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Abstract
There is ever increasing demand for flexible energy storage devices due to the development of wearable electronics and other small electronic devices. The electrode flexibility is best provided by a special set of nanomaterials, but the required methodology typically consists of multiple steps and are designed just for the specific materials. Here, a facile and scalable method of making flexible and mechanically robust planar supercapacitors with interdigital electrode structure made of commercial carbon nanomaterials and silver nanowires is presented. The capacitor structure is achieved with vacuum filtration through a micropatterned contact mask and finished with simple laser processing steps. A maximum specific capacitance of 4 F cm\(^{-3}\) was measured with cyclic voltammetry at scan rate of 5 mV s\(^{-1}\). The reliability and charge transfer properties of devices were further investigated with galvanostatic charge-discharge measurements and electrochemical impedance spectroscopy, respectively. Furthermore, mechanical bending tests confirmed the devices have excellent mechanical integrity, and the deformations have no adverse effects on the electrochemical charge-discharge behavior and stability.

Keywords: supercapacitors, vacuum filtration, laser processing, carbon nanotubes, reduced graphene oxide, silver nanowires

Supplementary material for this article is available online

(Some figures may appear in color only in the online journal)

1. Introduction

Due to the increasing popularity and market needs of portable electronics, wearable devices and Internet of Things (IoT), there is growing demand for the development of new advanced flexible energy storage devices [1, 2]. Carbon nanomaterials have numerous attractive properties that have been utilized in electrical applications [3]. Their electrical
conductivity, high surface area, chemical stability, light weight as well as mechanical flexibility make them the material of choice in electrodes for electrochemical energy storage, especially for supercapacitors with high power density and excellent cycle stability [4–6]. Among the carbon nanomaterials graphene/reduced graphene oxide (RGO) [7–34], carbon nanotubes (CNTs) [21–27, 35–46], carbon black [7, 8, 47], activated carbon [20, 25, 35, 48], carbon nanofibers [35, 49–51] as well as biomass derived carbon [51–53] have been the most prominent materials for supercapacitor electrodes. Moreover, silver in the form of nanowires [14, 35] and nanoparticles [54] have also been utilized in in the electrode material enhancing the performance of the electrode due to its superior intrinsic electrical conductivity [55].

As the field of flexible electronics grows rapidly, research on flexible supercapacitors have attracted significant attention [14–20, 23–37, 43–48]. Supercapacitors are usually assembled with a stacked configuration, where the two electrodes are wetted by a liquid electrolyte, separated with a spacer and connected by metal current collectors. All-solid-state flexible supercapacitors with a planar interdigital structure however offer a series of advantages, such as low volume, high stability and a control of the electrode design which has a strong influence on the device performance [14]. Most importantly, the devices can be placed in flexible and wearable systems as well as in other portable autonomous electrical circuits that can serve as back-up for uninterruptable power systems, small battery replacements and can be also applied in energy harvester devices to store the scavenged energy.

Vacuum filtration has long been used to make carbon-based films that are then utilized as electrodes. The process itself is facile and quick to produce highly porous carbon films that are either supported by the filter membrane [12–15, 27–29, 37–39] or completely freestanding after removal [7, 20, 23–25, 48]. The films can also be patterned into an interdigital structure by etching the filtered thin film [14] or by multi-step photo-lithography process [37]. Other reported methods of making carbon-based interdigital electrodes on flexible substrates have been inkjet printing [18, 35], mask-assisted spray coating [18] layer-by-layer assembly [22, 36], film transfer [17, 27, 29–31] and 3D-printing [46]. Moreover, laser processing is one of the most promising method to reduce process steps in microelectronics patterning as it does not require time consuming photolithography steps or vacuum equipment and is capable to process large areas. Laser has been used to process carbonaceous materials, such as graphene and graphene oxide [9–11, 16, 26, 31–33] as well to sinter nanostructured metals [54] and also to enable CNT growth on metal substrates [56].

