Fully Packaged Carbon Nanotube Supercapacitors by Direct Ink Writing on Flexible Substrates

Bolin Chen,† Yizhou Jiang,‡ Xiaohui Tang,† Yayue Pan,*† and Shan Hu*†‡

†Department of Mechanical Engineering, Iowa State University, Ames, Iowa 50011, United States
‡Department of Mechanical and Industrial Engineering, University of Illinois, Chicago, Illinois 60607, United States

ABSTRACT: The ability to print fully packaged integrated energy storage components (e.g., supercapacitors) is of critical importance for practical applications of printed electronics. Due to the limited variety of printable materials, most studies on printed supercapacitors focus on printing the electrode materials but rarely the full-packaged cell. This work presents for the first time the printing of a fully packaged single-wall carbon nanotube-based supercapacitor with direct ink writing (DIW) technology. Enabled by the developed ink formula, DIW setup, and cell architecture, the whole printing process is mask free, transfer free, and alignment free with precise and repeatable control on the spatial distribution of all constituent materials. Studies on cell design show that a wider electrode pattern and narrower gap distance between electrodes lead to higher specific capacitance. The as-printed fully packaged supercapacitors have energy and power performances that are among the best in recently reported planar carbon-based supercapacitors that are only partially printed or nonprinted.

KEYWORDS: direct ink writing, supercapacitor, carbon nanotube, 3D printing, flexible electronics

1. INTRODUCTION

Printed electronics offer the promise of low cost, rapid prototyping, easy customization, and even flexible and conformal devices when printed on soft substrates. Printed electronic components have been extensively studied, such as transistors,1,2 sensors,3 displays,4 etc. Since power supply is an indispensable component of any electronic systems, to fully realize the promise of printed electronics, printed power sources have to be developed. Recently, supercapacitors have emerged as a promising energy storage device, due to their high power, long cycle life, and ability to bridge the energy and power gap between batteries and conventional dielectric capacitors. Supercapacitors are widely used in electronic systems where fast and frequent charging/discharging is required. Hybrid power sources integrating batteries and supercapacitors together provide both high energy and high power at the same time.

A fully packaged supercapacitor consists of integrated functional parts and structural parts. Functional parts are components that contribute directly to the energy storage, including electrode, electrolyte, and separator, while structural parts are those that do not store energy but are necessary for achieving stable performance, including sealed casing, support substrates, etc. Since each component is made of very different materials, printing a fully packaged supercapacitor is challenging but necessary for practical applications of printed supercapacitors.

Additive Manufacturing (AM), also known as 3D Printing or Direct Digital Manufacturing, is a class of technologies that fabricate a three-dimensional physical model directly from its digital design, by accumulating materials, usually in a layer-by-layer way. Considering the unique strengths such as maskless, one-step operation and little material waste, AM technologies such as fused deposition modeling (FDM), inkjet printing, and direct ink writing (DIW) have been widely investigated for printed electronics in the past decade.5–14 The major drawback of inkjet printing is the limited variety of printable materials: materials need to have both flowability and jettableity, which means only low-viscosity materials can be processed in this technique. The major disadvantage of FDM is that its printing resolution is low, usually in the range of 50–200 μm. In addition, the feedstocks have to be filaments that can be melted in a low temperature (<200 °C). Compared with the inkjet printing and FDM technologies, the DIW technology has much wider choice of feedstocks ranging from highly viscous polymer gels to highly shear thinning colloidal suspensions, which allows an unrivaled freedom for choosing and preparing proper inks for the multimaterial supercapacitor design. The DIW method has demonstrated the capability of producing highly accurate, repeatable, and complex microstructures of any shape from a wide choice of materials in room temperatures, directly from digital design without the use of any masters or masks.15–18

Thus far, most studies on printed supercapacitors have focused on printing the electrode materials but rarely the fully...
packaged cell. The printed electrodes are later assembled with other nonprinted components. For example, Xie et al. used a laser to pattern graphene/nickel electrodes on poly(ethylene terephthalate) (PET) film and later manually casted poly(vinyl alcohol)/lithium chloride gel electrolyte onto the electrodes and sealed the cell with PET by hot-melt glue.19 Wang et al. used a commercial inkjet printer to pattern multiwall carbon nanotube-based nanocomposite electrodes on paper and later sandwiched the two electrodes with an electrolyte-soaked separator in between. Finally, they used adhesive tape to seal the cell.20 Tehrani et al. used a screen presser and a series of masks to print carbon—silver electrodes, gel electrolyte, and adhesives and finally assembled two electrolyte-loaded electrodes into a sealed cell with adhesives.21 Although excellent electrochemical performances have been reported from these studies, the printing/patterning techniques used in these studies (laser processing, inkjet printing, screen printing) involve multiple aligning, transferring, and assembling steps to finally produce a fully packaged supercapacitor. These intermediate steps add uncertainties to the volume, mass, and performance of the as-made supercapacitors and decrease repeatability, which is very important for practical manufacturing.

