Reduced Graphene Oxide as a Monolithic Multifunctional Conductive Binder for Activated Carbon Supercapacitors


†Electrical Engineering Division, Department of Engineering, University of Cambridge, 9 J. J. Thomson Avenue, CB3 0FA Cambridge, U.K.
‡Faculty of Physics, University of Vienna, Boltzmanngasse 5, A-1090 Vienna, Austria
§Institute of Materials Chemistry, Vienna University of Technology (TU Wien), Getreidemarkt 9/165, A-1060 Vienna, Austria

Supporting Information

ABSTRACT: Using reduced graphene oxide (r-GO) as a multifunctional conductive binder, a simple, cost-effective, and environmentally friendly approach is developed to fabricate activated carbon/reduced graphene oxide (AC/r-GO) composite electrodes for supercapacitors with outstanding performance. In such a composite, r-GO provides several much needed critical functions: r-GO not only serves as the binder material improving the AC particle/particle cohesion and electrode-film/substrate adhesion but also improves the electrical conductivity of the composite and provides additional surfaces for ion adsorption. Furthermore, during electrode fabrication, initial GO precursor functions as an effective dispersant for AC, resulting in a stable electrode material slurry. Employing characterization by advanced microscopy techniques, we show that AC and r-GO assemble into an interconnected network structure, resulting in a composite with high specific capacitance, excellent rate capability, and long cycling life stability. Such high-performance electrodes coupled with their relatively simple, scalable, and low-cost fabrication process thereby provide a clear pathway toward large-scale implementation of supercapacitors.

INTRODUCTION

One of the key challenges in the 21st century is unquestionably energy storage, where electrochemical double-layer capacitors (EDLCs), also called supercapacitors or ultracapacitors, offer a promising, low-cost, reliable, and environmentally friendly solution. Supercapacitors feature fast charge-discharge capability, long lifecycles, a wide range of operating temperatures, and operational safety and represent an emerging class of power sources for portable electronic devices.1 As the range of applications of supercapacitors spans from digital cameras, mobile phones, to hybrid vehicles, improving their performance, especially the energy density while maintaining the high power density and cycling stability, remains a primary research focus in the field. Because electrode materials are the key components to the performance of supercapacitors, rationally designing and fabricating high-quality electrode materials play a decisive role in developing next-generation high-performance supercapacitors.2−4

In the industrial cell-manufacturing processes, supercapacitor electrodes are cast onto a current collector from slurries that contain the active electrode material. Today, virtually all industrial supercapacitor manufacturers use coconut shell-derived activated carbon (AC) as the active electrode material due to its high surface area and porosity, stability during cycling, and low cost.5 However, AC alone cannot maintain the required electrical conductivity and mechanical stability of the electrode, and hence electrode slurries generally are also added to the binder system, which maintains both electrical and mechanical integrity of the electrodes once casted on to the current collector. Traditional binder systems are dual-component based, typically a polymer binder and a conductive additive, for two different functionalities. The organic polymer binder, such as poly(vinylidene fluoride) (PVDF),6−10 poly(tetrafluoroethylene) (PTFE),6−10 and carboxy methyl cellulose (CMC),11−13 is used to maintain the cohesion between AC particles and the adhesion of electrode film to the current collector. Conductive additives, such as carbon black (CB), are added to improve the conductivity of the electrode and hence the high rate performance of the supercapacitor. However, the organic polymer binders, themselves, are nonconductive and electrochemically inert, thus making no contribution to the total capacitance of the device. Therefore, they not only...
increase the electrical resistance of the electrode but also increase the "dead" weight of the electrode, which ultimately reduces the overall energy and power densities of the supercapacitor. Furthermore, in realizing greener processes for electrode fabrication, there is a need for replacing fluorine-containing binders such as PVDF and PTFE due to their environmental impact.\textsuperscript{14,15} On the other hand, addition of conductive additives, such as carbon black, may partly alleviate the problem of electrical conduction but can only establish electrical connection through a percolating point-to-point contact and also decreases the tensile strength and extensibility of the binders. Based on the above facts, an ideal binder material should be adhesive, electrically conductive, light, electrochemically active, and additionally form stable electrode slurries.