In this research, we are demonstrating a fast, scalable and straightforward method, in which planar supercapacitor electrodes with interdigital structure are prepared by vacuum filtration through micropatterned contact masks. Though a similar approach have been reported to make interdigital electrodes [27, 29], here no electrode film transfer was needed and a thin layer of silver nanowires (Ag-NWs) is deposited on the top of the carbon film, as a current collector layer to significantly lower the electrode resistance. Carbon nanotubes and RGO were chosen as electrode material due to their well-known properties such as excellent electrical conductivity, large surface and high stability under mechanical stress area which is why they are used in numerous electrochemical energy storage applications [5, 6, 57–59]. The as obtained nanocarbon-Ag capacitor structure is then further processed with a quick laser cleaning of any possible shorts, which is then followed by laser-assisted sintering the Ag-NWs to optimize the conductivity of the collector. By applying a PVA-based electrolyte we obtain solid-state supercapacitor devices that are mechanically robust and suitable for energy storage in wearable and other small electronics demanding flexible power sources.

2. Methods

100 mg of graphene oxide (Sigma Aldrich prod. No. 796034) was first mixed in 200 ml of de-ionized (DI) water. 2 g of NaBH₄ was then added to the solutions and stirred overnight with a magnetic stirrer at room temperature. The RGO was then filtered on PTFE filter paper (1 µm pore size), rinsed with DI-water seven times and dried at 70 °C overnight. Suspensions of multi-walled carbon nanotube (MWCNT) (Sigma Aldrich prod. No. 773840, 50 mg/L), single wall carbon nanotube (SWCNT) (Sigma Aldrich prod. No. 519308, 300 mg L⁻¹), RGO (300 mg L⁻¹) and Ag-NW (Nanostructured & Amorphous Materials Inc. prod. No #0475NW1 2.5 g l⁻¹) were done in isopropyl alcohol (IPA) and sonicated for 30 min. The SWCNT/RGO@IPA suspension were then made by mixing the two suspensions using a 1:1 ratio.

A design of four interdigital capacitor structures of 500 µm line width and 200 µm spacing (total electrode area of 0.49 cm²) was cut on biaxially-oriented polyethylene terephthalate (BoPET, thickness of 50 µm, 47 mm diameter) filter mask by a laser (LPKF Protolaser U3, λ = 355 nm). The filter mask is then placed on hydrophilic polyvinylidene fluoride (PVDF, Durapore Millipore GVWP4700, 47 mm diameter, 22 µm pore size) filter membrane for the vacuum filtering steps. 1 ml of MWCNT@IPA is first applied as a primer layer, which is then followed by 2 ml of SWCNT/RGO@IPA. The filtration assembly is then rinsed with IPA after which 220 µl of Ag-NW@IPA is applied as current collectors and the structures are left to dry overnight. All suspensions are sonicated for 3 min before applications. The capacitor structures are then trimmed (PRG = 0.9 W, f = 200 kHz) and sintered (Pavg = 2 W, f = 120 kHz, off-focus = 12 mm) by a UV-laser (LPKF Protolaser U3, λ = 355 nm). The filter membranes were then cut into separate capacitors. The H₃PO₄–PVA gel electrolyte was prepared by mixing PVA (Fluka prod. No. 10852, Mw ~ 61 000), and H₃PO₄ (SAFC, 85 wt.%) in DI-water in 1:2:10 ratio. The mixture was kept at 80 °C under stirring until a clear solution was obtained. The electrolyte was then cooled down to room temperature, applied on top of the capacitor structures and let to solidify overnight. The ionogel electrolyte was prepared by mixing PVA (Fluka prod. No. 10852, Mw ~ 61 000), 1H3PO4 (SAFC, 85 wt.%) in DI-water in 1:2:10 ratio. The mixture was kept at 80 °C under stirring until a clear solution was obtained. The electrolyte was then cooled down to room temperature, applied on top of the capacitor structures and let to solidify overnight. The ionogel electrolyte was prepared by mixing 110 mg of fumed silica and 3 ml of 1-ethyl-3-methylimidazolium tetrafluoroborate (EMIM-BF₄, Sigma Aldrich prod. No. 00768). The mixture was magnetically stirred for 3 h in a nitrogen atmosphere before applying to the capacitor structures.
The electrochemical measurements the capacitors were connected to a potentiostat-galvanostat (Princeton Applied Research VersaSTAT 3) with probes. The electrochemical performance of the capacitors was assessed by cyclic voltammetry (CV), galvanostatic charge-discharge (GDC) and electrochemical impedance spectroscopy (EIS) measurements. The specific volumetric capacitance $C_v$ was calculated from the averages of the integrated current-time hysteresis curves normalized by the electrode volume:

