cellulose peaks were observed at 14.7° and 22.8° for BC-PANI [51] and 7.91° and 20.53° for MC-PANI [38, 52], 8.87° and 19.88° for HPMC-PANI, and 8.87° and 20.62° for CMC-PANI [53, 54]. This suggested that cellulose structures of both BC and CC remained after the polymerization of PANI.
Characteristic peaks of PANI were also appeared in the XRD patterns of both BC-PANI and CC-PANI. For BC-PANI, the peak at 23.7° indicated that PANI inside BC was partially crystalline [55], and the peak at 26.7° was attributed to the parallel arrangement of PANI polymer chains [56]. For CC-PANI, the broad peak around at 20.26° was ascribed to the presence of the emeraldine state of PANI with good electrical conductivity [46–48]. There were more
peaks at 23.46˚ and 25.19˚ for HPMC-PANI, 23.5˚ and 25.46˚ for CMC-PANI, respectively, which are assembled peaks in XRD data of PANI [57]. The peak at 28.57˚ appeared in Table 4. Crease recovery of BC-PANI and CC-PANI (polymerization conditions for BC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 20˚C, 3 h, polymerization conditions for CC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 4˚C, 2 h).

| Material   | Crease recovery (%) |
|------------|---------------------|
| BC-PANI    | 25                  |
| MC-PANI    | 100                 |
| HPMC-PANI  | 86                  |
| CMC-PANI   | N/A                 |

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Fig 5. FT-IR spectra of (A) pure BC, and pure MC, (B) BC-PANI, and MC-PANI, (C) pure HPMC, and pure CMC, and (D) HPMC-PANI, and CMC-PANI (polymerization conditions for BC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 20˚C, 3 h, polymerization conditions for CC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 4˚C, 2 h).

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CMC-PANI was attributed to the periodicity perpendicular to the PANI chain direction [58]. These peaks demonstrated that PANI was successfully polymerized inside BC-PANI and CC-PANI.

Data on the degree of crystallinity of BC-PANI and CC-PANI, and the percent change before and after the polymerization of PANI are presented in Table 7. As shown in Table 7, the crystallinity of BC-PANI, MC-PANI, and HPMC-PANI were decreased by 19.1%, 21.8%, and 34.1%, respectively, whereas that of CMC-PANI was increased by 46.7%. This can be explained by the amorphous nature of PANI polymer. PANI has low crystallinity due to the repetitive formation of benzenoid and quinoid rings in PANI chains [59]. Thus, it is confirmed that the degree of crystallinity decreases as the cellulosic structure effectively anchors the PANI molecules. It can also explain why CMC-PANI exhibited the lowest electrical conductivity among the four conductive cellulose fabrics.

### Surface characterization

The surface morphologies of BC-PANI and CC-PANI were examined by SEM. Fig 7(A) shows the fibrous network structure of pure BC. A three-dimensional nanostructure built from cellulose nanofibers was clearly observed. In the SEM image of BC-PANI (Fig 7(B)), the BC fibers appeared thicker than those in the pure BC, due to the coating of PANI [60]. The images confirmed that BC was polymerized with PANI without significant changes in its fibrous structure. Before observing the surface morphology of CC-PANI, the morphology of a PANI solution diluted 500-fold with isopropanol and dried on a plastic film was examined (Fig 7(C)). The polymerized PANI particles were in spherical structures that were approximately 10 to 30 nm in diameter. This was due to the emulsion polymerization, since the size and shape of PANI particles corresponded to those of micelles [61]. Unlike pure BC, pure CC (Fig 7(E), 7(G) and 7(I)) did not show a fibrillar structure, but rather had a sheet type plane structure [62, 63]. Notably, in the SEM images of CC-PANI (Fig 7(D)), coagulated spherical structures were observed on the surface of CC, indicating the presence of PANI. The findings might reflect the influence of the spreading coefficient of the solvent used to prepare the PANI solution.