Although the role of the binder system is as important as that of the active material itself, in supercapacitor research, binders have received the least attention to date. Although several studies have tried to overcome the drawbacks of the binder systems by preparing binder-free self-standing thin film\textsuperscript{16−26} or aerogel electrodes,\textsuperscript{27−29} they are only compatible to an limited extent with large-scale electrode fabrication in industrial supercapacitor manufacturing processes.\textsuperscript{15} In contrast, the development of a monolithic binder material that intrinsically combines electrical conductivity and mechanical integrity by holding AC particles together and further bonding the electrode film to the current collector, while itself also.
acting as an electrochemically active double-layer material, remains strongly underdeveloped in the literature.30–32

Addressing the aforementioned issues, herein we present a rational method to fabricate AC-based supercapacitor electrodes, in which we introduce reduced graphene oxide (r-GO) as a highly efficient, monolithic, conductive binder material, which holds individual AC particles into a compact electrode layer adhered on to the current collector. Our approach synergistically combines mechanical integrity with good electrical conduction between AC particles, with the added benefit that r-GO also additionally serves as an active electrochemical double-layer material itself, thus circumventing a dead weight effect. Thereby, r-GO as a binder in AC supercapacitor electrodes enables much enhanced capacitance performance, rate capability, and cycling life stability compared to conventional dual-component organic polymer/conductive additive binder systems. Importantly, we link this enhanced performance to a comprehensive materials characterization of our novel AC/r-GO composite electrodes, thus developing an extensible framework toward optimization of multifunctional monolithic binders for supercapacitors.

■ RESULTS

For the preparation of the electrode material layer composed of r-GO and AC particles, a commercially available supercapacitor grade coconut shell AC (YP-50F from Kuraray Chemicals) was used. Graphene oxide (GO), synthesized by oxidizing vein graphite following Hummer’s method,33 was used as the starting material of the r-GO. A well-dispersed mixture of GO and AC in propylene carbonate (PC) was coated onto the Al current collector and GO within the electrode material was converted into r-GO by low-temperature thermal reduction. The initial ratio of GO to AC was set to 1:10 by weight. Weight ratio of r-GO to AC in the electrode layer obtained after the electrode layer is applied on to the current collector and the reduction is performed is about 0.65:10 by weight. This is because the weight of the starting GO is reduced by about 35% (from thermogravimetric analysis of GO) during the reduction of the graphene oxide due to the removal of oxide groups.2,34,35 Other weight ratios of GO to AC examined were 10:1, 1:1, and 1:20. For 10:1 and 1:1 weight ratios, the electrochemical performance of the resulting supercapacitors was lower than that for 1:10 weight ratio. For 1:20, the binding between the current collector and the electrode layer and binding between AC particles were poor.

To evaluate the electrochemical performance of AC/r-GO composite electrodes, symmetric two-electrode supercapacitors were fabricated with tetraethylammonium tetrafluoroborate (TEABF4) dissolved in PC as the electrolyte (referred as AC/r-GO supercapacitor). To facilitate direct comparisons, two comparative reference supercapacitors were prepared with electrodes composed of (a) a conventional organic binder, sodium carboxy methyl cellulose (CMC), with AC (1:10), referred to as AC/CMC, and (b) the conventional dual-component binder system CMC and carbon black (CB) with AC (1:1:10), referred to as AC/CMC/CB. For details on materials and methods see the Supporting Information. For further comparisons, the electrochemical performance of a supercapacitor prepared with pure r-GO measured under the same conditions is also given in the Supporting Information (referred to as r-GO supercapacitor) (Supporting Information Figure 1).

