Ag/MnO$_2$ Composite Sheath-Core Structured Yarn Supercapacitors

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One-dimensional (1D) yarn or fiber-shaped micro-supercapacitors that possess small diameter, volume and high mechanical degree of freedom, like flexibility or stretchability are highly needed due to the sudden high demand on appropriate power source for wearable electronics, micro-devices, and implantable medical devices. The composite sheath is fabricated on a commercially available CNT yarn substrate by alternating depositions of MnO$_2$ and Ag layers. Synergistic effect of high loading level of pseudocapacitive MnO$_2$ and reasonably improved rate-capability are achieved. In the composite sheath, the interconnected network provides electrical contact between MnO$_2$ aggregates and adjacent Ag layer. The conductive Ag inter layers shorten the solid-state charge diffusion length in the MnO$_2$. Moreover, generated electrons during the charge/discharge process can be collected rapidly by the adjacent Ag layer, therefore, the great extents of MnO$_2$ could be loaded onto the surface of CNT core fiber electrode without a significant rate-capability degradation. Due to the high MnO$_2$ loading level, the composite sheath-core yarn supercapacitor showed excellent specific areal capacitance (322.2 mF/cm$^2$) and according energy density (18.3 µWh/cm$^2$).
MnO$_2$ aggregates and adjacent Ag layer. The Ag conductive inter layers shorten the solid-state charge diffusion length in the MnO$_2$. Moreover, generated electrons during the discharging process can be effectively collected by the adjacent Ag layer, therefore, the great extents of MnO$_2$ (about 71.5 wt%) could be loaded onto the surface of CNT core yarn electrode without a significant rate-capability degradation or an impediment of mechanical flexibility. Thanks to high active material loading level, the composite sheath-core yarn supercapacitor showed excellent charge storage capabilities ($C_A = 322.3 \text{ mF cm}^{-2}$, $C_V = 208.1 \text{ F cm}^{-3}$) and according energy densities ($E_A = 18.3 \mu\text{Wh cm}^{-2}$, $E_V = 11.8 \text{ mWh cm}^{-3}$). More importantly, the composite sheath supercapacitor exhibited enhanced rate-capability, which is about 40% of the initial capacitance, and was retained when scan rate increased from 10 to 100 mV s$^{-1}$, which is two-fold higher than pristine MnO$_2$ sheath-core structured supercapacitor.

Results and Discussion
Previously reported literatures about yarn or fiber supercapacitors are mostly based on core-sheath structure configuration, where only the surface area of the electrically conductive core yarn is used as a loading site, increasing the MnO$_2$ loading level leads to a thick film formation with poor electrical conductivity. Unfortunately, from a view point of electrochemical reaction, this thick active material coating has several disadvantages such as longer solid-state ion diffusion length, high resistance, and low electrochemically active surface area. For this reason, previous reported plain core-sheath structured yarn supercapacitors which adopted the MnO$_2$ as an active material suffered from low MnO$_2$ loading levels mostly less than 20 wt%$^{2,18,19}$. The high active material loading without impairing the electrochemical performance or the mechanical strength is a very important research issue. Although a recent research achieved a record high MnO$_2$ loading level ($\sim 93$ wt%) inside the CNT yarn electrode by introducing liquid-state bscrolling technology, the limitation of using expensive, non-mass productive CNT sheets that are drawn from CNT forest still exists$^{20-22}$.