Herein, we report for the first time the use of DIW technology for printing fully packaged flexible supercapacitors.
40 mL of deionized (DI) water, and stirred at 300 rpm using a magnetic stirring bar for 5 min. The mixture was then placed inside an ice−water bath and subjected to 10 min of probe sonication (QSonica Q500, 50% power) to disperse SW-CNT uniformly in the DI water. To prepare PVA-based gel electrolyte, 6 g of PVA powder (Mowiol 18-88, Sigma-Aldrich, USA) and 12 g of lithium chloride powder (>99.0%, Sigma-Aldrich, USA) were added to 40 mL of DI water and stirred on a 85 °C hot plate until the mixture became a clear and glue-like gel. Silicone rubber was purchased from Nanda 705 RTV Silicon Rubber (Liyang, China) and used as received.

2.2. Direct Ink Writing Setup. In this study, a DIW testbed was developed to fabricate the fully packaged supercapacitor, including electrode, electrolyte, and sealed package. DIW was achieved by dispensing inks through syringe needles in close proximity to a moving platform in a layer-by-layer way. As shown in Figure 1, the testbed is equipped with a syringe needle, an air-pressure controller to regulate ink dispensing rate, a heated platform, a charge-coupled device (CCD) camera, an X−Z stage, and a Y stage. The syringe needle is mounted on an X−Z stage, and the platform is placed on a Y stage. The CCD camera is used to monitor the ink dispensing in real time and to measure the distance between tip and platform. The platform temperature is controlled and can be varied between 15 and 250 °C.

2.4. DIW of SW-CNT Supercapacitor. The DIW process for rapid production of the proposed supercapacitor consists of four steps as illustrated in Scheme 1a. During the entire printing process, the Kapton polyimide substrate (McMaster-Carr, USA) is fixed on the platform, which is controlled by the Y stage. First, 2 mL of SW-CNT ink is printed onto the Kapton substrate with a needle tip of 0.20 mm inner diameter using 1.0 psi dispensing air pressure and 200 mm/s writing speed (black region in Scheme 1a). With a platform temperature of 100 °C, the CNT solution is fully dried within 5 min. To remove surfactant and improve the conductivity of the CNT electrode, the printed part is soaked in concentrated nitric acid for 20 min, quenched in DI water, and dried in air. Second, the dispensing tip moves up by 0.1 mm in the Z direction to print the second layer. In the second layer, PVA electrolyte is then printed in the light green region in Scheme 1a. To print the PVA layer, a needle tip of 0.41 mm inner diameter is placed above the Kapton substrate with a 0.4 mm gap distance (also known as the standoff distance). The pumping pressure is set at 25 psi, and the substrate speed is set at 5 mm/s in the Y direction. Finally, a layer of silicone rubber is printed in the transparent region in Scheme 1a. The needle tip (0.84 mm inner diameter) is placed above the Kapton film substrate with a ~0.6 mm standoff distance. A pumping pressure of 30 psi and a platform moving speed of 10 mm/s are used to print the silicone rubber layer. The silicone rubber layer covers the whole supercapacitor, leaving only two SW-CNT electrode legs exposed for electrical connection.

To study how the electrode and electrolyte pattern design affect the overall performance of the supercapacitor, single-cell supercapacitor designs with different geometry are printed following the procedure given in Scheme 1a. Specifically, the width of the electrode ("W" in Scheme 1a) and the distance between electrodes ("D" in Scheme 1a) are varied, and their effects on the supercapacitor performances are compared. A multicell supercapacitor design is also printed. The design shown in Scheme 1b is an array of three identical supercapacitors connected in series.