Figure 1a shows the cyclic voltammograms (CV) for the AC/r-GO supercapacitor cycled between −2.5 and +2.5 V at scan rates 50, 100, 200, 300, 400, 500, 600, 700, 800, 900, 1000, and 1500 mV/s. Near-rectangular CV curves at all scan rates from 50 to 1500 mV/s with no major pseudocapacitance contribution indicate the ideal double-layer capacitor behavior (also see Supporting Information Figure 2). Additionally, the AC/r-GO electrode is robust enough to be charged and discharged over a wide range of scan rates (50–1500 mV/s) and still maintains its nearly ideal rectangular CV shape. Based on the area of the CV curve, we calculated the specific capacitance of the AC/r-GO supercapacitor as 77.6 F/g at 50 mV/s, and as the scan rate increases to 1500 mV/s, it retains 60.3 F/g with a capacitance retention ratio of 78% (Figure 1d). In contrast, the comparative supercapacitors exhibit lower specific capacitances of 60.3 F/g (AC/CMC) (Figure 1b) and 68.9 F/g (AC/CMC/CB) (Figure 1c) at 50 mV/s with capacitance retention ratio of only 55 and 58%, respectively, at 1500 mV/s (Figure 1d). Markedly distorted CV curves of AC/CMC (Figure 1b) and AC/CMC/CB (Figure 1c) supercapacitors at higher scan rates are a further evidence of their poorer rate capability.

Galvanostatic charge– discharge measurements at varying current densities further illustrate the excellent rate performance of the AC/r-GO electrodes. The specific capacitance, at 2 A/g, was estimated as 79.8 F/g for the AC/r-GO supercapacitor and is thereby higher than 57.5 F/g for the AC/CMC and 66.0 F/g for the AC/CMC/CB supercapacitors (Figure 1e). In addition, the sharp response and a small voltage drop (0.048 V) at the start of the discharge curve are indicative of the formation of an efficient EDL and fast ion transport within the AC/r-GO electrodes. The linear time dependence of the potential suggests the absence of major Faradic processes due to functional groups remaining on r-GO. As the current density increases from 2 to 5 A/g, the AC/r-GO supercapacitor consistently exhibits a higher specific capacitance and shows little change (4.8%) with increase in operating power compared to significant drop in the performance of the comparative counterparts (13.6 and 12.8% for AC/CMC and AC/CMC/CB, respectively) (Figure 1f).

The facilitated ion transport within the AC/r-GO electrodes can also be confirmed from the electrochemical impedance spectra (EIS). A complex plan plot of the impedance data of AC/r-GO along with comparative cells is shown in Figure 1g over the frequency range from 0.01 to 100 000 Hz. An expanded view of the high-frequency region is provided in the inset. The AC/r-GO supercapacitor gives the smallest high-frequency semicircle, indicating the smallest electrode–electrolyte charge transfer resistance. The small charge transfer resistance indicates a very fast charge-transport rate, which is in good agreement with CV and high current charge–discharge studies.36

Furthermore, AC/r-GO also demonstrated excellent cycling stability with ~80% capacitance retention over 20 000 charge–discharge cycles (Figure 1h), indicating a robust architecture of the electrode material. At the same time, ESR slowly increases during cycling.37

The electrochemical characterization results have shown excellent electrochemical response of the AC/r-GO electrodes compared to the comparative electrodes in terms of charge-discharge capacity, rate capability, and cyclic performance. To link this superior electrochemical performance of the novel AC/r-GO composite with its nano- and microstructure, the
composite is characterized using different complementary methods in the following paragraphs.

Scanning electron microscopy (SEM) images of the AC/r-GO composite electrodes indicate a good homogeneity of the AC/r-GO composite surface on a micrometer scale (Figure 2a,b). At higher magnification (Figure 2c), a composite structure of particulate features that are wrapped by sheets of partly electron transparent material is observed. The particulate features are ascribed to the AC particles and the wrapping sheets to the r-GO. It is important to note that hardly any unwrapped AC particles are observed on the electrode surface, suggesting a good interconnection of the AC grains with the r-GO sheets. The absence of charging during the SEM imaging indicates that the network of AC/r-GO is electrically conductive. Both observations are consistent with the good electrochemical performance of the AC/r-GO composites.

To confirm the intimate contact of AC and r-GO in the electrode, the morphological and structural features of the AC/r-GO composite were further examined using (scanning) transmission electron microscopy ((S)TEM). For this purpose, a as-fabricated electrode (from a working electrode batch) was scratched off the Al current collector and the resulting material was deposited onto amorphous carbon TEM grids with regular hole arrays. TEM images at intermediate magnifications of the thus-obtained specimen (Figure 3a,b) show an isolated AC particle, which is covered by an r-GO sheet. The edges of the AC particle present the typical highly porous structure of AC38−40 (Figure 3b) and the r-GO sheet is seen to intimately fold around the AC particle, consistent with the SEM results. The selected area electron diffraction (SAED) pattern in (Figure 3c,d) correspondingly indicates two superimposed phases with graphitic structure:41 one phase with higher graphitization and a partial in-plane orientation (hexagonal
spot pattern, white arrows), which we ascribe to the r-GO sheet, as well as another phase with lower graphitization and random orientation of nanocrystallites (rings in pattern), which is ascribed to the AC particle.

