RSC Advances



View Article Online

View Journal | View Issue

PAPER



Cite this: RSC Adv., 2020, 10, 38357

Received 9th September 2020 Accepted 7th October 2020 DOI: 10 1039/d0ra07748h

rsc.li/rsc-advances

Introduction 1

Recently, great interest has been focused on the application of nanostructured carbons for electrocatalysis1 and energy storage applications,²⁻⁵ particularly in electrochemical capacitors or supercapacitors (SCs).⁶⁻¹⁰ Due to a large specific surface area connected with high mesoporosity, nanostructured carbons have a high capability for charge accumulation at the electrode/ electrolyte interface (known as an electric double layer).11-13 Accumulation of an ionically conducting electrolyte solution on an electronically conducting electrode surface leads to the establishment of electric double-layer capacitance (EDLC). In brief, the electrochemical performance of these systems depends on the control of mass and charge transport (*i.e.*,

Opening the internal structure for transport of ions: improvement of the structural and chemical properties of single-walled carbon nanohorns for supercapacitor electrodes[†]

Wojciech Zieba, ២ ^a Piotr Olejnik, ២ ^b Stanislaw Koter, ២ ^c Piotr Kowalczyk, ២ ^d Marta E. Plonska-Brzezinska ២ *^b and Artur P. Terzyk 咆 *^a

We investigated the electrochemical performance of single-walled carbon nanohorns (SWCNHs) for use as supercapacitor electrodes. For the first time, we used acid-treatment for oxidation of SWCNHs and hole creation in their structure. A detailed study was performed on the correlation between the oxidation of SWCNHs via acid treatment and variable acid treatment times, the structural properties of the oxidized carbon nanostructures, and the specific capacitance of the SWCNH electrodes. We showed that simple functionalization of carbon nanostructures under controlled conditions leads to an almost 3-fold increase in their specific capacitance (from 65 to 180 F g^{-1} in 0.1 M H_2SO_4). This phenomenon indicates higher accessibility of the surface area of the electrodes by electrolyte ions as a result of gradual opening of the SWCNH internal channels.

> electronic and ionic mobility) and electron transfer kinetics in multiphase boundaries,¹⁴ and these are size dependent processes. To this end, much attention is focused on preparation of nanostructured carbon materials with controlled properties, including internal and external porosity, electrical conductivity and rich functional chemistry.15-17

> In literature, we find many reports of the use of nanostructured carbon materials as electrode materials in SCs,4,5,11 and the most popular are carbon nanotubes (CNT) and graphene electrodes.18 For example, multiwalled CNTs were heated to enhance their specific surface area and pore size distribution, and after thermal treatment, the CNT electrodes reached a maximum specific capacitance of 180 F g⁻¹ with a power density of 20 kW kg⁻¹.¹⁹ Even with special preparation of CNTs, their specific surface area is still much lower than that of mesoporous carbons, ordered mesoporous carbons or activated carbon electrodes (specific surface areas in the range of 600 and 3000 m² g⁻¹, respectively).²⁰⁻²² Surface modification of carbon nanostructures (CNs) via acidic or alkaline treatment, which introduces oxygen functional groups, has an additional effect and improves the electrochemical properties of SCs.13,23,24 It should also be emphasized that the gravimetric capacity of carbon materials does not increase linearly with the specific surface area, and based on the density functional theory model, it reaches a plateau for values of *ca.* 1200 m² g⁻¹.²² The authors suggested that this limitation of the gravimetric capacity can be ascribed to space constriction for charge accommodation inside the pore walls.²² In this context, a further increase in the specific

[&]quot;Faculty of Chemistry, Physicochemistry of Carbon Materials Research Group, Nicolaus Copernicus University in Toruń, Gagarin Street 7, 87-100 Toruń, Poland. E-mail: aterzyk@chem.uni.torun.pl

^bDepartment of Organic Chemistry, Faculty of Pharmacy with the Division of Laboratory Medicine, Medical University of Bialystok, Mickiewicza 2A, 15-222 Bialystok, Poland. E-mail: marta.plonska-brzezinska@umb.edu.pl