To address the limitations stated above, we propose a structural approach on yarn coating. The fabrication of the Ag/MnO$_2$ composite sheath on top of the commercially available CNT yarns aimed to maximize the pseudocapacitive active material (MnO$_2$) loading and maintain electrical conductivity, simultaneously. The pseudocapacitive MnO$_2$ and electrically conductive Ag were alternately deposited by electrochemical deposition method as shown in scheme Fig. 1A. We started from tens of meter-long, mechanically strong, and commercially available CNT yarns which required no further pre-treatment (Fig. 1B). Highly aligned CNT bundles characterized by SEM image shown in Fig. 1C can provide effective electron pathway and mechanical strength in yarn length direction$^{23}$. During the depositions, square wave potential was applied for core CNT yarn electrodes to form alternate MnO$_2$ and Ag based composite sheath. The magnified surface images of the Ag and MnO$_2$ deposits were shown in Fig. 1D,E, respectively. Rough surface Ag film was formed (Fig. 1D) while highly porous, even, and flower-like MnO$_2$ was observed (Fig. 1E), which is a morphological characteristic of the MnO$_2$ deposits$^{24}$.
Deposited amount of the sheath shell was controlled by deposition time or deposition number. As the deposition time for MnO₂ increases, the specific capacitance also increases while the rate-capability decreases, which is obtained by low-rate CV (at 10 mV s⁻¹) as shown in Fig. 3A1. Reversely, increasing Ag deposition number results in capacitance decrease and retention increase (Fig. S1B). For further electrochemical characterization, optimized deposition for composite sheath was applied (total of MnO₂, deposition time for 50 minutes, Ag deposition time 90 seconds.) Deposition of composite sheath was followed by repetitive deposition method that does not require any post treatment (vacuum or annealing treatment).

Cross-sections of composite sheath-core yarns were prepared by cutting them along their diameters using a Ga ion beam (7 nA beam current) in a Focused Ion Beam (FIB, Nova 200) operated at 30 kV. Then, the cut areas were cleaned via Ga ion polishing by etching several micrometers of yarn length with consecutively decreasing ion beam-currents ranging from 5.0 to 0.3 nA. The cleaned-cut yarns were next transferred to a Zeiss Supra 40 SEM to perform the microscopy (at 15 kV) and elemental EDAX mapping analysis (at 20 kV). Clear views of the yarn's cross-sections were attained by orienting the yarn's cut plane parallel to the electron-beam final aperture/detector in the SEM. Related SEM images and EDAX mapping results are provided in Figs 2A–C and S2. We define the sheath/core ratio as the ratio of the sheath thickness to the yarn diameter, as measured by using SEM images for twisted yarn. From SEM micrograph observation, it was estimated that the 50 μm composite yarn has a sheath/core ratio of about 0.106. EDX mapping and quantitative analyses (total of 3 areas per each sheath and core regions) showed almost no infiltration of the MnO₂ or Ag into the fiber core (Mn and Ag atomic concentration at sheath and core are 18.5(±2.7) at.% and 1.7(±0.3) at.% against <4 at.% and <0.5 at.% respectively, as shown in Fig. S2). XRD pattern of hybrid MnO₂/Pt supercapacitor is shown in Fig. S3. From the electrochemical impedance spectroscopy (EIS) measurement (Fig. 2D), normalized equilibrium series resistance (ESR) at 1 kHz for solid-state composite sheath-core supercapacitor is as small as 0.3 kΩ and it exhibited high slope of Nyquist curve, implying also good capacitive characteristic. On the other hand, the pristine MnO₂ sheath-core yarn supercapacitor showed larger ESR value (1.3 kΩ at 1 kHz) and lower Nyquist slope, which originated from long solid-state ion diffusion length and limited ion accessibility by thick MnO₂ coating. As a result, the composite sheath-structured supercapacitor exhibited improved rate-capability that 42.6% capacitance was retained when the scan rate was increased from 10 to 100 mV s⁻¹ while the pristine MnO₂ retained only 21.2% not showing box-like CV curves (Figs 2E and S4). The CV curves for composite and pristine sheath structured supercapacitors compared in Fig. 2F also support the improved rate-capability performance.