### Table 5. Characteristic peaks of cellulose in the FT-IR spectra of pure BC, pure MC, pure HPMC, and pure CMC.

| Wavenumber (cm⁻¹) | Peak no. | BC | MC | HPMC | CMC | Peak assignment |
|------------------|---------|----|----|------|-----|----------------|
| 1                |         | 3342| 3450| 3480 | 3332| O—H stretching [37] |
| 2                |         | 2895| 2894| 2904 | 2914| C—H stretching [38–40] |
|                  |         | 2829| 2837|      | 1589| COO⁻ asymmetric stretching [41] |

### Table 6. Characteristic peaks of PANI in the FT-IR spectra of BC-PANI and CC-PANI (polymerization conditions for BC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 20°C, 3 h, polymerization conditions for CC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 4°C, 2 h).

| Wavenumber (cm⁻¹) | Peak no. | BC-PANI | MC-PANI | HPMC-PANI | CMC-PANI | Peak assignment |
|------------------|---------|---------|---------|-----------|----------|----------------|
| 1                |         | 3240    | 3282    | 3230      | 3232     | N—H stretching [42] |
| 2                |         | 1612    | 1606    | 1606      |          | C=\N stretching of quinoid ring [42, 44] |
|                  |         | 1568    | 1567    | 1571      |          | C=\C stretching of benzenoid ring [45–48] |
| 4                |         | 1425    | 1436    | 1423      | 1411     | C=\C stretching of quinoid ring [43, 45–48] |
| 5                |         | 1313    | 1307    | 1309      | 1313     | C—N stretching of secondary amine [46–50] |

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Discussion

In this study, we produced BC-PANI and CC-PANI composites with good electrical conductivity by using PANI in the emeraldine-salt state. Interestingly, we directly fabricated BC-PANI and CC-PANI-based conductive fabric material.

Table 7. Crystallinity degrees of BC-PANI and CC-PANI (polymerization conditions for BC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 20°C, 3 h, polymerization conditions for CC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 4°C, 2 h).

|                | Crystallinity (%) | Percent change (%) |
|----------------|-------------------|--------------------|
|                | Before polymerization | After polymerization |
| BC-PANI        | 77.4               | 65.0               | -19.1 |
| MC-PANI        | 60.9               | 50.0               | -21.8 |
| HPMC-PANI      | 52.7               | 39.3               | -34.1 |
| CMC-PANI       | 32.8               | 61.5               | 46.7  |

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Fig 6. XRD patterns of (A) pure BC and BC-PANI, (B) pure MC and MC-PANI, (C) pure HPMC and HPMC-PANI, and (D) pure CMC and CMC-PANI (polymerization conditions for BC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 20°C, 3 h, polymerization conditions for CC-PANI: pH level of DBSA solution = 2.0, DBSA:aniline:APS = 1:1:1, 4°C, 2 h).

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and CC-PANI by mixing BC and CC powders with a PANI solution. Importantly, it was confirmed that polymerization of PANI on MC enables the fabrication of conductive cellulose with improved electrical conductivity than BC. The findings support the view that the MC-PANI is a good conductive fabric that could potentially be used in wearable devices and sensors.

