Highly transparent, conducting, body-attachable metallized fibers as a flexible and stretchable film

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Abstract

silver-electroplated nickel microfibers Core-shell-structured were fabricated via electrospinning and subsequent electroplating for applications including transparent conductive films (TCFs) and heaters. The fabrication protocol generated self-fused junctions at the intersections of overlapping micro-nanofibers, yielding remarkably high electrical conductivity, which is highly desirable for the aforementioned applications. A very low sheet resistance of less than 0.2 Ω sq⁻¹ with a high transmittance of over 92% was achieved in these structures. A cactus-like morphology of silver-plated microfibers, which dramatically increases surface-to-volume (S/V) ratio which should produce electric field concentration at silver nanowire tips, was also demonstrated. This unique surface morphology could be promising for energy and environmental applications that require large interfacial areas and electric field concentration, but yielded lower transmittance. These cactus-like microfibers could be further coated with Cu and Pt to produce hierarchically-structured multimetallic microfibers. The lowresistivity transparent silver micro-nanofiber films exhibited good heating and mechanical properties, as demonstrated in bending and stretching tests. A record high temperature of 209 °C was achieved with a transparent heater based on the Ag microfibers.

Keywords: Silver-plated fibers, Core-shell fibers, Flexible electronics, Transparent conducting films and heaters

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1. Introduction

One-dimensional (1D) Ag microstructures, such as Ag nanowires (AgNWs), are of great interest because of their potential applicability in various electronic devices [1-8]. Nanoscale 1D Ag, which possesses the best electrical and thermal properties among known metals, is among the most valuable materials for transparent conductive electrodes (TCEs) [9, 10]. Micro- and nanoscale electrode performance and reliability depend on the percolative structures of TCE materials [11, 12]. An ideal percolative cluster with a continuous conducting network structure can confer high electrical conductivity to electrodes without significant losses in transparency [9, 13].

Several methods have been employed to fabricate 1D Ag microstructures, including hydrothermal [14, 15], microwave-assisted [16, 17], electrochemical [18], UV- [19, 20], and polyol-based ones [21, 22]. The polyol method is currently recognized as the most promising approach in terms of cost and scalability. This process is based on the growth of Ag nanoparticles (NPs) via a reaction between metallic salts and polyol. However, the approach is insufficiently mature for commercial-level implementation.

We recently explored the fabrication of percolative TCEs via electrospinning and electroplating [11, 12]. Electrospinning produces polymer nanofibers (NFs) that can be applied in various industrial fields, including biomedical- and energy-related industries [23-28]. Electrospun NFs can be electroplated to form metallized NFs when their surfaces are seeded with metal NPs by either dip-coating or sputtering. Subsequent oxidation of the metal fibers converts them into metal oxides that can be used in various energy and environmental applications requiring semi-conducting properties and high interfacial areas [11, 12, 28-34]. A percolation model that describes the electrical and optical behaviors of 1D nanomaterials was

recently reported based on the renormalization group theory technique of the classical percolation theory [11, 35, 36].

Note also that transparent conducting networks of NFs (about 100 nm in diameter) can be formed via the electrospinning and solution blowing of polymer blends containing conducting polymers [37]. Such a NF network reached a moderate sheet resistance (R_s) value of 8.4 k Ω sq⁻¹ with 84% optical transmittance.

Sinha-Ray *et al.* [29] presented initial studies on the production of Ag-plated NFs by electrospinning and electroplating. These fibers were used as heat-removing mats for microelectronic cooling applications. However, the electrical properties of these NFs were not reported and transparent conductive films (TCFs) and heating applications were not discussed. We demonstrate herein the enhanced electrical properties of electrospun and Ag-plated NFs, surpassing those of other 1D nanomaterials in terms of transparency and electrical conductivity [11]. In addition, a cactus-like nanotextured surface on Ag microfibers (MFs) is achieved by the manipulation of the electroplating precursor formulation. Because of the dramatically increased surface area of the fibers, the nanotextured AgMFs show great promise for applications requiring large interfacial area of Ag or electric field concentration at sharp Ag tips.

2. Experimental

2.1. Materials

2.1.1. Electrospinning. Polyacrylonitrile (PAN, $M_w = 150$ kDa) and *N*,*N*-dimethylformamide (DMF, 99.8%) were purchased from Sigma-Aldrich (USA). Ni foil (99.9%, thickness of 20 μ m), used as a material for the Ni frame (as a collector of electrospun fibers) and as the anode electrode (in the Ni-electroplating process), was purchased from Wellcos Corporation (Republic of Korea).

