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Ultrahigh-power micrometre-sized supercapacitors based on onion-like carbon

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Electrochemical capacitors, also called supercapacitors, store energy in two closely spaced layers with opposing charges, and are used to power hybrid electric vehicles, portable electronic equipment and other devices1. By offering fast charging and discharging rates, and the ability to sustain millions of cycles2–4, electrochemical capacitors bridge the gap between batteries, which offer high energy densities but are slow, and conventional electrolytic capacitors, which are fast but have low energy densities. Here, we demonstrate microsupercapacitors with powers per volume that are comparable to electrolytic capacitors, capacitances that are four orders of magnitude higher, and energies per volume that are an order of magnitude higher. We also measured discharge rates of up to 200 V s⁻¹, which is three orders of magnitude higher than conventional supercapacitors. The microsupercapacitors are produced by the electrophoretic deposition of a several-micrometre-thick layer of nanostructured carbon onions5–7 with diameters of 6–7 nm. Integration of these nanoparticles in a microdevice with a high surface-to-volume ratio, without the use of organic binders and polymer separators, improves performance because of the ease with which ions can access the active material. Increasing the energy density and charge rates of supercapacitors will enable them to compete with batteries and conventional electrolytic capacitors in a number of applications.

The recent boom in multifunction portable electronic equipment and the increasing need for wireless sensor networks for the development of smart environments has raised the problem of developing sufficiently compact and/or flexible energy storage. Designing efficient, miniaturized energy-storage devices that can achieve high energy delivery or harvesting at high discharge rates with a lifetime that matches or exceeds that of the machine being powered remains a challenge1. Integrating the storage element as close as possible to the electronic circuit (directly on a chip) is another challenge. Because electrochemical energy storage in batteries occurs by means of volumetric reactions8, the charge discharge rate and specific power of the best batteries (lithium-ion) are limited by the rate of solid-state diffusion. Also, the redox reactions and expansion–contraction of the active material limit battery lifetime to just hundreds or thousands of cycles. These problems can only be partially resolved by using nanostructured materials10,11. The properties of thin-film batteries, despite their excellent performance per unit volume, drop dramatically in the micrometre range12.

Electrochemical capacitors (ECs) store energy using an accumulation of ions of opposite charge in a double layer at electrochemically stable, high specific surface area electrodes. The high surface-to-volume ratio of the active material leads to the high energy and power densities of ECs; this is further enhanced in microsupercapacitors13,14. Porous activated, templated15 and carbide-derived carbons16, multi- and single-walled carbon nanotubes17 and multilayer graphene18 have been used as the electrode materials in supercapacitors. However, very few studies have been performed using spherical carbon nanoparticles19–22, which do not have the narrow pores that in activated carbons or nanotubes may cause the transport of ions to be the rate-controlling factor limiting the charge/discharge rate. This Letter reports the electrochemical performance of onion-like carbon (OLC) electrodes assembled in a micromanufactured device, which was able to cycle at a scan rate of 200 V s⁻¹.

OLCs are quasi-spherical nanoparticles consisting of concentric graphitic shells23. Although there are many methods by which OLCs may be produced, the annealing of detonation nanodiamond powders (Supplementary Fig. S1a–d) is inexpensive and the only widely used method that allows the synthesis of large amounts of OLC24. In an ideal case, they could be considered as multishell giant fullerenes, but real particles have discontinuous and defective shells when synthesized at temperatures below 1,800 °C (as shown in Supplementary Fig. S1d) or polygonized shells when higher temperatures or longer times are used.

Carbon onions offer a moderate specific surface area (~500 m² g⁻¹; Supplementary Table S1) compared to that of activated carbons, but this surface is fully accessible to ion adsorption (Fig. 1a) because there is no porous network inside the particles. According to OLCs have demonstrated modest gravimetric capacitance in previous studies, at about one-third that of activated carbons24. Based on these results, we anticipated that the accessible external surface area of the carbon onions would be appealing in the design of thin electrodes for microsystems, where the limited volume available for device integration onto chips drives the technology.