Aberration-corrected STEM elucidates the structure of the r-GO sheets with atomic resolution. In the STEM images (Figure 3e,f), r-GO flakes are found to be typically comprised of few layers of r-GO that are partly turbostratically restacked. Monolayer r-GO regions generally show a hexagonal "honeycomb" lattice, albeit with extensive disorder (Figure 3g), which is fully consistent with previous reports on the atomic structure of r-GO. Successful formation of r-GO in the composite from the GO precursor is further confirmed by X-ray diffractometry (XRD) measurements and Fourier-transform infrared (FTIR) spectroscopy measurement of the AC/r-GO composites and its individual components (Supporting Information Figures 3 and 4).

Conductive tip atomic force microscopy (c-AFM) was used to further evaluate the topography of the AC/r-GO composite and the homogeneity of its electrical conductivity. Topography AFM images (two-dimensional (2D) and three-dimensional (3D)) (plots) and the corresponding c-AFM current maps of an as-fabricated AC/r-GO electrode are shown in Figure 4. The observed topography of the AC/r-GO composite (Figure 4a,b) is found to be in good agreement with the SEM observations above (Figure 2). Global height variations on a low-micrometer scale are found with smaller local variations in the nanometer range. The latter are ascribed to the wrinkling of r-GO sheets and AC particle protrusions. Importantly, the current map (at 1 V bias) in Figure 4c indicates a homogeneous electrical conductivity of the AC/r-GO composite over micrometer lateral length scales, which are consistent with measurements over several AC grains. In particular, no regions with significantly reduced electrical conductivity are observed. This suggests that the r-GO binder successfully electrically connects the individual AC particles and prevents particle isolation. The very small spread in the histogram over the measured current values (Figure 4d) further confirms this. Consistent with electrochemical characterization and the electron microscopy data above, the AC/r-GO electrodes are thus found to be homogeneously electrically conductive and r-GO is thereby identified as an efficient conductive binder material for AC.
**DISCUSSION**

In addressing the shortcomings related to conventional binder systems in supercapacitors, we introduced r-GO as a monolithic multifunctional conductive binder that can replace the conventional dual-component binder system in facile, solution-based electrode coating or printing (on to current collector) process to realize large-scale and low-cost EDLCs. The near-rectangular CV curves of the AC/r-GO electrodes, even at an ultrafast scan rate of 1500 mV/s, linear voltage—time profile, and the highly symmetric charge—discharge characteristics indicate a fast charge propagation and an efficient EDL formation within the AC/r-GO structure. This high-power capability along with excellent cycling stability clearly highlights the superior supercapacitive performance of the AC/r-GO electrodes. Energy density and power density of supercapacitors are also important parameters for their real-life applications.1 For our devices with AC/r-GO as the electrodes, in the TEABF4/PC electrolyte system, a power density of about 134 kW/kg was achieved with an energy density of 17 Wh/kg (specific capacitance of 76 F/g) at the current density of 5 A/g (see the Supporting Information for calculations). Under the same conditions, AC/CMC gave a power density of 21 kW/kg and an energy density of 11 Wh/kg (specific capacitance of 49 F/g), whereas those for AC/CMC/CB were 33 kW/kg and 13 Wh/kg (specific capacitance of 58 F/g). Whereas the power density of comparative supercapacitors, AC/CMC and AC/CMC/CB, remains close to the range for normal AC supercapacitors with conventional binder systems26,29,48 (5–20 kW/kg), that of AC/r-GO devices is far beyond this range.