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References
1. Barakzeh i M, Montazer M, Sharif F, Norby T, Chatzitakis A. A textile-based wearable supercapacitor using reduced graphene oxide/polypryrole composite. Electrochimica Acta. 2019; 305:187–196. https://doi.org/10.1016/j.electacta.2019.03.058
2. Yun Y, Hong W, Kim W, Jun Y, Kim B. A Novel Method for Applying Reduced Graphene Oxide Directly to Electronic Textiles from Yarns to Fabrics. Advanced Materials. 2013; 25(40):5701–5705. https://doi.org/10.1002/adma.201303225 PMID: 23946273
3. Lee Y, Kim J, Noh J, Lee I, Kim H, Choi S et al. Wearable Textile Battery Rechargeable by Solar Energy. Nano Letters. 2013; 13(11):5753–5761. https://doi.org/10.1021/nl403860k PMID: 24164580
4. Majumder S, Mondal T, Deen M. Wearable Sensors for Remote Health Monitoring. Sensors. 2017; 17 (12):130. https://doi.org/10.3390/s17010130 PMID: 28085085
5. Huang Q, Wang D, Zheng Z. Textile-Based Electrochemical Energy Storage Devices. Advanced Energy Materials. 2016; 6(22):1600783. https://doi.org/10.1002/aenm.201600783
6. Zysset C., Cherenack K., Kinkeldei T. and Tröster G., “Weaving integrated circuits into textiles,” International Symposium on Wearable Computers (ISWC) 2010, Seoul, 2010: 1–8. https://doi.org/10.1109/ISWC.2010.5656874
7. Mičušik M, Nedelčev T, Omastová M, Krupa I, Olejníková K, Fedorko P et al. Conductive polymer-coated textiles: The role of fabric treatment by pyrrole-functionalized triethoxysilane. Synthetic Metals. 2007; 157(22–23):914–923. https://doi.org/10.1016/j.synthmet.2007.09.001
8. Kaur G, Adhikari R, Cass P, Bown M, Gunatillake P. Electrically conductive polymers and composites for biomedical applications. RSC Advances. 2015; 5(47):37553–37567. https://doi.org/10.1039/c5ra01851j
9. Rahman M, Kumar P, Park D, Shim Y. Electrochemical Sensors Based on Organic Conjugated Polymers. Sensors. 2008; 8(1):118–141. https://doi.org/10.3390/s8010118 PMID: 27879698
10. Palaniappan S, John A. Polyaniylene materials by emulsion polymerization pathway. Progress in Polymer Science. 2008; 33(7):732–758. https://doi.org/10.1016/j.progpolymsci.2008.02.002
11. Kim B, Koncar V, Dufour C. Polyaniylene-coated PET conductive yarns; Study of electrical, mechanical, and electro-mechanical properties. Journal of Applied Polymer Science. 2006; 101(3):1252–1256. https://doi.org/10.1002/app.22799
12. Tang S, Wang A, Lin S, Huang K, Yang C, Yeh J et al. Polymerization of aniline under various concentrations of APS and HCl. Polymer Journal. 2011; 43(8):667–675. https://doi.org/10.1038/pj.2011.43
13. Ćirić-Majlanović G. Recent advances in polyaniline research: Polymerization mechanisms, structural aspects, properties and applications. Synthetic Metals. 2013; 177:1–47. https://doi.org/10.1016/j.synthmet.2013.06.004
14. Bhadra J, Al-Thani N, Madi N, Al-Maadeed M. Effects of aniline concentrations on the electrical and mechanical properties of polyaniline polyvinyl alcohol blends. Arabian Journal of Chemistry. 2017; 10 (5):664–672. https://doi.org/10.1016/j.arabjc.2015.04.017
15. Cochrane C, Koncar V, Lewandowski M, Dufour C. Design and Development of a Flexible Strain Sensor for Textile Structures Based on a Conductive Polymer Composite. Sensors. 2007; 7(4):473–492. https://doi.org/10.3390/s7040473

16. George J, Sabapathi SN. Cellulose nanocrystals: synthesis, functional properties, and applications. Nanotechnology, Science and Applications. 2015; 8:45–54. https://doi.org/10.1039/c6nt09494e PMID: 28116390

17. Petersson L, Oksman K. Biopolymer based nanocomposites: Comparing layered silicates and micro-crystalline cellulose as nanoreinforcement. Composites Science and Technology. 2006; 66(13):2187–2196. https://doi.org/10.1016/j.compscitech.2005.12.010

18. Trache D, Hussein M, Haafiz M, Thakur V. Recent progress in cellulose nanocrystals: sources and production. Nanoscale. 2017; 9(5):1763–1786. https://doi.org/10.1039/c6nr09494e PMID: 28116390

19. Ashjaran A, Yazdanshenas M, Rashidi A, Khajavi R, Rezaee A. Overview of bio nanofabric from bacterial cellulose. Journal of the Textile Institute. 2013; 104(2):121–131. https://doi.org/10.1080/00405000.2012.703796

20. Properties Ashjaran A. and Applications of Bacterial Cellulose as a Biological Non-woven Fabric. Asian Journal of Chemistry. 2013; 25(2):783–788. https://doi.org/10.14233/ajchem.2013.12889

21. Cochis A, Grad S, Stoddart MJ, Faré S, Altmare L, Azzimonti B, et al. Bioreactor mechanically guided 3D mesenchymal stem cell chondrogenesis using a biocompatible novel thermo-reversible methylcellulose-based hydrogel. Scientific Reports. 2017; 7(1):45018. https://doi.org/10.1038/srep45018 PMID: 28332587

22. Park J-S, Ruckenstein EJCP. Viscoelastic properties of plasticized methylcellulose and chemically crosslinked methylcellulose. 2001; 46(4):373–81.