OLC particles were produced by annealing nanodiamond powder at 1,800 °C (Fig. 1b) then deposited from colloidal suspensions using an electrophoretic deposition technique (EPD) onto interdigital gold current collectors patterned on silicon wafers. This method allows electrodes to be prepared without the use of an organic binder (Fig. 1c; see also Supplementary Table S2). An adherent layer of OLC was obtained on the gold current collectors, with a well-defined pattern and no short circuit between the electrodes (Fig. 1d). The microdevice was constructed with 16 interdigital electrodes with thicknesses of 7 µm (eight electrodes per polarity), as shown in Fig. 1e.

Cyclic voltammograms (CVs) were recorded at scan rates from 1 to 200 V s⁻¹ to test the power capability of the microsystem. A reproducible and stable capacitive behaviour (up to 10,000

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cycles; Supplementary Fig. S2) was obtained for the microdevice over a 3 V potential window in a 1 M solution of tetraethylammonium tetrafluoroborate in propylene carbonate, with a linear dependence of the discharge current on the scan rate and low resistive contributions up to 100 V s\(^{-1}\) (Fig. 2). This scan rate is more than two orders of magnitude higher than any result reported with alternative devices\(^{13,29}\), including microdevices\(^{13,34}\) and microcavity electrodes\(^{26,28}\). Furthermore, the more interdigital electrodes per unit area, the more power is extracted from the microdevice. This is explained by a significant reduction in the mean ionic diffusion path between the two electrodes (Supplementary Fig. S3 and Table S2).

Figure 1 | Design of the interdigital microsupercapacitor with OLC electrodes. a, Cross-section of a charged zero-dimensional OLC (grey) capacitor, consisting of two layers of charges (blue and pink) forming the inner and outer spheres, respectively. b, Transmission electron microscopy image of a carbon onion produced at 1800 °C. Lattice spacing between the bent graphene layers in the onions is close to 0.35 nm. c, Schematic of the microdevice (25 mm\(^2\)). Two gold current collectors made of 16 interdigital fingers were deposited by evaporation on an oxidized silicon substrate and patterned using a conventional photolithography/etching process. Carbon onions (active material) were then deposited by electrophoretic deposition onto the gold current collectors. d, Optical image of the interdigital fingers with 100-µm spacing. e, Scanning electron microscopy image of the cross-section of the carbon onion electrode. A volumetric power density of 1 kW cm\(^{-3}\) was obtained with a deposited layer thickness in the micrometre range, not the nanometre range.

Figure 2 | Electrochemical characterizations of the microdevices. a, CVs obtained at different scan rates in a 1 M Li\(_2\)NBF\(_4\)/anhydrous propylene carbonate on a 16-interdigital electrochemical microcapacitor with a 7-µm-thick OLC deposit. A typical rectangular shape, as expected for double-layer capacitive materials, is observed at an ultrahigh scan rate over a 3 V potential window. b, Evolution of the discharge current versus scan rate. A linear dependence is obtained up to at least 100 V s\(^{-1}\) in the capacitive region, indicating an ultrahigh power ability for the microdevices.

The significant (7 µm) thickness of the active film (Fig. 1). The combination of carbon onions with a fully accessible surface area, with a binder-free deposition technique and a micro-interdigital device design led to this high power/energy performance.

To highlight the effect of the endohedral structure of carbon onions on cell performance, a 5-µm-thick high-surface-area activated carbon (AC) (1,700—1,800 m\(^2\) g\(^{-1}\), Kuraray Chemical Co.) was used as the active material in a microdevice made of 16 interdigital electrodes. The deposition technique and the cell assembly were kept the same. Figure 3a,b shows a comparison of the stack capacitance and energy of microdevices with AC and OLC. Despite a higher capacitance (9.0 F cm\(^{-3}\), compared to 1.3 F cm\(^{-3}\) for OLC) and a capacitive behaviour up to 1 V s\(^{-1}\) (Supplementary Fig. S4), the capacitance of microdevices with AC fell quickly at a higher scan rate, indicating a severe decrease in the instantaneous power of AC relative to OLC-based microdevices because of the limitation of ion transfer in the inner porous network of the AC. Both microdevices exhibited a low equivalent series resistance (ESR;
Figure 3 | Comparison of microsupercapacitors and other energy storage devices. a. Evolution of the stack capacitance versus scan rate. Carbon onion microsupercapacitors can sustain very high scan rates, like electrolytic capacitors. The stack capacitance is, however, four orders of magnitude higher than that of the electrolytic capacitors. b. Evolution of the volumetric energy of different energy-storage devices. c, d. Evolution of the real and imaginary part (C' and C") of the stack capacitance of a 16-interdigital electrochemical microcapacitor based on OLC (c) and AC (d). An extremely low relaxation time constant $\tau_0$ (26 ms) was obtained for the OLC, revealing fast accessibility of the ions for electroosmosis.