2.1.2. Ni-electroplating. Ni (II) sulfamate tetrahydrate [Ni(SO₃NH₂)₂·4H₂O, 98%], boric acid (H₃BO₃, 99.97%), sodium hydroxide solution (NaOH, 1.0 M), and formaldehyde (CH₂O, 35%) were purchased from Sigma-Aldrich (USA). Single-distilled deionized (DI) water purchased from Samchun Pure Chemical (Republic of Korea) was used for the preparation of the electroplating solutions.

2.1.3. First Ag-electroplating. Ag nitrate (AgNO₃, 99%), potassium hydroxide (KOH, 85%), ammonium hydroxide (NH4OH, 99.9%), and nitric acid (HNO₃, 70%) were purchased from Sigma-Aldrich (USA). Sugar (Beksul, Republic of Korea) was obtained from a local market. Platinum (Pt) plates, used as the anode material (in the Ag/Cu/Pt electroplating process), were obtained from HanTeck PMC (Republic of Korea).

2.1.4. Second Ag-electroplating. The KS-700 solution, potassium argentocyanide $[KAg(CN)_2]$, and potassium cyanide (KCN, 98%) were obtained from Poongwon Chemical (Republic of Korea). The KS-700 solution consisted of 6% (v/v) sodium borate (B4Na₂O₇), 79% (v/v) water, 10% (v/v) potassium phosphate (KH₂PO₄), and 5% (v/v) phosphates.

2.1.5 Cu-electroplating. Sulfuric acid (H₂SO₄, 99%) was purchased from Matsunoen Chemicals (Japan), and hydrochloric acid (HCl, 37%) and Cu(II) sulfate (CuSO₄, 99%) were purchased from Sigma-Aldrich (USA).

2.1.6. Pt-electroplating. A chloroplatinic acid solution (H₂PtCl₆, 5 wt% in distilled water) was obtained from HanTeck PMC (Republic of Korea).

2.2. Prefabrication of polyacrylonitrile nanofibers

First, an 8 wt% yellow PAN solution was prepared by dissolving PAN powder in DMF. The solution was stirred for 24 h at 25°C until it became homogenous. The PAN solution was then injected with an 18-gauge needle (Nordson EFD, USA) using a syringe pump (Legato 100, KD Scientific Inc., USA) with a fixed flow rate of $Q_{es} = 200 \ \mu L \ h^{-1}$. A direct current (DC) voltage of $V_{es} = 6.5 \ kV$ was applied to the needle using a DC power supply (EP20P2, Glassman High Voltage Inc., USA). Electrospinning of the Taylor cone-like PAN jet at the needle tip was produced a continuous single nanofiber (NF), which was deposited on aa Ni frame with a square open area of 3 cm × 3 cm and a frame width of 1 cm. The deposition time (or electrospinning time), t_{es} , was set to 5 s to obtain a thin NF mat with high transparency. Note that the frame-type substrate was used to obtain freestanding NFs.

2.3. Fabrication of Ni and Ni/Ag-electroplated fibers

The Ni-electroplating solution was prepared by blending 80 g Ni(SO₃NH₂)₂·4H₂O, 6 g H₃BO₃, and 200 mL DI water [29]. The pH of the solution was adjusted to 4.5 by adding NaOH and measured using pH paper (Advantec, Japan). The solution was blended using a magnetic

stirrer for 24 h at room temperature before use. Note that all electroplating solutions were stored in dark brown glass bottles at room temperature and used less than 20 times.

For the first Ag-electroplating solution, two different solutions were prepared: solution A and solution B [29]. For preparing solution A, 0.5 g AgNO₃ and 0.5 g KOH were dissolved in 5 mL of DI water, respectively. Next, these two solutions were mixed, and 1.25 mL of NH₄OH was added. Solution A was stirred for 24 h. For preparing solution B, sugar (2.25 g), 0.1 mL HNO₃, and 25 mL DI water were mixed. Solution B was boiled at 100 °C and stirred by a magnetic bar for 24 h. Finally, solution A and solution B were blended and stirred for 10 min. After the plating solution was prepared, the Ag-electroplating experiment proceeded immediately.