2.5. Material and Electrochemical Characterization. The viscosity of the CNT ink, PVA electrolyte ink, and silicon rubber

Figure 2. (a) SEM of as-printed CNT electrode surface without acid treatment; inserted optical image shows the developed SW-CNT ink. (b) SEM of acid-treated electrode surface with inserted high-magnification image (scale bar 200 nm). (c) Change of resistance of printed traces with ink volume used before and after nitric acid treatment. (d) Full XPS spectrum of raw SW-CNT and as-printed SW-CNT without acid treatment and with acid treatment.
ink are measured by Malvern Kinexus ultra+ rheometer at 25 °C, with shear rate ranging from 1 to 100 s⁻¹. A microscopic image of the as-printed SW-CNT electrode is taken from a FEI Quanta 250 FEG Scanning Electron Microscope (SEM) at an operating voltage of 10 kV. The XPS measurements of surface composition of the electrodes are performed using a Kratos Amicus/ESCA 3400 instrument. The sample is irradiated with 240 W unmonochromated Mg Kα X-rays, and photoelectrons that emit at 0° from the surface normal are energy analyzed using a DuPont type analyzer. The pass energy is set at 150 eV, and either a Shirley or a linear baseline is removed from all reported spectra. CasaXPS is used to process raw data files.

Galanostatic charge–discharge (GCD), cyclic voltammetry (CV), and electrochemical impedance spectroscopy (EIS) are performed on the printed supercapacitors with a Gamry Reference 3000 electro-chemical station. The capacitances (C) of each device at different current densities are calculated from the discharge curves obtained from GCD tests using the formula

$$ C = \frac{I}{\Delta U} $$

(1)

where I is the applied discharge current (amp), \( \Delta U \) is discharge time (second), and \( \Delta U \) (volts) is the discharge voltage after IR drop is removed.

The gravimetric, areal, and volumetric specific capacitances of each device at different current densities are calculated from the discharge curves obtained from GCD tests using the following equations

$$ C_{\text{sp, g}} = \frac{C}{m} $$

(2)

$$ C_{\text{sp, areal}} = \frac{C}{A} $$

(3)

$$ C_{\text{sp, vol}} = \frac{C}{V} $$

(4)

where m (g), A (cm²), and V (cm³) are the mass, area, and volume of the active materials of all electrodes. Here, active materials are the printed CNTs (dark gray region in Scheme 1) that overlap with the electrolyte (green region in Scheme 1).

As pointed out by Gogotsi and Simon et al., volumetric energy density and power density can provide more reliable performance metrics for porous nanomaterial-based thin film devices compared to gravimetric capacitance. As a result, the volumetric energy density (Wh cm⁻³) of each device is calculated using

$$ E = \frac{0.5 \times C \times \Delta U^2}{3600 \times V} $$

(5)

The volumetric power density (W cm⁻³) of the device is calculated from

$$ P = \frac{E}{\Delta t} \times 3600 $$

(6)

3. RESULTS AND DISCUSSION

The viscosity measurements of the inks for the SW-CNT electrode, PVA electrolyte, and silicone-sealed package are plotted in Figure S1. The morphology of printed SW-CNT electrode before and after acid treatment is shown in Figure 2a and 2b. In as-printed electrodes, a layer of surfactant on top of the SW-CNT can be seen in Figure 2a. After acid treatment, the surfactant top layer seems to be removed, and the highly porous structure of the SW-CNT can be clearly seen in Figure 2b. The inserted image in Figure 2a shows the prepared SW-CNT ink. Figure S2 shows the cross-section SEM of SW-CNT electrode with an average thickness of 1.26 μm. Different volumes of SW-CNT ink (0.2, 0.3, and 0.4 mL) are used to print traces 10 mm long and 3 mm wide. Resistances of traces with and without nitric acid treatment are measured and compared after the traces dry (Figure 2c). The resistance drops as the volume of ink used increases. The resistance drop at low ink volume is mainly explained by percolation theory, which says that for nanotube- and nanowire-type materials to form a conductive 2D thin film, the nanotubes/nanowires must form a connected network.23,24 When the volume of CNT ink used is still low, a continuous connected network of CNT has not formed yet across the whole electrode area and the reduction in electrode resistance with more ink is mainly ascribed to the gradual buildup of such a percolated network. With high ink volume, the deposited CNTs are more than enough to form a percolated network. Hence, the resistance drop is mainly due to an increase of electrode thickness (t), following Ohm’s law $R_{\text{CNT}} = \rho_{\text{CNT}} \frac{W}{l}$, where $W$ is the width of the electrode and $l$ is the length of the electrode. After nitric acid treatment, the resistance of the electrode traces all drop compared with untreated ones. The full-range XPS of printed SW-CNT with/without acid treatment are compared in Figure 2d, with the XPS of raw SW-CNT added as a reference. The high sodium (Na) content in the SW-CNT without acid treatment comes from the Na of the SDS surfactant used for preparing the SW-CNT printing ink. More importantly, after acid treatment, no sodium (Na) Auger (~495 eV) signal is detected, which suggests significant removal of the SDS surfactant. Removing the highly insulating SDS surfactant improves the electrical contact of nearby nanotube bundles and reduces the overall electrical resistance of SW-CNT electrodes. High-resolution XPS of the C 1s, O 1s, and N 1s of the raw SW-CNT and the printed SW-CNT electrode with and without acid treatment are shown in Figures S3–S. Compared with the C 1s of the raw SW-CNT and the as-printed SW-CNT without acid treatment, there is a 0.3 eV upshift of the C 1s binding energy after acid treatment, indicating oxidation of SW-CNT electrode by the nitric acid.25 The O 1s peak ($531.2$ eV) in as-printed SW-CNT electrode without acid treatment is attributed to the oxygen from the sulfate (SO₄²⁻) groups in the SDS surfactant. In acid-treated samples, the O 1s peak is shifted to $534.2$ eV, which is attributed to oxidation of SW-CNT by nitric acid. As a result, the volumetric energy density (Wh cm⁻³) of each device is calculated using