To further compare these values with the literature data for monoblock electrodes (Figure 5) such as self-standing thin films and aerogels developed with the purpose of overcoming the drawbacks of conventional binders, the power density of our AC/r-GO supercapacitors is much higher than the maximum power density of 5.1 kW/kg reported for a graphene paper electrode pillared by carbon black nanoparticles in aqueous electrolyte,16 the power density of 7.3 kW/kg for AC/carbon nanotube paper electrode in aqueous electrolyte,26 and that of 6.2 kW/kg obtained for composite organogels of r-GO and AC in organic electrolyte of TEABF4/PC.29 Moreover, we compare the power density of AC/r-GO devices with that of other composite electrodes based on carbon spheres and r-GO prepared using conventional binders (Figure 5). Power densities of 4.2, 10, and 15.4 kW/kg have been reported for 3D hierarchical composite electrode of slightly crumpled r-GO sheets intercalated with mesoporous carbon spheres (prepared using PTFE) in an aqueous electrolyte,49 for an electrode of r-GO/AC (prepared using PTFE) in ionic liquid electrolyte,50 and a nanostructured electrode comprising functionalized carbon spheres and r-GO sheets (prepared using PTFE) in aqueous electrolyte, respectively,51 again confirming the high-power delivery of the AC/r-GO electrodes.

The excellent electrochemical performance of our AC/r-GO electrodes stems from the role r-GO plays within the composite as well as the highly interconnected 3D structure that favors fast charge transportation of the electrode layer. In general, this is shown schematically in Figure 6.

![Figure 6. Schematic illustration of the AC/r-GO electrode.](image)

r-GO is well-known for its outstanding electrical conductivity and high surface area for ion adsorption.52,53 When processed properly, these useful properties can be harnessed and such a structure provides many benefits for the improvement of energy storage applications. Within the AC/r-GO composite, r-GO has several critical functions for efficient use of AC for energy storage; r-GO facilitates the mechanical connection among AC particles in the electrode, serves to enable the mechanical integrity of electrode layer and its adhesion to the current collector and provides additional surface for EDL ion adsorption, and finally/hence replaces the need for conductive additives by improving the electrical conductivity of the composite.

Comprehensive materials characterization of our AC/r-GO electrodes clearly highlights the highly interconnected 3D structure of the composite. The r-GO layers are capable of surface contact with AC particles with low contact resistance, as shown here by our combination of SEM, TEM, and c-AFM. Conductive r-GO layers that wrap around AC particles three-dimensionally not only provide additional electron transport pathways but also decrease the ion diffusion length that favors fast charge transportation resulting in high rate performance.
Furthermore, holes created on the r-GO sheets due to the removal of in-plane C atoms during the reduction process may provide cross-plane diffusion channels for electrolyte ions.54–55 Functioning as the ion diffusion shortcuts between different layers of graphene, they greatly speed up the ion transport across the entire electrode film. In sharp contrast, in conventional binder systems, nonconductive polymers and conductive carbon particles that make electrical connection through a percolating point-to-point contact greatly retard mass transfer and thereby place limitations on the high-power performance of EDLCs. (As a result, the CV curves of such electrodes distort with increasing sweep rate, as also seen here for the conventional r-GO-free comparative supercapacitors.)

Functional groups remaining on the r-GO layers after reducing at 300 °C, such as carbonyls, ethers, and phenols, play a key role in determining the adhesive properties of r-GO as the layers make surface contact with AC particles and the current collector (Supporting Information Figure 4). The noncovalent interactions (such as hydrogen bonding and π−π interactions), which are established between r-GO layers themselves, and between r-GO and AC particles and current collector, assist r-GO to maintain the structural stability of the electrode as the binder material, by keeping the AC particles bound together to the current collector. This is consistent with the reports that polymers with high adhesive properties, such as alginate and poly(acrylic acid), form strong hydrogen bonds with the oxide surface on the active materials.58 Also, incorporation of such noncovalent interactions is expected to be a useful tool in tuning network properties between binder polymers and active particles.59,60 Additionally, they have been seen to improve the cycling stability of the electrodes.58 Moreover, these functional groups are also beneficial to the superior supercapacitive performances by improving electrolyte wettability and electrical conductivity.61 The unique combination of adhesive, conductive, and high surface area properties of r-GO into a single electrode results in excellent electrochemical performance for AC/r-GO supercapacitors.