The second Ag-electroplating solution was prepared as follows: 250 mL KS-700 solution was mixed with 200 mL DI water and maintained at 45 °C. Next, 60 g KAg(CN)₂ was added to the solution and the total volume was adjusted to 500 mL while the temperature was maintained at 45 °C. Then, this solution was heated to 65 °C and 100 g of KCN was added to it. The pH of the solution was also adjusted to 4.5 by adding NaOH. This solution was also blended and stored in a dark brown glass bottle at room temperature, like the Ni-electroplating solution, and used less than 20 times for electroplating.

Prior to the electroplating process, the freestanding PAN NFs were lightly sputtered with Pt (MSP-1S, Vacuum Device Inc., Japan) [29, 38]. According to our previous work [11], a Pt layer of a few nanometers in thickness layer sputtered on the PAN NFs was sufficient to provide the electrical conductivity required to trigger the electroplating of the polymer NFs. The PAN NFs were first electroplated with Ni, followed by Ag. The Ni-electroplated fibers (NiEFs) were prepared as follows: The PAN NFs (suspended on a Ni frame) and a piece of Ni foil (3 cm \times 3 cm) were immersed in the electroplating solution as the cathode and anode,

respectively. An electric current (current density $I_{ep} = 0.20$ A cm⁻² based on the total mat area of the PAN NFs) was sustained by a power supply (SPS-1820, GW Instek, Taiwan) for 60 s. The distance between the frame and the foil was 2 cm. The plated sample was rinsed with 40% (v/v) CH₂O (diluted in DI water) and DI water for 10 min and 5 s, respectively. Finally, the rinsed sample was dried under an Ar atmosphere for a few minutes to prevent oxidation. As a result, the electroplating of Ni yielded Ni-wrapped PAN NFs. The NiEFs were then electroplated with Ag.

Most of the experimental conditions for the Ag-electroplating were the same as those used for the Ni-electroplating. The Ag-electroplating solution and the Pt plate for the anode were used instead of the Ni-electroplating solution and Ni foil, respectively. The first Agelectroplating was conducted with $I_{ep} = 0.125$ A cm⁻² and different electroplating times of t_{ep} = 10 and 60 s. Electroplating of Ag yielded uniquely shaped Ag-wrapped NiEFs, as shown in Figs. 1. The second Ag-electroplating was conducted on NiEFs with different electroplating voltages of $V_{ep} = 2$, 6, and 8 V, and the electroplating times of $t_{ep} = 1$, 3, 5, and 10 s. In addition, according to the different Ag-electroplating methods, two types of Ag microfibers (MFs) were fabricated as shown in Fig. 2.

2.4. Fabrication of Ni/Ag/Cu/Pt-electroplated fibers

The Cu-electroplating solution was prepared by mixing 10 g H₂SO₄, 1 g HCl, 32 g CuSO₄, and 20 g CH₂O with 200 mL of DI water [29, 39]. The solution was blended using a magnetic stirrer for 24 h at room temperature. The Cu-electroplating was performed on the first AgMFs by applying a voltage of 3 V for 30 s. A Pt plate was used as the anode. The Ni/Ag/Cu-electroplated fibers were then rinsed with 40% (v/v) CH₂O (diluted in DI water) and DI water

for 10 min and 5 s, respectively. The mat was then dried under Ar atmosphere for a few minutes to prevent oxidation.

The Pt-electroplating solution was obtained by adding DI water to 160 mL of 5% H₂PtCl₆ until the total solution volume reached 1 L [12]. The solution was magnetically stirred for one day at room temperature to stabilize. The pH was adjusted to 1.5 by adding H₂SO₄ or NaOH. The Pt-electroplating conditions for the Ni/Ag/Cu/Pt were $I_{ep} = 0.2$ A cm⁻² and $t_{ep} = 90$ min, using a Pt plate as the anode.

2.5. Characterization

The surface morphologies of the samples were analyzed by scanning electron microscopy (SEM) using a field-emission SEM system with energy-dispersive X-ray spectroscopy (FE-SEM/EDX, Quanta 250 FEG, FEI). The average diameter of the AgMFs was measured by using I[^]MEASURE3 software (IngPlus, Republic of Korea); 100 fibers from the SEM images were measured and the average value was used. The roughness values R_a of the AgMFs were obtained by an optical profiler (OP, NT-1100, Veeco, USA). The arithmetic average roughness values were obtained using Vision 3.0 software (Veeco, USA); 10 fibers from the optical images were measured and the average value was used. X-ray diffraction (XRD, SmartLab, Rigaku) and X-ray photoelectron spectroscopy (XPS, X-tool, ULVAC-PHI) were used to analyze the crystalline structures and surface elemental compositions of the AgMFs. The transparency T_r , which is based on air, was measured for a square area of 2 cm × 0.8 cm in air using ultraviolet-visible spectrophotometry (Optizen POP, Mecasys). The R_s values were obtained by the four-point probe method with a sheet resistance meter (FPP-400, Dasol Eng) based on a dual-configuration procedure. The value of R_s was the average value of