$$ E = \frac{0.5 \times C \times \Delta U^2}{3600 \times V} $$

(5)

The volumetric power density (W cm⁻³) of the device is calculated from

$$ P = \frac{E}{\Delta t} \times 3600 $$

(6)
of S11–S13 between 0 and 1 V at a current density of 1.32 A cm$^{-3}$ and the calculated specific capacitance are shown in Figure 3a. All GCD curves have symmetric charge and discharge profiles; however it is obvious that S11 with the smallest electrode–gap distance of 1 mm has the highest specific capacitance of 13.37 F cm$^{-3}$ (1.9 F g$^{-1}$ and 1.68 mF cm$^{-2}$) followed by S12 and S13. There is a clear decreasing trend for the specific capacitance as the electrode–gap distance increases. The reason for this observed trend could be understood from the EIS of the three supercapacitors (Figure 3b). The Nyquist plots are almost-vertical straight lines at low frequencies, characteristic of capacitive behaviors. Intersect of the Nyquist plot with the real axis at high frequency is the equivalent series resistance (ESR) of the supercapacitor, which consists of resistances from both the electrode and the electrolyte. Large ESR will cause a large drop of voltage at the beginning of discharge, which decreases the usable discharge voltage window and capacity of the supercapacitor. As shown in Figure 3b, as the electrode–gap distance is increasing, the ESR increases from 95 ohms for S11 to 478 ohms for S13, mainly caused by the increased electrolyte resistance since the electrodes in all three cells are identical. The larger electrolyte resistance in S13 is caused by the larger electrode gap as explained in Table S1 and associated discussion in the Supporting Information. In short, for planar supercapacitors the electrolyte resistance ($R_s$) can be expressed

Table 1. Geometries Used in Electrode and Electrolyte Design Studies

<table>
<thead>
<tr>
<th>geometries $^a$</th>
<th>study 1</th>
<th>study 2</th>
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<tr>
<td>width, $W$ (cm)</td>
<td>S11 0.2</td>
<td>S12 0.2</td>
</tr>
<tr>
<td>length, $L$ (cm)</td>
<td>S21 0.15</td>
<td>S12 0.2</td>
</tr>
<tr>
<td>gap distance, $D$ (cm)</td>
<td>S11 0.1</td>
<td>S12 0.2</td>
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</table>

$^a$Refer to Scheme 1a for how width ($W$), length ($L$), and gap distance ($D$) are defined.

Figure 3. (a) Galvanostatic charge/discharge at 1.32 A cm$^{-3}$ for cells in study 1, and calculated specific cell capacitances. (b) Nyquist plot of study 1 supercapacitors from potentiostat EIS with inserted zoom-in at high-frequency range, and table of ESR values. (c) Galvanostatic charge/discharge at 1.32 A cm$^{-3}$ for cells in study 2, and calculated specific cell capacitances. (d) Nyquist plot of study 2 supercapacitors from potentiostat EIS with inserted zoom-in at high-frequency range, and table of ESR values. (e) Five thousand cycles GCD of cell S22 at 1.32 A cm$^{-3}$. (f) Ragone plot of cell S22 compared with recent relevant work.
as $R_s = \rho \kappa$, where $\rho$ is the resistivity of the electrolyte and $\kappa$ is the cell constant. As the gap distance increases, the cell constant also increases, leading to higher electrolyte resistance $R_s$ and degraded capacitive performance. From this study it can be concluded that reducing the electrode gap distance can enhance the capacitance of the as-printed supercapacitor. However, the printing resolution of DIW with the developed CNT ink limits how small the gap distance can be. Printing an extra small gap distance beyond the resolution of DIW may lead to partial overlapping of two adjacent electrodes and the shorting of the two electrodes, which results in a completely malfunctioned device.