Electrochemical energy storage performances of the solid-state, Ag/MnO₂ composite yarn supercapacitor comprising of two symmetric yarn electrodes coated by PVA-LiCl gel electrolyte are shown in Fig. 3. Figure 3A shows CV curve comparison (at 10 mV s⁻¹) between pristine CNT and Ag/MnO₂ composite sheath-deposited CNT yarn supercapacitors. Box-like rectangular CV curve of composite sheath yarn supercapacitor without any faradic redox peaks indicates excellent capacitive characteristic. The charge storage capability of the yarn supercapacitor could be dramatically improved by present composite sheath yeast deposition. It is notable that the contribution of the sheath coating to total charge storage performance is impressively dominant that about 50 times larger CV area is obtained. Accordingly, calculated specific areal capacitances of the single electrode are roughly proportional to the loading amount of MnO₂ by electrochemical deposition (inset of Fig. 3D). Unless otherwise noted, capacitances are single-electrode values that are normalized with respect to the total dimension of the electrochemical active material (presently the Ag/MnO₂ composite sheath-CNT core yarn). The areal capacitance of pristine CNT yarn supercapacitor was about 4.9 mF cm⁻² at 10 mV s⁻¹, which is comparable to the values from previous researches (1.97–8.66 mF cm⁻²) and it dramatically increased to 322.2 mF cm⁻² when 71.5 wt% MnO₂ loaded for the composite sheath yarn supercapacitor. CV curves with various scan rates (at 2, 30, and 100 mV s⁻¹) and galvanostatic charge/discharge curves with various current densities (at 50, 100, and 200 mA cm⁻²) for composite sheath yarn supercapacitor are plotted in Fig. 3B, C, respectively. Moreover, this pseudocapacitive characteristic can also be obtained when working voltage range is extended up to 1.2 V (Fig. S5). Linear time-dependent change characteristic without any redox reaction at a constant current density in galvanostatic curves also indicates the capacitive characteristic of present composite yarn supercapacitor. The areal capacitance of yarn type supercapacitors shows only moderate capacitance enhancement (18.6−512%) compared with previously reported plain core-shell structured yarn supercapacitors.

Areal and volumetric specific capacitances are plotted in Fig. 3E. The highest values of the areal and volume normalized specific capacitances were calculated to be 322.2 mF cm⁻² and 208.1 F cm⁻³, respectively, from CV curves at 2 mV s⁻¹ scan rate. Resulting energy and power densities of the composite sheath supercapacitor are compared with previously reported other yarn supercapacitors as shown in Fig. 3F and Table S1. At 1.2 V, the calculated areal and volumetric energy densities of present composite sheath yarn supercapacitor were 18.3 J Wh cm⁻² and 11.8 mWh cm⁻³, respectively (normalized by total device volume including both yarn electrodes and PVA-LiCl gel electrolyte coating). The composite sheath yarn supercapacitor's energy density is higher than other flexible yarn type supercapacitors as shown in Table S1. The volumetric energy density value is higher than previously reported ZnO-doped MnO₂ Core/shell nanocables (0.04 mWh cm⁻³)⁷, poly(vinyl alcohol)/graphene hybrid fiber (5.97 mWh cm⁻³)⁸, carbon black/graphene hybrid fiber (2.8 mWh cm⁻³)⁹ and twisted CNT/MnO₂ wire (1.73 mWh cm⁻³)⁹.

Capacitance retention performance of the solid-state, composite sheath yarn supercapacitor versus charge/discharge cycle was characterized in Fig. 4A. Area of initially measured CV curve was retained about 80% after 1000 charge/discharge cycle at 10 mV/s (inset of Fig. 4A). Despite the high loading of brittle active materials, present sheath-core supercapacitor mechanically strong and flexible. About 130 MPa yarn strength is achieved
at 1.6% tensile deformation as shown in Fig. S6. To demonstrate the excellent flexibility, capacitance retention was also measured during dynamically applied bending with 170 bending degree and found to be about 90% capacitance retention after 1000 times bending test as shown in Fig. 4B. The solid-state supercapacitors can be connected in serial or parallel directions. The proportional increase in voltage and current were observed in CV curves for serial and parallel connected supercapacitors, respectively (Fig. 4C). Moreover, to demonstrate the possibility of wearable energy storage application, 2 cm long yarns were sewn into in commercial textile as shown in Fig. 4D. Due to high mechanical strength, the arrays of original yarns in the textile were successfully replaced by the embedded yarn electrodes.
Conclusion
Herein, we achieved synergistic effect of high loading level of pseudocapacitive MnO₂ and reasonably improved rate-capability. The interconnected networks provide electrical contact between MnO₂ aggregates and adjacent Ag layer in the composite sheath. The conductive Ag inter layers shorten the solid-state charge diffusion length.