10 randomly measured R_s values at different locations. The AgMF sample for cross-sectional imaging was prepared using a dual-beam focused ion beam (FIB) system (SCIOS, FEI) and the sample was analyzed by transmission electron microscopy (TEM, JEM 2100F, JEOL Inc.). In the FIB process, rough cuts were performed with a beam current of 7–15 nA at 30 kV, and final cleaning was conducted with a beam current of 3-7 nA at 30 kV to ensure minimal ion damage. For the stretching test (cf. Movie S1), an AgMF mat was first transferred onto an Ecoflex film with 1 ml of eco-flex for adhesion and was dried for 24 h. One end of the sample was fixed and the other end of the sample was stretched using a syringe pump (LEGATO100, KDScientific, USA) with a fixed speed of cyclic motion. Simultaneously, the resistance changes were recorded by a digital multimeter (3256, Hioki) during stretching. For the bending cycle test, an AgMF mat was transferred onto a polyethylene terephthalate (PET) film; a drop of ethanol was first dripped onto the film and then the AgMF mat was placed onto the PET film using tweezers. In a short time, the ethanol between the fibers and PET film was evaporated, thus facilitating the adhesion of the fibers to the PET film. The AgMF mat adhered to the PET film during the bending test and did not delaminate. The sample was repetitively bent around one axis with a bending radius of 4 mm and frequency of 2 Hz. The resistances between several pairs of points, equally spaced in the direction perpendicular to the axis, were measured. For LED operation, LEDs were purchased from GreenMax (0.24 W at 12 V, Republic of Korea). Six LEDs were connected in series and the voltage of 6 V was applied to the circuit. The heating performance was examined using a power supply (SPS-1820, GW Instek, Taiwan), thermocouple, and data recorder (MV1000, Yokogawa). To minimize the IR loss, 2.3-mm-diameter 10-cm-long Cu wires were used in the electrochemical experiments with the maximum iR loss of only 0.002%. The heating performance under stretching (cf. Movie S2) was observed by an infrared (IR) camera (FLIR-E63900, FLIR, Wilsonvile, USA), where AgMFs on Eco-flex were stretched by a syringe pump at a speed of movement. Note that all the mechanical and optical experiments mentioned above were conducted at ambient conditions of room temperature and 20-30% humidity.

3. Results and discussion

Scanning electron microscopy (SEM) images of the silver microfibers (AgMFs) are presented in Fig. 2 (cf. Low magnification SEM images are also presented in Fig. S1). Discernibly unique morphologies are produced using the first Ag-electroplating solution, as revealed in Figs. 2a and 2b. In contrast to the second Ag-electroplating process (cf. Figs. 2c– 2k), highly textured cactus-like surfaces were observed with the first method. Although the second Ag-electroplating process also produced unique textured surfaces, the first induced a distinctly different morphology. Different trends were observed with respect to T_r and R_s for the two fiber types (see Tables 1 and 2). Thus, the choice of the Ag-electroplating method should be dictated by the intended applications of the AgMFs. For example, the first process appears to be more appropriate for catalytic and electrocatalytic applications, while the second process may be appropriate for transparent conductive electrodes (TCEs) where a balance between high T_r and low R_s is necessary. The mechanical and thermal experiments in this study were based on the second AgMFs for application to TCEs.

Using the second Ag-electroplating method with the electroplating voltage of $V_{ep} = 2$ V (Figs. 2c–2e), granular island-like clusters were initially formed on the outer surfaces of the Ni-electroplated fibers (NiEFs). In addition, as the electroplating time (t_{ep}) increased from 1 to 10 s, the overall surface of the fibers was gradually covered by Ag (Figs. 2c–2e). In contrast, at $V_{ep} = 6$ and 8 V, a short $t_{ep} = 1$ s was sufficient to fully cover the entire outer surface of the NiEFs with Ag (Figs. 2f–2k); at $t_{ep} = 1$ s ($V_{ep} = 5$ V), increased surface roughness (R_a) was observed compared with the basic NiEFs ($t_{ep} = 0$ s) (Table S1). As t_{ep} increased to 5 s, the Ag layer became thicker while the R_a value decreased. However, the value of R_a significantly increases again at $t_{ep} = 10$ s. The complete Ag plating of the Ni fibers is attributed to the enhanced metallic bonding of the fibers at the fiber junctions, as well as the higher conductivity