23. Chang C, Zhang L. Cellulose-based hydrogels: Present status and application prospects. Carbohydrate Polymers. 2011; 84(1):40–53. https://doi.org/10.1016/j.carbpol.2010.12.023

24. Chattopadhyay D, Mandal B. Methyl Cellulose Stabilized Polyaniiline Dispersions. Langmuir. 1996; 12(6):1585–1588. https://doi.org/10.1021/la950523y

25. Jia B, Hino T, Kuramoto N. Synthesis and chiroptical properties of water-processable polyaniiline using methylcellulose as a molecular template. Reactive and Functional Polymers. 2007; 67(9):836–843. https://doi.org/10.1016/j.reactfunctpolym.2007.01.005

26. Han J, Shim E, Kim H. Effects of cultivation, washing, and bleaching conditions on bacterial cellulose fabric production. Textile Research Journal. 2018; 89(6):1094–1104. https://doi.org/10.1177/0040517518763989

27. Song J, Su J, Loureiro A, Martins M, Cavaco-Paulo A, Kim HR et al. Ultrasound-assisted swelling of bacterial cellulose. Engineering in Life Science. 2017; 17(10):1108–1117. https://doi.org/10.1002/elsc.201700085

28. Kim H, Song J, Silva C, Kim HR. Production of conductive bacterial cellulose-polyaniiline membranes in the presence of metal salts. Textile Research Journal. Epub 2019/12/16. https://doi.org/10.1177/0040517519893717

29. Yoon S-B, Yoon E-H, Kim K-B. Electrochemical properties of leucoemer aldine, emeraldine, and pernigraniline forms of polyaniiline/multi-wall carbon nanotube nanocomposites for supercapacitor applications. Journal of Power Sources. 2011; 196(24):10791–7. https://doi.org/10.1016/j.jpowsour.2011.08.107

30. Sambasevam KP, Mohamad S, Phang S-W. Enhancement of polyaniiline properties by different polymerization temperatures in hydrazine detection. Journal of Applied Polymer Science. 2015; 132(13):n/a-n/a. https://doi.org/10.1002/app.41746

31. Ji Y, Qin C, Niu H, Sun L, Jin Z, Bai X. Electrochemical and electrochromic behaviors of polyaniiline-graphene oxide composites on the glass substrate/Ag nano-film electrodes prepared by vertical target pulsed laser deposition. Dyes and Pigments. 2015; 117:72–82. https://doi.org/10.1016/j.dyepig.2015.01.026

32. Rao PS, Subrahmanyas, Sathyaranayana DN. Inverse emulsion polymerization: a new route for the synthesis of conducting polyaniiline. Synthetic Metals. 2002; 128(3):311–6. https://doi.org/10.1016/S0379-6779(02)00014-4

33. Cho MS, Park SY, Hwang JY, Choi HJ. Synthesis and electrical properties of polymer composites with polyaniiline nanoparticles. Materials Science and Engineering: C. 2004; 24(1–2):15–8. https://doi.org/10.1016/j.msec.2003.09.003

34. Liu L, Grunlan JC. Clay Assisted Dispersion of Carbon Nanotubes in Conductive Epoxy Nanocomposites. Advanced Functional Materials. 2007; 17(14):2343–8. https://doi.org/10.1002/adfm.200600785

35. Arboleya J, Wilde P. Competitive adsorption of proteins with methylcellulose and hydroxypropyl methylcellulose. Food Hydrocolloids. 2005; 19(3):485–91. https://doi.org/10.1016/j.foodhyd.2004.10.013
36. Zhou N and Ghosh TK. On-Line Measurement of Fabric Bending Behavior, Textile Research Journal 68(7), 533–542. https://doi.org/10.1177/004051759806800711

37. Maiti D, Mollick MM, Mondal D, Bhowmick B, Bain MK, Bankura K et al. Synthesis of methylcellulose-silver nanocomposite and investigation of mechanical and antimicrobial properties. Carbohydrate Polymers. 2012; 90(4):1818–25. Epub 2012/09/05. https://doi.org/10.1016/j.carbpol.2012.07.082 PMID: 22944452.