Figure 3. (A) CV curves of pristine CNT and Ag/MnO₂ composite sheath supercapacitor measured at 10 mV s⁻¹. (B) CV (measured from 2 to 100 mV s⁻¹) and (C) Galvano-static curves (measured from 50 to 200 mA cm⁻²) of Ag/MnO₂ composite sheath supercapacitor. The supercapacitors comprise two symmetric electrodes coated by PVA/LiCl gel electrolyte. (D) Areal capacitances of yarn supercapacitor versus active material loading of composite sheath yarn supercapacitor compared with reported yarn supercapacitors with various amounts of active materials loading: (a) 1.45 wt% MnO₂ coated CNT/nylon coil fiber, (b) 4.1 wt% MnO₂ coated CNT yarn, (c) 17.2 wt% MnO₂ coated CNT coil yarn, (d) 50 wt% polyaniline coated CNT/rubber fiber. The inset shows areal capacitances with increasing MnO₂ loading in pristine MnO₂ sheath supercapacitor. (E) Areal and volumetric capacitances from 2 to 100 mV s⁻¹. (F) Ragone plots comparing volumetric energy and power densities for the present all-solid-state supercapacitor: (a) 2.44 mW cm⁻³–0.04 mWh cm⁻³ ZnO-doped MnO₂ Core/shell nanocables, (b) 26.9 mW cm⁻³–5.97 mWh cm⁻³ poly(vinyl alcohol)/graphene hybrid fiber, (c) 1.2 W cm⁻³–2.8 mWh cm⁻³ carbon black/graphene hybrid fiber, (d) 0.79 W cm⁻³–1.73 mW cm⁻³ twisted CNT/MnO₂ wire.
in the MnO$_2$. Moreover, generated electrons during the charge/discharge process can be collected rapidly by the adjacent Ag layer, therefore, the great extents of MnO$_2$ could be loaded onto the surface of CNT core fiber electrode without a significant rate-capability degradation (42.6% in comparison to pristine MnO$_2$ sheath’s 21.2%). Moreover, thanks to high MnO$_2$ loading level of the composite sheath-core yarn supercapacitor, excellent charge storage capabilities (specific areal and volumetric capacitances: 322.2 mF cm$^{-2}$ and 208.1 F cm$^{-3}$) and according energy density (area and volumetric energy densities: 15.3 μWh cm$^{-2}$ and 8.3 mWh cm$^{-3}$) were achieved.

**Method**

**Preparation of Ag/MnO$_2$ Composite Sheath Supercapacitor.** Commercial CNT yarn (Muratec, Japan) is used as a current collector. One end of the 30–40 μm diameter CNT yarn electrode was connected electrically to Cu wire using silver paste and cover it with epoxy. First of all, the electrochemical deposition of MnO$_2$ onto the CNT yarn electrode was performed using a potentiostatic method. About 1.3 V (vs Ag/AgCl as a reference electrode and Pt mesh as a counter electrode in a three-electrode system) was applied for 5 minutes using an electrochemical analyzer (CHI 627b system, CH Instruments, Austin, TX) in a solution containing 0.02 M MnSO$_4$.5H$_2$O and 0.2 M Na$_2$SO$_4$. After the first MnO$_2$ deposition, the deposition of Ag performed onto MnO$_2$-depositied CNT yarn electrode using a potentiostatic method. About $-1$ V (vs Ag/AgCl as a reference electrode and Pt mesh as a counter electrode in a three-electrode system) was applied for 10 seconds using an electrochemical analyzer in solution containing 0.02 M AgNO$_3$ and 0.2 M Na$_2$SO$_4$. The total MnO$_2$ and Ag alternative depositions were performed 10 and 9 times, respectively. The PVA–LiCl gel electrolyte containing 3 g PVA (Mw 146,000–186 000) and 6 g LiCl in 30 mL deionized water was prepared. The solution was heated on 160–180 °C until they dissolve in the water. Fabrication of the Ag/MnO$_2$ composite sheath yarn supercapacitor was completed by coating the 10 wt% PVA–LiCl gel electrolyte on parallel, symmetric Ag/MnO$_2$ composite sheath electrodes. PVA wt% is key parameter to control the viscosity of prepared gel electrolyte. For ideal case, the gel electrolyte needs to have proper viscosity so that can be easily coated on the surface of the yarn electrode when coating. At the same time, it should maintain yarn shape with enough viscosity after drying process. We experimentally find out that 10 wt% is the optimized concentration condition. And we used PVA-LiCl electrolyte
because MnO$_2$ is degraded because it reacts with acids and bases. The reduction reaction with the acid cause MnO$_2$ to be degraded by the following reaction.