of Ag relative to that of Ni, which caused a significant decrease in the electrical contact resistance at the junctions. For example, at $V_{ep} = 5$ V, R_s decreased significantly from 3.01 to 0.49 Ω sq⁻¹ when t_{ep} increased from 0 (which corresponds to the bare NiEFs) to 1 s (Table 2). Note that R_s decreased to 0.05 Ω sq⁻¹ when t_{ep} increased from 0 to 10 s, while the average thickness (D_{avg}) of the AgMFs increased from 3.47 to 12.6 µm. As V_{ep} increased, the grain size and corresponding growth rate also increased considerably. The formation of a bump-like Ag texture became more pronounced as V_{ep} increased from 6 to 8 V (cf. Figs. 2f–2k).

To better understand the effect of Ag presence on the fiber surface, XRD analysis was conducted (Fig. S2). The XRD sample described here was the second AgMF with $V_{ep} = 5$ V and $t_{ep} = 3$ s. The XRD profile revealed four main peaks at $2\theta = 38.0^{\circ}$, 44.1°, 64.3°, and 77.3°, corresponding to the diffraction peaks of the (111), (200), (220), and (311) planes of Ag, respectively (JCPDS No. 04-0783). In addition, the surface compositions of the AgMFs and the chemical states of Ag were characterized by XPS analysis. The complete survey spectra for the AgMFs are shown in Fig. S3a, revealing the presence of Ag, C, and O. The C 1*s* peak at 284 eV was used as a reference for calibration of binding energies. The small O peak arose from some oxidation during the manufacturing processes [29, 40]. The Ag 3*d* XPS spectrum contains two sharp peaks (Fig. S3b) at 368.1 and 374.3 eV, corresponding to Ag $2d_{3/2}$ peaks was 5.8 eV, corresponding metallic Ag⁰ [41].

Fig. 3 shows cross-sectional transmission electron microscopy (TEM) and corresponding EDS elemental mapping images of the second AgMFs and Ni/Ag/Cu/Pt MFs (where the first Ag-electroplating method was used). The samples for TEM analysis were prepared by the focused ion beam (FIB) milling method (cf. Sec. 2.5.). The sequential electroplating of Ni and Ag yielded intact core–shell-structured metallized fibers. The

elemental mapping results reveal that the fiber contains a middle Ni shell on the interior PAN core and an outer Ag shell. The thicknesses of the outermost Ag, middle Ni, and innermost PAN layers are 1.53, 1.05, and 0.75 μ m (0.37 μ m radius of PAN fiber), respectively (Fig. 3a). The thickness of both the Ni and Ag layers can be controlled by changing the electroplating conditions (cf. Sec. 2.3.). The selected-area electron diffraction (SAED) pattern (shown in Fig. S4) also confirmed that the electroplated fibers were covered by Ag with a face-centered cubic structure [42].

Figs. 3e–3j show cross-sectional TEM and elemental mapping images of the Ni/Ag/Cu/Pt multilayered MF. Outside the Ni core, the cactus-like Ag layer is formed by the first Ag-electroplating (Fig. 3g). Then, a thick but discontinuous Cu layer was deposited on the Ag layer, and Pt was plated where the Cu layer had not grown. Therefore, the overall shape of the fibers is smoothly rounded despite the hierarchical structure of the Cu layer, as shown in Fig. S5. The thickness of each metal layer can be varied by adjusting plating parameters (cf. Sec. 2.4.).