In the second study, the electrode–gap distance (“D” in Scheme 1a) and the height of the electrode pattern (“L” in Scheme 1a) are kept the same but the width of the electrode pattern (“W” in Scheme 1a) is changed from 1.5 to 2 and 2.5 mm, hereafter denoted as S21, S12, and S22 respectively. The exact geometry of the cells can be found in Table 1. The volume of SW-CNT ink is adjusted so that the density and thickness of SW-CNT is kept the same for S21, S12, and S22. Specifically, the GCD curves of S21, S12, and S22 at varied current densities. (f) Cyclic performance of 3-cell supercapacitor at a constant 5.06 A cm$^{-3}$ current density GCD.

<table>
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<tr>
<th>active material</th>
<th>electrolyte</th>
<th>technique</th>
<th>capacitance</th>
<th>energy density</th>
<th>power density</th>
<th>year</th>
<th>ref</th>
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<td>graphene</td>
<td>LiCl/PVA</td>
<td>laser process</td>
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<td>0.98 mWh cm$^{-3}$</td>
<td>0.3 W cm$^{-3}$</td>
<td>2016</td>
<td>19</td>
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<td>carbon onion</td>
<td>Et$_4$NBF$_4$/anhydrous propylene carbonate</td>
<td>laser process</td>
<td>1.3 F cm$^{-3}$</td>
<td>1.5 mWh cm$^{-3}$</td>
<td>35 W cm$^{-3}$</td>
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<td>graphene</td>
<td>H$_2$PO$_4$/PVA</td>
<td>inkjet printing</td>
<td>99 $\mu$F cm$^{-2}$</td>
<td>N/A</td>
<td>N/A</td>
<td>2017</td>
<td>37</td>
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<td>MEMS</td>
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<td>495 W cm$^{-3}$</td>
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<td>0.3 W cm$^{-3}$</td>
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<td>DIW</td>
<td>15.34 F cm$^{-3}$</td>
<td>1.2 mWh cm$^{-3}$</td>
<td>11.8 W cm$^{-3}$</td>
<td>2017</td>
<td>this work</td>
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Figure 4. (a) Optical picture of the as-printed 3-cell supercapacitor array; 160° bended supercapacitor array. (b) CV comparison of bent and flat cells. (c) CV profiles of 3-cell supercapacitor at different scan rates. (d) Calculated specific capacitance at different current densities. (e) Galvanostatic charge/discharge of 3-cell supercapacitor at varied current densities. (f) Cyclic performance of 3-cell supercapacitor at a constant 5.06 A cm$^{-3}$ current density GCD.
and 1 V at a current density of 1.32 A cm\(^{-3}\) and the calculated specific capacitance are shown in Figure 3c. It is found that the specific capacitance increases as the electrode width increases from S21 (7.15 F cm\(^{-3}\)) to S22 (15.34 F cm\(^{-3}\)). The Nyquist plots in Figure 3d show that the ESR decreases as the electrode width increases. Different from study 1, this change in ESR in study 2 is caused by the changes of both the electrode resistance and the electrolyte resistance as the electrode width changes. Obviously, wider electrode patterns can lower the electrode resistance and enhance the capacitance of the printed supercapacitors. In addition, increasing electrode width also leads to the decrease of electrolyte resistance for planar supercapacitors,\(^{18}\) according to the analysis presented in Table S1 and the associated discussion in the Supporting Information. It should be noted that for optimal supercapacitor performance, the electrode width should not be arbitrarily large. As shown in the simulated electric field near the planar electrodes in Figure S6, the electric field becomes weaker at locations away from the edge of the electrodes and same as the electrostatic force driving the formation of electric double layers (EDL). Thus, the active materials away from the electrode edges do not contribute as much EDL capacitance as those near the edges and the overall specific capacitance is decreased. With relatively narrow width, the benefits of increasing electrode width (i.e., lower electrode and electrolyte resistance) will overshadow the effect of nonuniform electric field and the specific capacitance will increase with electrode width (as shown in our study 2 results). It is expected that with an extra wide electrode, the effect of nonuniform electric field will eventually become more significant than the benefits of the wide electrode and the specific capacitance will start decreasing with the increase of electrode width.