$$\text{MnO}_2 + 2\text{H}_2\text{SO}_4 \rightarrow 2\text{MnSO}_4 + \text{O}_2 + 2\text{H}_2\text{O}$$

$$\text{MnO}_2 + 4\text{HCl} \rightarrow \text{MnCl}_2 + \text{Cl}_2 + 2\text{H}_2\text{O}$$

The oxidation reaction with the alkali cause MnO$_2$ to be degraded by the following reaction.

$$2\text{MnO}_2 + 4\text{KOH} + \text{O}_2 \rightarrow 2\text{K}_2\text{MnO}_4 + 2\text{H}_2\text{O}$$

All products used in this experiment were purchased from Sigma-Aldrich without additional chemical treatments. The weight of the CNT yarn and electrodes was measured with an electronic balance. The mass loading weight was calculated from the formula

$$\text{loaded mass weight (sheath)} = \frac{\text{Total electrode weight} - \text{CNT yarn weight (core)}}{\text{Total electrode weight}}.$$}

**Characterization.** All electrochemical measurements for the solid-state yarn supercapacitor used the two-electrode configuration and the electrochemical analyzer. X-ray diffraction data were measured by x-ray diffractometer (SmartLab, Rigaku). Cyclic voltammetry and chronopotentiometry measurements for all investigated supercapacitors were made using a electrochemical analyzer (CHI 627b, CH Instrument). Nyquist curves were measured using another electrochemical analyzer (Reference 600, Gamry Instrument). Scanning electron microscope images were obtained using a Zeiss Supra 40 SEM.

**Specific Capacitance Calculation.** The capacitance of Ag/MnO$_2$ composite sheath supercapacitor was calculated from the CV curves. From

$$C = \frac{I}{(dV/dt)}$$

where $I$ is average current and $dV/dt$ is the voltage scan rate. The specific areal capacitance for each electrode in a supercapacitor having equal anode and cathode capacitances was calculated using $C_s = 4C/A$, where $A$ is the total surface area of the anode and cathode. Also, the specific volumetric capacitance for each electrode in a supercapacitor having equal anode and cathode capacitances was calculated using $C_v = 4C/V$, where $V$ is the total volume of the anode and cathode. The specific energy density was calculated from the equation

$$E = \frac{1}{2}CV^2.$$}

**Cross-sectional Characterization.** The Ag/MnO$_2$ composite sheath yarn was cut and polished along its diameter using Ga ions in a Focused Ion Beam (FIB NOVA 200). Microstructural and chemical analyses were carried out at Zeiss SUPRA 40 Gemini EDAX and Zeiss-LEO Model 1530. Samples were coated by sputtering with gold for imaging purposes. ImageJ was used as a tool for image-based quantitative digital analysis of the porosity of the cross-section of the composite yarn.