The Ag shell provides high electrical conductivity to the metallized fibers, which exhibit remarkably low R_s values (Fig. 4 and Table 2). High transparency is also observed (Figs. 4a and 4c). Note that the $t_{ep} = 0$ s case in Fig. 4 corresponds to nickel electroplating only, without silver-electroplating. These characteristics are illustrated by the electrical conductivity–transmittance plot in Fig. 4b. A common figure of merit (FoM) used to assess and compare the performance of transparent electrodes is defined as FoM = T_r^{10}/R_s (Ω^{-1}) [43]. The FoM of the AgMFs prepared by the second method is up to 1.2 Ω^{-1} . The reprted FoM of Cu-electroplated wires [11] reached 1.8 Ω^{-1} ; FoM of Cu nanofibers [44] was 0.15 Ω^{-1} ; FoM of Ag nanowires [45] was 0.012 Ω^{-1} ; FoM of gold nanofibers [46] was 0.0094 Ω^{-1} ; FoM of graphene [47] was 0.009 Ω^{-1} ; FoM of PEDOT:PSS [48] was 0.0042 Ω^{-1} ; FoM of single-walled carbon nanotubes [49] was 0.00084 Ω^{-1} . Clearly, the FoM of the AgMFs is among the highest reported, very similar to the maximum FoM reported for CuEWs, and with a different T_r vs. R_s tradeoff curve from the CuEWs, such that the AgMFs achieve previously unachievable low values of R_s for high T_r .

Moreover, our AgMFs possess high T_r and low R_s compared to the conventional AgNW samples, cf. Table S2, which compares the substrates used in each method of producing transparent conducting films. These substrates have transparency values in the 80 - 91% range, while the AgMF film is free-standing and does not require a substrate, as demonstrated in Fig. 5 [50-52]. This free-standing feature is important, because it facilitates flexibility while minimizing the mass of the electrode. The T_r values of the AgMFs in the 400–800 nm wavelength (λ) range are almost constant, which is advantageous for their use in various optoelectronic applications, including solar energy harvesting, lighting, and display applications [53].

Table 3 compares the T_r and R_s values measured here with those from previous studies by An *et al.* [11, 12, 38]. The differences in the electrospinning and electroplating conditions, such as t_{es} (electrospinning time, cf. Sec. 2.), t_{ep} , V_{ep} , and I_{ep} (current density, cf. Sec. 2.), used for the different metal fibers arise from the corresponding differences in the extent of electroplating, metal type, *etc.* For the CuEWs, the average diameter, D_{avg} , values were not reported, whereas the R_s values are as low as 0.36–0.42 Ω sq⁻¹ with $T_r = 92–97\%$. The maximum T_r value of 97% for the CuEWs is 2% higher than that of the AgMFs ($T_r = 95\%$). The comparison of the R_s values of the AgMFs and CuEWs at the same $T_r = 92\%$, shows that the R_s value for the AgMFs ($R_s = 0.18 \Omega \text{ sq}^{-1}$) is one-half of that of the CuEWs ($R_s = 0.36 \Omega \text{ sq}^{-1}$). For the PtNFs, the minimum t_{es} of 90 s is 90 times that of the AgMFs ($t_{es} = 1$ s), resulting in a T_r of 55%, lower than that of the AgMFs. In addition, the minimum t_{ep} is 900 s, 900 times longer than that of the AgMFs ($t_{ep} = 1$ s). Nevertheless, the D_{avg} value of the PtNFs is 0.64 µm, considerably lower than that of the AgMFs ($D_{avg} = 3.47 \mu m$), which may be attributed to the experimental differences in the electroplating rates of Pt versus Ag. The R_s values of the PtNFs when $T_r = 55\%$ and 48% are 14.1 and 4.32 Ω sq⁻¹, respectively, which are larger than those of the other materials. Even though a rigorous comparison was not conducted in the present study, it seems that these results can be attributed to the different material properties and experimental conditions. The conditions used previously for the fabrication of NiEFs, $t_{es} = 5$ s and $I_{ep} = 0.22$ A cm⁻², (Table 3) were used in the present study to fabricate the framework of the AgMFs (cf. Sec. 2.3.). Even though the same t_{es} and t_{ep} values were used, the obtained T_r and R_s values for the NiEFs in the previous and present studies (Table 3) are different because the other experimental conditions, such as the devices, temperature, and humidity, varied somewhat (not shown here).

Stretching and bending tests were performed to examine the stretchability and flexibility of the AgMFs prepared by the second electroplating method. Fig. 5a shows data for the two different stretching tests. One applies uniaxial strain until the fiber mat breaks, whereas the other is a stretching cycle test over 80 cycles with repetitive stretching to 100% strain. In the uniaxial strain test, the resistance changes sharply above 140% strain and the fibers are broken at 160% strain (see Movie S1). In the stretching cycle test, a small change in the resistance is observed over 80 cycles. Fig. 5b shows the bending stability of the AgMFs according to the bending radius r_b . The electrical properties of the mat are maintained until r_b is set to 1 mm from a flat mat. Note that light scattering from the background lighting makes the stand-alone fibers distinctly visible in Fig. 5b. However, when the fibers are deposited on a transparent substrate as in Figs. 4c and 5c, they look quite transparent. A typical haze value of the metallized fiber for a transparency of 91.3% is about 5%.