In addition, from the perspective of the electrode fabrication process, the initial GO precursor, which is present in the electrode slurry, functions as an effective dispersant for AC in PC, resulting in a stable electrode material slurry which facilitates the electrode film fabrication. We find that the AC/GO slurry in PC can be applied to a collector uniformly so as to form a smooth layer on the collector, which contains the active substance in a uniform composition. This is consistent with previous reports on GO as an excellent two-dimensional polymeric dispersing agent for dispersion of various carbon-based materials.52–54

**CONCLUSIONS**

Development of an efficient binder system is a frequently overlooked bottleneck in realizing high-performance supercapacitors. In conventional methods of electrode fabrication, dual-component conductive additive–polymer binder system is added to improve the electrode stability at the expense of electrical conductivity and capacitance/weight ratio of the electrode. Therefore, a binder material that combines mechanical stability, high electrical conductivity, and significant ion-adsorption capability and can also disperse the active materials and stabilize them during solution-based electrode fabrication is urgently needed. This study reported the use of r-GO as such an ideal binder for AC-based high-power supercapacitor electrodes. With its adhesive and conductive properties and high surface area, r-GO serves as a single-component conductive binder with multifunctionality to maintain high electronic conductivity and mechanical adhesion while functioning as an electrochemically active material for charge storage. Its electrochemical properties were demonstrated by the outstanding performance of the AC/r-GO composite electrodes prepared without any additional polymer binders or conductive additives. The specific capacitance of the AC/r-GO electrodes increased by ∼20–38% at 2 A/g compared to comparative supercapacitors, with power density values as high as 134 kW/kg at 5 A/g. These AC/r-GO composite electrodes exhibit excellent cycling performance with 80% capacitance retention over 20 000 charge–discharge cycles. SEM, TEM, STEM observations, and c-AFM measurements together demonstrate structurally and electrically interconnected and homogeneous “wrap around” structure of AC/r-GO composites, linking the superior electrochemical performance to the microstructure of the AC/r-GO composites. The initial GO precursor, which is present in the electrode slurry, also functions as an effective dispersant for AC, resulting in stable electrode material slurry during solution-based electrode fabrication. Importantly, such a facile solution-based process combined with electrode coating or printing on to current collector provides a practical route for making large-scale and low-cost devices.65 This combination of unique properties makes the reported monolithic multifunctional r-GO binder suitable for significantly enhancing the power capacity of industrial-type AC supercapacitors used in applications such as electric vehicles.

**EXPERIMENTAL SECTION**

Fabrication of AC/r-GO Electrodes and Supercapacitors. Graphene oxide (GO), synthesized using graphite oxide derived from vein graphite (purity >99%, Bogala Graphite, Sri Lanka) following Hummers’ method,53 is used as the starting material of the reduced graphene oxide (r-GO), which serves as the multifunctional binder and conductive additive in the electrode composite. The as-synthesized graphite oxide was washed with a diluted HCl solution to completely remove residual salts. It was further washed with water until the pH of the rinse water became neutral (pH = 7). Ultrapure Milli-Q water was used in all the experiments. The as-purified graphite oxide suspensions were then dispersed in water. The exfoliation of graphite oxide to GO was achieved by ultrasonication of the dispersion. Graphene oxide dispersions prepared according to the above procedure were dried in an oven maintained at a temperature of 50 °C. Completely dried GO was used for further characterization and supercapacitor electrode fabrication. As produced, completely dried GO was characterized using different complementary methods6 and used for activated carbon (AC)/r-GO electrode fabrication. The results clearly confirmed the successful fabrication of GO decorated with hydroxyl, carbonyl, ether, and carboxyl groups, which in its restacked dried form keeps an interlayer distance of ~0.79 nm.

A commercially available, supercapacitor grade, steam-activated, coconut shell AC (YP-50F from Kuraray Carbons, surface area in the range of 1500–1800 m2/g, pore volume 0.7 cm3/g and ash content <1%) was used as received without further processing to prepare the composite.