**References**

1. Choi, C. et al. Flexible supercapacitor made of carbon nanotube yarn with internal pores. *Adv. Mater.* 26, 2059 (2014).
2. Ren, J. et al. Twisting carbon nanotube fibers for both wire-shaped micro-supercapacitor and micro-battery. *Adv. Mater.* 25, 1155 (2013).
3. Yu, D. et al. Controlled functionalization of carbonaceous fibers for asymmetric solid-state micro-supercapacitors with high volumetric energy density. *Adv. Mater.* 26, 6790 (2013).
4. Choi, C. et al. Elastomeric and dynamic MnO2/CNT core-shell structure coiled yarn supercapacitor. *Adv. Energy Mater.* 6, 1502119 (2016).
5. Choi, C. et al. Stretchable, weaveable coiled carbon nanotube/MnO$_2$/polymer fiber solid-state supercapacitor. *Sci. Rep.* 5, 9387 (2015).
6. Zhang, Z. et al. Superelastic supercapacitors with high performances during stretching. *Adv. Mater.* 27, 356 (2015).
7. Lee, J. A. et al. Ultrafast charge and discharge bisrolled yarn supercapacitors for textiles and microdevices. *Nat. Commun.* 4, 1970 (2013).
8. Yang, P. et al. Hydrogenated Zno Core-shell nanocables for flexible supercapacitors and self-powered systems. *ACS Nano* 7, 2617 (2013).
9. Chen, S. et al. Conductive, tough, hydrophilic poly(vinyl alcohol)/graphene hybrid fibers for wearable supercapacitors. *J. Power Sources* 319, 271 (2016).
10. Ma, W. et al. Hierarchically porous carbon black/graphene hybrid fibers for high performance flexible supercapacitors. *RSC Adv.* 6, 501112 (2016).
11. Xie, K. & Wei, B. Materials and structures for stretchable energy storage and conversion devices. *Adv. Mater.* 26, 3592 (2011).
12. Xie, K. et al. Polyallylene nanowire array encapsulated in titania nanotubes as a superior electrode for supercapacitors. *Nanoscale.* 3, 2202 (2011).
13. Xie, K. et al. Highly ordered iron oxide nanotube arrays as electrodes for electrochemical energy storage. *Electrochem. Commun.* 13, 657 (2011).
14. Wei, W., Cui, X., Chen, W. & Ivey, D. G. Manganese oxide-based materials as electrochemical supercapacitor electrodes. *Chem. Soc. Rev.* 40, 1697 (2011).
15. He, Y. et al. Freestanding three-dimensional graphene/MnO$_2$ composite networks as ultralight and flexible supercapacitor electrodes. *ACS Nano* 7, 174 (2013).
16. Xu, C., Kang, F., Li, B. & Du, H. Recent progress on manganese dioxide based supercapacitors. *J. Mater. Res.* 25, 1412 (2013).
17. Gogotsi, Y. & Simson, P. True performance metrics in electrochemical energy storage. *Science* 334, 917 (2011).
18. Zhang, M., Atkinson, K. R. & Baughman, R. H. Multifunctional carbon nanotube yarns by downsizing an ancient technology. *Science* 306, 1358 (2004).
19. Dong, X. et al. Synthesis of a MnO$_2$-graphene foam hybrid with controlled MnO$_2$ particle shape and its use as a supercapacitor electrode. *Carbon* 50, 4865 (2012).
20. Ren, J., Bai, W., Guan, G., Zhang, Y. & Peng, H. Flexible and weaveable capacitor wire based on a carbon nanocomposite fiber. *Adv. Mater.* 25, 5965 (2013).
21. Choi, C. et al. Improvement of system capacitance via weaveable superelastic bisrolled yarn supercapacitors. *Nat. Commun.* 7, 13811 (2016).
22. Choi, C. et al. Weavable asymmetric carbon nanotube yarn supercapacitor for electronic textiles. *RSC Adv.* 8 13112 (2018).
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Author Contributions
J.H.K. and C.S.C. conceived the idea and designed the experiments; J.H.K., C.S.C., J.M.L. and M.J.A. contributed mechanical/electrochemical characterization; J.M.L., C.S.C., S.J.K. and R.H.B. wrote the paper. All authors discussed the results and commented on the manuscript.

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