$$C_v = \frac{\int I(U) dV}{\Delta U / \Delta t \cdot v}$$  \hspace{1cm} (1)

where $I(U)$ is the charging current, $dU/dt$ is the scan rate, $\Delta U$ is the used voltage range between $U_1$ and $U_2$ and $v$ is the volume of the device. The capacitance was calculated from charge-discharge measurements with:

$$C_v = \frac{I}{\Delta U / \Delta t \cdot v}$$  \hspace{1cm} (2)

where $I$ is the used current, $\Delta U$ the voltage window, $\Delta t$ is the discharge time and $v$ is the volume of the capacitor. Energy density values of the on-chip devices are obtained from:

$$E_d = \frac{CL^2}{2v}$$  \hspace{1cm} (3)

where $C$ is the calculated capacitance from charge-discharge measurements, $U$ is the used voltage range and $v$ is the volume of the device. The power density is calculated using the equation:

$$P_d = \frac{E_d}{\Delta t}$$  \hspace{1cm} (4)

where $E_d$ is the calculated energy density from charge-discharge measurements and $\Delta t$ is the discharge time.

X-ray micro-computed tomography analysis was done with Bruker Skyscan 2211 instrument (Projected images were reconstructed using CtVox software). The electrical resistivity measurements were done with a multimeter (Fluke 289 true RMS multimeter) The structure and the materials were assessed with field emission scanning electron microscope (FESEM, Zeiss Ultra Plus and Zeiss Sigma, light microscope (Olympus BX51 equipped with Colorview imaging system), Raman spectroscopy (Horiba Jobin–Yvon LabRAM HR, $\lambda = 488$ nm) and x-ray photoelectron spectroscopy (XPS, Thermo Fisher Scientific Escalab 250 XI system with Al K$\alpha$ x-ray source, 1486.6 eV, data evaluation using Avantage software).

### 3. Results and discussion

The interdigital capacitor structures were prepared using a standard filtration setup (figure 1(a)) combined with a filtration mask. In the process steps (figure 1(b)), the filter mask with the capacitor patterns was first placed on the top of a PVDF filter membrane. The carbon nanomaterial, consisting of SWCNTs and RGO dispersed in IPA, was then filtered through the mask and membrane thus depositing on the open areas of the filtration mask. Using a mixture of CNTs and RGO prevents the graphene sheets from stacking, thus enhancing the effective surface area and performance in a supercapacitor [60]. Next the Ag-NWs were deposited on the top of the carbon nanomaterials to function as a current collector layer. Though the filtration process is simple, quick and effective, the carbon nanomaterial frequently deposited underneath the filter mask because of a slight underflow of the dispersion. This issue was tackled by depositing a thin layer of MWCNTs as a primer layer. The MWCNTs had a much poorer dispersion in IPA which prevented them from flowing underneath the filtration mask.

Despite the deposited Ag-NWs, the resistivity of the electrodes was still too high ($R_s > 100 \Omega/□$) to function properly in a supercapacitor, which was likely caused by the imperfect junctions between the crossing nanowires in the network. To improve the conductivity of the Ag-NW current collector layer, we applied a sintering strategy demonstrated earlier [35, 52]. Here, we used a scanned pulsed laser beam set to a strong off-focus (to increase the effective beam spot size and to avoid ablation of the material) to heat and anneal the crystalline Ag-NWs, which underwent partial melting and fused together at the junctions (figures 2(a) and (b)) thus eliminating the contact resistance and consequently lowering the resistivity of the Ag-NW network (figure 2(c)). In the first laser-assisted sintering cycle, the resistivity decreased over two orders of magnitude from $R_s > 100 \Omega/□$ to $R_s > 0.3 \Omega/□$ after which the subsequent sintering cycles resulted in only very moderate further drop of resistivity. In an attempt to optimize further the conductivity of the collectors, we tested whether thicker Ag-NW networks could have any advantage. The results showed that the resistivity did not significantly improve with higher Ag-NW loadings showing only linear relationship in the analyzed surface loading range between 0.5 mg cm$^{-2}$ and 1.25 mg cm$^{-2}$ with corresponding sheet resistances of 0.28 $\Omega/□$ and 0.12 $\Omega/□$. Based on the results above, in the further part of the work, we applied 100 cycles for laser-assisted sintering of Ag-NW networks with 1.0 mg cm$^{-2}$ surface loading to be used for the electrodes of supercapacitors.