Fig. 5c shows the bending cycle data for the AgMFs. The electrical performance of the AgMFs remains constant for 2000 cycles even when the bending direction is changed (see the inset image of Fig. 5c), which induces expansion or compression of the sample [54]. The photograph in Fig. 5d shows LED operation using the highly transparent AgMFs, thus demonstrating the electrical stability of the AgMFs. The excellent mechanical and electrical stability of the AgMFs is expected to allow durability and facilitate application in various flexible electrical devices.

To characterize the heating performance of the second AgMFs as a transparent heater, both the heat generation and heat cycle performances were characterized as functions of the applied voltage and time, as depicted in Fig. 6. Fig. 6a shows the change in temperature (T) of the AgMFs at different applied voltages (V_a). The value of T increased from 27.3 to 209.4 °C as $V_{\rm a}$ increased from 0 to 1.6 V. Reversible heating of the AgMFs was observed in the 0–1.6 V range, although the AgMFs are snapped at voltage values $V_a > 2.0$ V. Furthermore, the performance of the AgMFs (*i.e.*, T = 209.4 °C at $V_a = 1.6$ V) is comparable to those of AgNWs (T = 48-160 °C) [48,:55-58] and CNTs (T = 77-160 °C) [9,:59-61] used as transparent heaters in previous studies, as well as those of graphene-based complexes (T = 90-140 °C) [62, 63]. After a DC voltage $V_c = 0.8$ V is applied, T of the AgMFs rises sharply to 80 °C and is maintained continuously (the red line in Fig. 6a,). There was no surface change in the SEM image (Fig. S6). The O content of the AgMFs before and after heating [measured by energydispersive X-ray spectroscopy (EDX)] was 4.40 wt% and 4.48 wt%, respectively. In the heat cycle test (Fig. 6b), the durability of the AgMFs was maintained even after 30 cycles. Fig. 6c shows the stretchability of AgMFs at a constant voltage. The heating performance is maintained until the AgMFs are stretched from 0% to 160% (see Movie S2). This flexible, stretchable, and transparent heater can be used as a smart window material for residences or

automobiles that require high optical clarity [64]. In addition, the high thermal stability is expected to permit general applicability in wearable devices [65].

4. Conclusion

Core-shell-structured silver microfibers (AgMFs) with an intermediate Ni shell on PAN cores were fabricated using a combination of electrospinning and electroplating. The AgMFs produced using two different Ag-electroplating approaches were morphologically different, showing that the fabrication process can be tailored to produce specific morphologies to meet requirements for the intended end use of the MFs. The cactus-like morphology of the Ag fibers that maximizes the S/V ratio holds great promise for energy and environmental applications requiring maximal interfacial areas. In addition, the addition of other metallic layers was demonstrated, highlighting the potential for additional functions of such fibers. The AgMFs plated by the second method, with smoother surface morphology, exhibited remarkably high transmittance and low sheet resistance values, providing lower sheet resistance at transmittance up to 91% than any previously reported transparent electrodes. The AgMFs exhibited excellent mechanical performance and revealed no changes in electrical properties at 100% strain or after 80 repetitive stretching cycles or 2000 bending cycles. The AgMFs allowed resistive heating to 209 °C, which appears to be a record value compared to prior reports using Ag nanomaterials for transparent heaters. A heater temperature of 80 °C remained stable for over 500 s and 30 repetitive heating-cooling cycles. The thermal properties of the second type of AgMFs were thus significantly enhanced compared to those of previously reported materials. The second AgMFs have great potential for TCEs, smart windows, and wearable applications that require enhanced optical, mechanical, and thermal properties.

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Fig. 1. Schematic of the fabrication process of the AgMFs.



Fig. 2. SEM and (in the insets) silver elemental mapping images of AgMFs for the different Ag-electroplating methods, and different values of t_{ep} , and V_{ep} : (a–b) The first Ag-electroplating method with a fixed $V_{ep} = 8$ V and different (a) $t_{ep} = 10$ s and (b) $t_{ep} = 60$ s. (c–e) The second Ag-electroplating method with a fixed $V_{ep} = 2$ V and different (c) $t_{ep} = 1$ s, (d) $t_{ep} = 3$ s, and (e) $t_{ep} = 10$ s. (f–h) The second Ag-electroplating method with a fixed $V_{ep} = 10$ s. (i–h) The second Ag-electroplating method with a fixed $V_{ep} = 5$ V and different (f) $t_{ep} = 1$ s, (g) $t_{ep} = 3$ s, and (h) $t_{ep} = 10$ s. (i–k) The second Ag-electroplating method with a fixed $V_{ep} = 5$ V and different (i) $t_{ep} = 1$ s, (j) $t_{ep} = 3$, and (k) $t_{ep} = 10$ s. Yellow dots in the elemental mapping images represent Ag.