For supercapacitor electrode fabrication, GO and AC were mixed in propylene carbonate (PC) (Sigma-Aldrich, 99.7%) to...
prepare a slurry. The investigated ratios by weight of GO to AC were 10:1, 1:1, 1:10, and 1:20, where the best supercapacitor performance was observed for 1:10. The mixed solution was sonicated for further dispersion of GO and the electrodes were prepared by slurry casting onto conductive coated Al current collector. The AC/GO composite coated on the current collector was heated at 300 °C in an inert (Ar) atmosphere, at low pressure (1 kPa) for 30 min to convert the GO within the composite electrode material into r-GO. Electrodes from GO/AC mixtures of 10:1, 1:1, and 1:10 were structurally stable after electrode fabrication and reduction. For 1:20, the binding between current collector and the electrode layer and binding between AC particles were poor.

Fabrication of Comparative Supercapacitors. Comparative supercapacitors were fabricated in a manner similar to that of the above-described test supercapacitor with AC/r-GO composite electrodes (referred as AC/r-GO supercapacitor).

For comparison, the following supercapacitors electrodes were fabricated:

1. Electrodes having an electrode material layer composed of the AC and sodium carboxy methyl cellulose (CMC, Sigma-Aldrich, average molecular weight (Mn) ~90 000), which is a conventional binder, instead of r-GO obtained by reducing GO (referred as AC/CMC supercapacitor), with the weight ratio of CMC/AC = 1:10.

2. Electrodes having an electrode material layer composed of the AC and sodium carboxy methyl cellulose (CMC) and carbon black (CB, Cabot Corporation), which is a conventional conductive additive, instead of r-GO obtained by reducing GO (referred as AC/CMC/CB supercapacitor), with weight ratio of CB/CMC/AC = 1:1:10.

3. Electrodes having an electrode material layer composed only of r-GO obtained by reducing GO under the same reduction conditions used for the reduction of GO in the AC/GO composite (referred as r-GO supercapacitor).

Materials Characterization. The structure of AC/r-GO composite was characterized by field-emission scanning electron microscopy (SEM, Hitachi S-5500), conductive atomic force microscopy (c-AFM, NT-MDT NTEGRA Spectra), transmission electron microscopy (TEM, Philips CM200 at 200 kV), bright field TEM and selected area electron diffraction (SAED), and scanning transmission electron microscopy (STEM, aberration-corrected Nion UltraSTEM 100 at 60 kV, medium angle annular dark field imaging). For TEM and STEM, the examined material was obtained directly from an as-fabricated AC/r-GO composite electrode coated on Al foil (from a working electrode batch). The composite was manually scratched off the Al foil and the thus-obtained material was dispersed in isopropyl alcohol, and then via drop cast transferred to TEM grids (Quantifoil holey amorphous carbon). For SEM, the electrode was used as it is without any further processing.

Electrochemical Testing. The electrochemical behavior of the as-prepared composite electrodes was characterized by cyclic voltammetry (CV), galvanostatic charge–discharge measurements, and electrochemical impedance spectroscopy (EIS) with two symmetric electrodes in coin cell geometry using an Autolab electrochemical interface instrument (PGSTAT 302N). A solution of 1M TEABF4 (Sigma) in PC (Sigma) was used as the electrolyte and a filter paper (Whatman, grade no. 1) was used as the separator. The coin cell preparation was carried out in a nitrogen-filled glovebox (MBrAun) with oxygen and moisture levels of <1 ppm. The electrochemical measurements were taken using a two-electrode system at ambient temperature.

Galvanostatic charge–discharge cycling was conducted between the potential limits of 0 to +2.5 V at different current densities (2, 3, 4, and 5 A/g) based on the total mass of active materials. The cell capacitance at different current densities was calculated according to $C = I/(\Delta \nu/\Delta t)$, where $C$ is the measured capacitance for the two-electrode cell (F), $I$ is the constant discharging current (A), $\Delta \nu$ is the voltage difference from 90 to 30% of the discharge voltage range (V), and $\Delta t$ is the time required to go from 90 to 30% of the discharge voltage range (s).67

The CV curves were recorded in the potential range from −2.5 to +2.5 V at different scan rates (50, 100, 200, 300, 400, 500, 600, 700, 800, 900, 1000, and 1500 mV/s). The cell capacitance at different scan rates was calculated according to the equation $C = (\int \Delta \nu/V, 26)$, where $C$ is the cell capacitance (F), $I$ is the response current, $V$ is the potential window, and $\nu$ is the CV scan rate (mV/s).