Figure 4 | Comparison, in a Ragone plot, of the specific energy and power density (per cm$^3$ of stack) of typical electrolytic capacitors, supercapacitors and batteries with the microdevices. All the devices (macro and micro) were tested under the same dynamic conditions. A very high energy density was obtained with the AC-based microsupercapacitor, whereas ultrahigh power density was obtained with the OLC-based microsupercapacitor.

Supplementary Fig. S5), but because the entire outer surface of the OLC is fully accessible to ion adsorption/desorption, it is characterized by an extremely small characteristic relaxation time constant $\tau_0$ of 26 ms ($\tau_0$ being the minimum time needed to discharge all the energy from the device with an efficiency of greater than 50% (ref. 30); Fig. 3c), which is much lower than that of the AC-based microdevice ($\tau_0 = 700$ ms; Fig. 3d) or OLC-based macroscopic devices ($\tau_0 > 1$ s)$^{34}$. It has been shown theoretically that the positive curvature causes the normalized capacitance to increase with decreasing particle size$^{31}$. Therefore, OLC has the potential for delivering high power and energy as one of the smallest endohedral particles available.

A comparison of the different energy storage devices designed for power microelectronics applications (a 500-μAh thin-film lithium battery, a 25-mF supercapacitor and an electrolytic capacitor of the same absolute capacitance), tested under the same dynamic conditions, is presented in the Ragone plot of Fig. 4. This shows the relationship between the volumetric energy density and power density of the stack (comprising the current collectors, the active material and the separator) for the energy storage devices tested. The specific power and specific energy were calculated by integrating the CVs at different rates (see Supplementary Information for
The electrolytic capacitors were able to perform at a very high scan rate, but had a stack capacitance that was four orders of magnitude lower than the microdevices, and a specific energy that was more than one order of magnitude lower. Meanwhile, the lithium battery and conventional supercapacitors could not provide the ultrastable discharge rate demonstrated by the microdevice. Electrochemical microcapacitors with AGs demonstrate an energy density of 1 x 10^-3 W cm^-2 (10 Wh l^-1 using conventional units). With OLC, a maximum power density is obtained that is 100 times higher, at close to 1 kW cm^-2 (1 MW l^-1). These therefore represent a new generation of devices, approaching the power of electrolytic capacitors, but having orders of magnitude larger energy density. To the best of our knowledge, there are no other devices available, including state-of-the-art microcapacitors that show the same performance characteristics.

This finding addresses the need for microscale energy storage in numerous areas where electrolytic capacitors cannot provide sufficient volumetric energy density, such as nomad electronics, wireless sensor networks, biomedical implants, active radiofrequency identification (RFID) tags and embedded microsensors. These applications can be extended to larger devices by scaling up the electrode surface and using high-resolution ink-jet, transfer printing and other high-throughput techniques that are suitable for mass production. Further improvements in the configuration of the patterned electrodes (for example, decreasing the space between the electrodes down to the nanoscale), in the material properties (for example, decreasing the onion size), the density and homogeneity of the deposited electrode, and viscosity/conductivity of electrolytes are expected to allow the design of high-energy supercapacitors with even higher power and energy characteristics.

References


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Author contributions

M.B. and D.P. conceived and designed the experiments for the elaboration of the electrochemical microcapacitor. D.P. established the EPD process. Y.G. was involved in material synthesis and characterization. V.M. carried out the simulation of OLC formation. D.P., P.H., P.L.T. and P.S. performed the electrochemical characterizations. D.F., M.B., P.S. and V.G. co-wrote the paper, and all authors discussed the results and commented on the manuscript.

Additional information

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