Though the amount of material deposited underneath the filtration mask was significantly reduced by the MWCNT primer layer, there still was an occasional occurrence of carbon material depositing on the interdigital electrode spacings, which was shorting the capacitor structure (figure S1 (available online at stacks.iop.org/NANO31/495403/mmedia)). The electrode spacings were therefore always cleaned by a laser trimming process using a pulsed UV process laser with a low power adding only one extra phase in the lasering process. As the carbon material effectively absorbs the UV radiation, the trimming process can easily damage the membrane substrate if the power is too high. This was especially prominent when the carbon layer was thick and well absorbing so that the generated heat could partially degrade the PVDF filter underneath [61]. Due to this, and to the apparent material loss, it was not feasible to just filter a uniform layer on carbon material and pattern it by using only laser processing. In this study, the total electrode area of one capacitor was 0.49 cm$^2$ and a total of
Figure 1. (a) Filtration setup. (b) Illustration of the capacitor assembly steps. (i) Carbon nanomaterial deposition on filter membrane through a filtration mask, and (ii) subsequent silver nanowire deposition. (iii) Sintering and trimming of the interdigital electrode pattern by a UV-laser. (iv) Applying PVA gel electrolyte on the electrodes.

Figure 2. (a) SEM image of the pristine silver nanowires on the electrode. (b) SEM image of the silver nanowires after 100 sintering cycles showing the necks of sintered joints at contacts. (c) Effect of laser sintering on the electrode resistance with different Ag-NW loadings. (d) High magnification SEM image of the electrode material taken from the cross-section of the electrode. The inset panel shows a lower magnification cross-section (scale bar is 5 µm). (e) Optical camera image illustrating the size and the structure of the supercapacitor and (f) flexibility of the device.

650 µg of carbon material was deposited as electrode material in one capacitor resulting in a film thickness of ~10 µm (figure 2(d)) which is much thicker than usually reported. However, when investigating the quality of the used commercial carbon nanomaterials, Raman spectroscopy indicate significant concentration of defects in the lattices of both RGO and MWCNT materials [62], which likely reduced the capacity performance (figure S2). XPS analysis of the as-prepared electrode confirm Ag-NWs to be metallic and also show large concentration of carbon-oxygen bonds within the material [63], which is expected (figure S3).

One of the main advantages of this method is that the electrode thickness can be much greater than what is typically reported in planar interdigital capacitor structures made with etching [14, 34], aerosol spraying [18, 33], layer-by-layer assembly [37] or inkjet printing [18, 35]. As a last step the H3PO4/PVA gel electrolyte was applied on the capacitors and left to dry overnight providing a flexible and robust
Figure 3. X-ray tomography images of the H$_3$PO$_4$-PVA electrolyte covered electrode under bending. The scalebar is 1 mm.

Figure 4. (a) Cyclic voltammetry curves of the supercapacitor (b) Corresponding calculated specific capacitances with different scan rates. (c) Charge-discharge curves with different current densities. (d) Ragone plot of energy and power densities.

The electrochemical behavior of the supercapacitors was measured in the voltage window between 0 V and 0.5 V (calculation methods and equations are provided in the supporting information). By CV with scanning rates from 5 mV s$^{-1}$ to 1000 mV s$^{-1}$. The CV curves have rounded shape even at slower scan rates (figure 4(a)) indicating ohmic losses in the electrode structure. The highest volumetric capacitance 4 F cm$^{-3}$ was measured at 5 mV s$^{-1}$, and as typical for supercapacitors, decreased at higher scan rates. (figure 4(b)). The GDC curves follow the typical triangular shape (figure 4(c)) and show capacitances very close to those measured with CV (3.7 F cm$^{-3}$ at the lowest current density of 40 mA cm$^{-2}$). Similar to the CV analysis, the measured capacitances dropped at higher current densities due to the increasing voltage drop at the start of the discharge step. The calculated energy and power densities are illustrated in Ragone plot (figure 4(d)). The highest measured energy and power densities were 130 $\mu$Wh cm$^{-3}$ and 2.7 W cm$^{-3}$ respectively as the energy and power performance was significantly limited by the narrow bias window of 0.5 V. The performance of the capacitors is however still well comparable with the state-of-the-art in flexible planar supercapacitors (Supplementary table S1). The methodology reported here have however significant advantages as it is facile, scalable and could be used with multiple different electrode materials.