38. Trivedi MK, Branton A, Trivedi D, Nayak G, Mishra RK, Jana S. Characterization of physicochemical and thermal properties of biofield treated ethyl cellulose and methyl cellulose. International Journal of Biomedical Materials Research. 2015; 3(6):83–91. https://doi.org/10.11648/j.ijbmr.20150306.12

39. Singh RK. Methylcellulose synthesis from corn cobs. Journal of Thermal Analysis and Calorimetry. 2013; 114(2):809–19. https://doi.org/10.1007/s10973-013-3032-4

40. Mruthyunjaya Swamy TM, Ramaraj B, Siddaramaiah. Thermal and morphological properties of SA/HPMC blends. Journal of Applied Polymer Science. 2009; 112(4):2235–40. https://doi.org/10.1002/app.29738

41. Hebeish A, Hashem M, EI-Hady MM, Sharaf S. Development of CMC hydrogels loaded with silver nanoparticles for medical applications. Carbohydrate Polymers. 2013; 92(1):407–13. Epub 2012/12/12. https://doi.org/10.1016/j.carbpol.2012.08.094 PMID: 23218313.

42. Jafari Y, Ghoreishi SM, Shabani-Nooshabadi M. Characterization and Corrosion Inhibition Study of DBSA-Doped Polyaniline Coating on 310 Stainless Steel. Iranian Journal of Chemistry & Chemical Engineering, 2017; 36(5):23–32.

43. Butoi B, Groza A, Dinca P, Balan A, Barna V. Morphological and Structural Analysis of Polyaniline and Poly(o-anisidine) Layers Generated in a DC Glow Discharge Plasma by Using an Oblique Angle Electrode Deposition Configuration. Polymers. 2017; 9(12):732. https://doi.org/10.3390/polym9120732 PMID: 30966033

44. Jeevananda T, Siddaramaiah, Kim N, Heo S, Lee J. Synthesis and characterization of polyaniline-multi-wall carbon nanotube nanocomposites in the presence of sodium dodecyl sulfate. Polymers for Advanced Technologies. 2008; 19(12):1754–1762. https://doi.org/10.1002/pat.1191

45. Kızıldag N, Ucar N, Oztoksoy ME, Garimestani H, Wang Y, Dahmen K-H. Poly(vinyl alcohol)/Polyaniline (PVA/PANI) conductive nanofibers by electrospinning. In: The International Istanbul Textile Congress 2013, Istanbul, Turkey, 30 May-1 June 2013, pp. 1–6. Istanbul: The International Istanbul Textile Congress.

46. Radhakrishnan S, Rao CRK, Vijayan M. Performance of conducting polyaniline-DBSA and polyaniline-DBSA/Fe3O4 composites as electrode materials for aqueous redox supercapacitors. Journal of Applied Polymer Science. 2011; 122(3):1510–8. https://doi.org/10.1002/app.34236

47. Misoon O, Seok K. Effect of dodecyl benzene sulfonic acid on the preparation of polyaniline/activated carbon composites by in situ emulsion polymerization. Electrochimica Acta. 2012; 59:196–201. https://doi.org/10.1016/j.electacta.2011.10.058

48. Chen T, Dong C, Li X, Gao J. Thermal degradation mechanism of dodecylbenzene sulfonic acid-hydrochloric acid co-doped polyaniline. Polymer Degradation and Stability. 2009; 94(10):1788–94. https://doi.org/10.1016/j.polymdegradstab.2009.06.011

49. Kakde K. Synthesis and Characterization of Polyaniline Doped with HCl, H2SO4 and PVA as Secondary Dopant for Toxic Gas (Ammonia) Sensor. Indian Journal of Science and Technology. 2017; 10(20):1–4. https://doi.org/10.17485/ijst/2017/v10i20/112920

50. Karaoğlan N, Bindal C. Synthesis and optical characterization of benzene sulfonic acid doped polyaniline. Engineering Science and Technology, an International Journal. 2018; 21(6):1152–1158. https://doi.org/10.1016/j.ijestch.2018.09.010