Cell S22 with the highest specific capacitance is used for further studies. A cyclic stability study (Figure 3c) showed that S22 can maintain ~96.5% of its initial capacitance after 5000 cycles of GCD at 1.32 A cm\(^{-3}\). The energy density and power density of cell S22 are calculated using eqs 5 and 6 at varied current density from 0.02 to 20 A cm\(^{-3}\) and compared with recent planar carbon-based electric double-layer capacitors (EDLCs) in Figure 3f. The S22 cell has a high energy density of 1.18 mWh cm\(^{-3}\) at a power density of 11.8 W cm\(^{-3}\). The energy and power performances of the as-printed fully packaged supercapacitor are among the best in recently reported planar electric double-layer capacitors and even higher than some graphene-based supercapacitors (Table 2), which involved a high-cost fabrication process such as lithography patterning and high-temperature annealing.

An array of three supercapacitors connected in series is printed. A picture of the printed supercapacitor array is shown in Figure 4a. The SW-CNT electrode can be clearly seen in the picture, whereas the transparent electrolyte and silicone seal cannot be discerned in the picture. The supercapacitor is connected to electrical wires by silver paste. Since all components including the electrodes, the electrolyte, the substrate, and the seal are flexible, the as-printed supercapacitor showed excellent flexibility. A CV test is performed when the cell is bent, and the bent cell retains similar performance compared to unbent flat cell (Figure 4b), which proves the mechanical flexibility of the cell.

A series of CV tests is performed with a potential window of 0–3 V and varied scan rates of 10, 20, 30, 40, 50, and 100 mV s\(^{-1}\). The CV curves (Figure 4c) all have rectangle shapes, which confirms the printed device is a pure electric double-layer capacitor. The GCD curves measured at current densities 0.72, 2.17, 3.61, 5.06, 6.50, and 10.84 A cm\(^{-3}\) are shown in Figure 4e. The charge and discharge curves are highly symmetric. Specific capacitances of the supercapacitor array are calculated by eq 1 and plotted at different current densities (Figure 4d). The specific capacitance only drops by 34.9% when current density increases from 0.72 to 10.84 A cm\(^{-3}\), demonstrating good rate capability.

The cyclic stability is investigated by performing GCD at a current density of 5.06 A cm\(^{-3}\) (Figure 4f). There is no significant decrease of cell capacitance (~2.42 F cm\(^{-3}\)), and the capacitance is retained 97.7% after 1000 cycles of charge/discharge. It is noted that the capacitance increases after a few cycles at the beginning of the cyclic test. This increase can be attributed to the increase of SW-CNT surface area induced by the surface activation effect of the charge/discharge process.

4. CONCLUSION

In this study, we report for the first time the direct ink writing (DIW) of fully packaged supercapacitors on flexible polymer substrate. Different supercapacitor designs have been implemented by the DIW, and electrochemical study results show that smaller gap spacing between two electrodes and a wider electrode facilitate the fast ion transport and electron transport, respectively, which improve the overall cell performance. A multicell supercapacitor array with three identical cells connected in series is printed to demonstrate the scalability of the developed technology. The direct-written fully packaged SW-CNT supercapacitor has comparable energy and power performances to recently reported planar carbon-based supercapacitors fabricated by other printing or nonprinting methods. The developed ink formula and direct ink writing process enable the mask-free, transfer-free, and alignment-free printing with precise and repeatable control on the spatial distribution of different functional materials. It paves the way for rapid, low-cost, and scalable manufacturing of high-performance fully packaged supercapacitors for practical applications.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.7b06804.

Viscosity of each printing material at shear rate 1–100 S\(^{-1}\) at 25 °C temperature, cross-section SEM of SW-CNT electrode in S1 and S2 study, C1s spectrum of raw SW-CNT and the as-printed electrodes without and with acid treatment, O1s spectrum of raw SW-CNT and the as-printed electrodes without and with acid treatment, N1s spectrum of raw SW-CNT and the as-printed electrodes without and with acid treatment, finite element simulation of equipotential lines near a coplanar electrode, calculated cell (k) constant for different electrode geometries in study 1 and study 2, discussion of results (PDF)

**AUTHOR INFORMATION**

*Corresponding Authors*

E-mail: yayuepan@uic.edu.
E-mail: shanhu@iastate.edu.

**ORCID**

Shan Hu: 0000-0003-4842-5748

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