When using a 0.8 V potential window it was noted that the performance of the capacitor dropped to 85% after 1000 measurement cycles and to 42% after 5000 cycles (figure 5(a)). After 5000 cycles the changes were also clearly visible in the positive electrode that turned dark indicating that the metallic silver have oxidized. Experiments with EMIM-BF$_4$-based gelated ionic liquid (ionogel) that allow for a higher potential window (up to 2 V) enabled higher currents in CV measurements due to the oxidation reaction of the Ag-NWs (figure S4) therefore leading to an even more accelerated performance degradation. This founding was surprising as the Ag-NWs have been previously reported as an electrode material with both H$_3$PO$_4$/PVA-based electrolytes [64] and with ionogels [35], though it has to be noted that Ag/Ag$^+$ has a redox potential of 0.799 V vs normal hydrogen electrode which also
Figure 5. (a) Retention of the capacitance with different with and without silver nanowires in 0.5 V and 0.8 V potential ranges. (b) Nyquist diagram according to EIS measurement on the supercapacitor. (c) Cyclic voltammetry curves under different bending conditions. (d) Retention of capacitance as a function of repeated bending cycles.

decreases according to the size of the nanoparticle [65]. One possible explanation could be that the role of the Ag-NWs was not as significant as in this research as a current collector and therefore the degradation did not significantly affect the performance. Without using the Ag-NWs the capacitive performance was still 89% after 10 000 cycles indicating that there still were some irreversible chemical reactions occurring on the carbon material surfaces likely due to impurities. The capacitor performance without the Ag-NWs as a current collector was however rather poor as the series resistance was almost two orders of magnitude higher (figure S5). With 0.5 V potential range the capacitive performance dropped initially to 96% after the first 1000 cycles after which the performance slowly improved and was 106% of the original performance after 10 000 cycles indicating a good material reliability when using a 0.5 V potential range. No visible aggregation of the electrode material was seen after the retention tests. From EIS measurements the series resistance was evaluated to be ~33 Ω (figure 5(b)). The Nyquist diagram does not show a clear half-circle indicating low charge transfer resistance. As the resistance of the electrode after Ag-NW sintering was ~0.1 Ω, the resistivity is likely due to the carbon materials used in this research. This was also indicated by the lower series resistance of capacitors made with thinner electrode material layers.

Mechanical bending tests conjunction with CV measurements were conducted to assess the reliability of the flexible capacitors. The capacitors exhibited excellent stability. Even with 0.7 mm bending radius, the shape of the CV curve does not change, and the overall capacitance is only 1% lower (figure 5(c)). The same behavior is also visible with retention tests that show the performance was still 98% of the original value after 1000 bending cycles with ~1 mm bending radius (figure 5(d)) indicating excellent overall stability under mechanical stress as expected from the x-ray topology analysis.

4. Summary and conclusions

Here we have demonstrated a fast and easily up and down scalable method of producing interdigital supercapacitor structures by straightforward filtering steps using commercially available materials such as SWCNTs and MWCNTs, RGO and Ag-NWs on PVDF filter substrates. We have shown that laser-assisted sintering of Ag-NWs deposited on nanostructured carbon-based interdigital porous electrode films allow for the fabrication of highly conductive current collectors of high performance flexible planar supercapacitors. Using H₃PO₄/PVA electrolytes, volumetric capacitances of up to 4 F cm⁻³ were achieved. According to our study, the suitability of Ag-NWs as an electrode material in supercapacitors is limited to a small electrochemical window due to the oxidation of silver above 0.5 V.

While our work shows a simple method to achieving planar flexible capacitor devices, we envisage a number of possibilities that could improve the overall performance of the devices. For instance, the collector may be optimized further by replacing silver with gold (or with gold coated core–shell nanowires) to expand the voltage window. Another possible method to decrease the series resistance could be to use the filtering mask also as a shadow mask for physical vapor deposition of a metal thin film for the current collector. This would of course demand very careful work not to move the mask between the steps. Moreover, the performance could be also significantly improved by utilizing nanostructured carbons with higher specific surface area and conductivity than those applied in this study. It shall be also noted here, that the capacitance of the devices may be increased further significantly by adding pseudocapacitive materials such as oxides of manganese or ruthenium on the electrodes [40, 66, 67].

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