Fig. 3. Cross-sectional TEM images of (a) the second AgMF, and (e) the Ni/Ag/Cu/Pt multilayered MF. Elemental mapping images of (b–d) the second AgMF, and (f–j) the

Ni/Ag/Cu/Pt multilayered MF. Red, yellow, pink, and cyan dots in the elemental mapping images represent Ni, Ag, Cu, and Pt, respectively.



Fig. 4. (a) T_r spectra of the second AgMFs prepared using $V_{ep} = 5$ V with variation of t_{ep} . (b) T_r and R_s of the AgMFs as a function of t_{ep} : $t_{ep} = 10, 5, 3$, and 1 s from left to right, red stars. The T_r values of CuEWs, CuNFs, AgNWs, AuNFs, graphene, PEDOT:PSS, and SWNTs (single wall nanotube) are shown for comparison. (c) Photographs of the second AgMFs ($t_{ep} = 10, 5, 3$, and 1 s from left to right).



Fig. 5. (a) Results corresponding to uniaxial strain (red line) and stretching cycle tests (blue line). (b) Bending radius and (c) bending cycle tests [each color represents bending direction; the AgMFs were expanded (red line) and compressed (blue line)]. (d) A photograph of LED operation using the second AgMFs on a human hand ($t_{ep} = 3$ s, $V_{ep} = 5$ V).



Fig. 6. Results of (a) heating tests and (b) heating cycle tests for the second AgMFs ($t_{ep} = 3$ s and $V_{ep} = 5$ V). (c) IR images during stretching.

Case	Experimental result			
t _{ep} (s)	<i>T</i> _r (%)	$R_{\rm s} \left(\Omega \ { m sq}^{-1} ight)$		
10	25	$0.150{\pm}0.019$		
60	6	0.221±0.034		

Table 1. T_r at the wavelength $\lambda = 550$ nm and the R_s values for the 1st AgMFs ($I_{ep} = 0.125$ A cm⁻²).

Table 2. T_r at the wavelength $\lambda = 550$ nm and R_s values for the 2nd AgMFs ($V_{ep} = 5$ V).

Case	Experimental result				
$t_{\rm ep}$ (s)	D _{avg} (μm)	<i>T</i> _r (%)	$R_{\rm s}~(\Omega~{ m sq}^{-1})$		
0	3.47±0.17	96	3.01±1.05		
1	5.17±0.26	95	$0.49{\pm}0.01$		
3	6.50±0.40	92	0.18±0.03		
5	8.35±0.54	91	0.11 ± 0.04		
10	12.60±2.27	84	0.05 ± 0.01		

Table 3. T_r at the wavelength $\lambda = 550$ nm and the R_s values for comparison with previous studies by An *et al.* [11, 12, 38].

Case				Experimental result			Rof
Туре	$t_{\rm es}$ (s)	$t_{ep}(s)$	$V_{ m ep}$ or $I_{ m ep}$	$D_{\mathrm{avg}}(\mu \mathrm{m})$	$T_{\rm r}$ (%)	$R_{\rm s} \left(\Omega \ { m sq}^{-1} \right)$	К
AgMFs (2 nd)	5	0	5 V	$3.47{\pm}0.17$	96	$3.01{\pm}1.05$	Present study
		1		5.17±0.26	95	$0.49{\pm}0.01$	
		3		6.50 ± 0.40	92	0.18 ± 0.03	
		5		8.35 ± 0.54	91	0.11 ± 0.04	
CuEWs	1		3 V	-	97	0.42	[11]
	5	3			93	0.37	
	6				92	0.36	
PtNFs	90	900	0.11 A cm ⁻²	0.64	55	14.1	[12]
		1800		0.82	48	4.32	
		2700		1.22	36	0.88	
NiEFs	5	60	0.22 A cm^{-2}	4.75	93	0.73	[38]
	30			3.85	83	0.49	
	60			3.77	64	0.30	