The specific capacitances were calculated from the following equation.

$$C_{sp} (F/g) = 4 \times C/m$$

Specific capacitance $C_{sp}$ is the capacitance per unit mass for one electrode, where $C$ is the measured capacitance for the two-electrode cell (F) and $m$ is the total mass of the active material (g) in both electrodes. The multipler of 4 adjusts the capacitance of the cell and the combined mass of two electrodes to the capacitance and mass of a single electrode.

Frequency response analysis was carried out in the range of 0.01–100 000 Hz with a DC bias of 10 mV. Extrapolating the curve on the Nyquist plot to intersect the X-axis yields ESR values.67

The power density of the AC/r-GO device was calculated from the galvanostatic charge–discharge curves at different charge–discharge current densities using the following formula.

$$P = \frac{(\Delta V)^2}{4R_{ESR} m}$$

where $P$ is the power density (kW/kg), $\Delta V$ is the operating voltage window, and $m$ is the total mass of the active material in both electrodes. $R_{ESR}$ is the internal resistance of the device estimated from the voltage drop (IR drop) at the beginning of the discharge curve and calculated from the change in voltage ($V_{drop}$) divided by the total change in the current applied using the formula

$$R_{ESR} = \frac{V_{drop}}{I}$$

The energy density of the device was obtained from the following formula.

$$E = C_{sp} \times (\Delta V)^2/(2 \times 3600)$$

where $E$ is the energy density (Wh/kg), $C_{sp}$ is the specific capacitance (F/g), and $\Delta V$ is the operating voltage window.
The energy density ($E$) can also be obtained by measuring the discharge time at a certain current density and calculated according to the following equation.68

$$E = \int_{t_1}^{t_2} IV \, dt$$

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b01075

Cyclic voltammograms and Nyquist plots for the r-GO supercapacitor (Figure 1); CV profiles and dependence of the capacitive current on the applied scan rates between 50 and 1500 mV/s for the AC/r-GO supercapacitor (Figure 2); XRD patterns of r-GO, AC/r-GO composite and AC (Figure 3); FTIR spectra of GO, r-GO, AC/r-GO, and AC (Figure 4) (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

*E-mail: dtlg2@cam.ac.uk.*

**ORCID**

Dona T. L. Galhena: 0000-0002-8619-164X
Bernhard C. Bayer: 0000-0002-4829-3207
Jannik C. Meyer: 0000-0003-4023-0778
Stefan Hofmann: 0000-0001-6375-1459
Gehan A. J. Amaratunga: 0000-0002-8614-2864

**Author Contributions**

D.T.L.G. synthesized graphene oxide, prepared supercapacitor cells, performed electrochemical experiments, SEM, XRD, and FTIR characterization, and analyzed the data. B.C.B and J.C.M. performed (S)TEM and AFM characterization. S.H. and G.A.J.A. supervised the project. D.T.L.G. and B.C.B. wrote the article. All the authors discussed the results and commented on the manuscript.

**Notes**

The authors declare no competing financial interest.

The authors declare that the data supporting the findings of this study are available within the article and its Supporting Information files.

**ACKNOWLEDGMENTS**

D.T.L.G. acknowledges the technical support by J. N. R. Grundy (University of Cambridge) and A. Osborne (Metrohm U.K.) and financial support from Newnham College, Cambridge, and the Cambridge Commonwealth Trust. B.C.B. acknowledges funding from the European Union’s Horizon 2020 research and innovation program under the Marie Skłodowska-Curie grant agreement 656214-2DInterFOX. J.C.M. acknowledges support from the Austrian Science Fund (FWF, P25721-N20).

**REFERENCES**


(60) Choi, J. W.; Aurbach, D. Promise and reality of post-lithium-ion batteries with high energy densities. Nat. Rev. Mater. 2016, 1, No. 16013.


(64) Li, Y.; Yang, J.; Zhao, Q.; Li, Y. Dispersing carbon-based nanomaterials in aqueous phase by graphene oxides. Langmuir 2013, 29, 13527–13534.


DOI 10.1021/acsomega.8b01075
ACS Omega 2018, 3, 9246–9255