51. Fei G, Wang Y, Wang H, Ma Y, Guo Q, Huang W et al. Fabrication of Bacterial Cellulose/Polyaniline Nanocomposite Paper with Excellent Conductivity, Strength, and Flexibility. ACS Sustainable Chemistry & Engineering. 2019; 7(9):8215–8225. https://doi.org/10.1021/acssuschemeng.8b06306

52. Maiti D, Mollick MM, Mondal D, Bhowmick B, Bain MK, Bankura K et al. Synthesis of methylcellulose-silver nanocomposite and investigation of mechanical and antimicrobial properties. Carbohydrate Polymers. 2012; 90(4):1818–25. Epub 2012/09/05. https://doi.org/10.1016/j.carbpol.2012.07.082 PMID: 22944452.

53. George J, Kumar R, Sajeevkumar VA, Ramana KV, Rajamanickam R, Abhishek V et al. Hybrid HPMC nanocomposites containing bacterial cellulose nanocrystals and silver nanoparticles. Carbohydrate Polymers. 2014; 105:285–92. Epub 2014/04/09. https://doi.org/10.1016/j.carbpol.2014.01.057 PMID: 24709882.

54. Salama A. Preparation of CMC-g-P(SPMA) super adsorbent hydrogels: Exploring their capacity for MB removal from waste water. International Journal of Biological Macromolecules. 2018; 106:940–6. Epub 2017/08/24. https://doi.org/10.1016/j.ijbiomac.2017.08.097 PMID: 28834704.
55. Guo G. Facile Synthesis of Conducting Hydrogels Based on Polyaniline Fiber and Graphene Oxide for Application in all-solid-state Supercapacitors. International Journal of Electrochemical Science. 2019; 5899–5912. https://doi.org/10.20964/2019.06.220

56. Mu B, Tang J, Zhang L, Wang A. Facile fabrication of superparamagnetic graphene/polyaniline/Fe₃O₄ nanocomposites for fast magnetic separation and efficient removal of dye. Scientific Reports. 2017; 7(1). https://doi.org/10.1038/s41598-017-05755-6 PMID: 28706284

57. Han D, Chu Y, Yang L, Liu Y, Lv Z. Reversed micelle polymerization: a new route for the synthesis of DBSA–polyaniline nanoparticles. Colloids and Surfaces A: Physicochemical and Engineering Aspects. 2005; 259(1–3):179–87. https://doi.org/10.1016/j.colsurfa.2005.02.017

58. Mokhtar N, Chye D, Phang S. Morphology, Conductivity and Microwave Absorption Behavior of Polyaniline Nanocomposites after Chemical Treatment. Polymers and Polymer Composites. 2017; 25(7):545–554. https://doi.org/10.1016/j.colsurfa.2005.02.017

59. Mostafaei A, Zolriasatein A. Synthesis and characterization of conducting polyaniline nanocomposites containing ZnO nanorods. Progress in Natural Science: Materials International. 2012; 22(4):273–280. https://doi.org/10.1016/j.pnsc.2012.07.002

60. Yue L, Xie Y, Zheng Y, He W, Guo S, Sun Y et al. Sulfonated bacterial cellulose/polyaniline composite membrane for use as gel polymer electrolyte. Composites Science and Technology. 2017; 145:122–131. https://doi.org/10.1016/j.compsci.2017.04.002

61. Han Y-G, Kusunose T, Sekino T. One-step reverse micelle polymerization of organic dispersible polyaniline nanoparticles. Synthetic Metals. 2009; 159(1–2):123–31. https://doi.org/10.1016/j.synthmet.2008.08.011

62. Khan RA, Salmieri S, Dussault D, Sharmin N, Lacroix M. Mechanical, barrier, and interfacial properties of biodegradable composite films made of methylcellulose and poly(caprolactone). Journal of Applied Polymer Science. 2012; 123(3):1690–7. https://doi.org/10.1002/app.34655

63. Khan RA, Dussault D, Salmieri S, Safrany A, Lacroix M. Improvement of the mechanical and barrier properties of methylcellulose-based films by treatment with HEMA and silane monomers under gamma radiation. Radiation Physics and Chemistry. 2012; 81(8):927–31. https://doi.org/10.1016/j.radphyschem.2012.02.044