^{&#}x27;Faculty of Chemistry, Department of Physical Chemistry, Nicolaus Copernicus University in Toruń, Gagarin Street 7, 87-100 Toruń, Poland

^dCollege of Science, Health, Engineering and Education, Murdoch University, WA, 6150. Australia

[†] Electronic supplementary information (ESI) available: Selected GCMC simulated local N₂ (77 K) adsorption isotherms and the fit to experimental data, high-resolution C 1s XPS and N 1s spectra, deconvolution of Raman spectra, CVs of SWCNH-ox-1.5 in 0.1 M different supporting electrolytes, specific capacitance of SWCNH, SWCNH-ox and r-SWCNH-ox in aqueous solutions of different supporting electrolytes. See DOI: 10.1039/d0ra07748h

surface of carbonaceous materials does not affect the increase in gravimetric capacity.

Single-walled carbon nanohorns (SWCNH) belong to the class of novel carbon nanomaterials. Thousands of cone-end shaped tubules with diameters of 2-5 nm are agglomerated in spheres with a diameter in the range of 60-120 nm. The spheres can occur in few different forms: dahlia, petal-dahlia, bud and seed. Dahlia and petal-dahlia shapes have long cone endings on a spherical surface in contrast to bud and seed shapes.²⁵ The material is produced by CO₂ laser ablation of graphite and is catalyst-free with notably high purity of up to 90%. To obtain access to the internal volume of SWCNH, various oxidation reactions have been applied, e.g., heat treatment under different atmospheres, such as air,26,27 CO2 28 and O2.29-32 Less popular methods are the so-called "wet methods" using, e.g., H_2O_2 , ³³⁻³⁶ concentrated acids or their mixtures, e.g., nitric acid³⁷⁻⁴¹ or "piranha" solution.42,43 Oxidation changes selected chemical and physical properties of the material, thus increasing the specific surface area and pore volume. Surface oxygen groups increase hydrophilicity, enabling the formation of stable colloids in water.39 Generally, these functionalization approaches increase the specific effective surface of the SWCNHs while optimizing the structural parameters that influence the capacitive properties of the SWCNH electrodes.^{27,29,31,44} Therefore, due to a large surface area, homogeneous structure and capacity for modifications, SWCNHs have found application in many electrochemical areas, such as Li-ion batteries, 30,45-48 SCs, 26,27,29,31,44,49,50 sensors 41,51-53 and/or fuel cells.54-56 Moreover, SWCNHs could also be used as composites to construct complicated and hierarchical systems that exhibit optimized targeted properties.33

For the first time, we report the structural changes of SWCNHs that were performed using oxidation by nitric acid. A

variable acidic treatment time was applied, and the correlations between the structural changes and the electrochemical properties of the obtained CNs were examined. We showed that simple functionalization of CNs under controlled conditions leads to an almost 3-fold increase in their specific capacitance due to an 'opening structure' for ion mobility. Additionally, the resulting SWCNH electrode has excellent electrochemical stability.

2 Results and discussion

2.1. Structural and textural characterization of SWCNHs

The physicochemical properties of CNs are closely related to their structure, type of carbon atoms, number of defects on the surface, type of functional groups, etc. Therefore, we performed systematic experimental and theoretical studies of the structure and surface of the SWCNHs modified in the oxidation process. Fig. 1A presents the N₂ adsorption-desorption isotherms measured for the pristine and the oxidized SWCNHs. The shape of the isotherm measured for the pristine SWCNHs (Fig. 1A) suggests the presence of mesopores. The BET surface area (S_{BET}) was calculated using the Brunauer-Emmett-Teller57 adsorption isotherm equation. However, to make the porosity parameters more realistic, we used a series of local isotherms for cylinders calculated using grand canonical Monte Carlo (GCMC) simulations, and this procedure is described in detail in the Experimental section (fit quality is shown in Fig. S1 and S2^{\dagger}). The S_{BET} value for SWCNHs (Table 1) is in good agreement with the values reported for this system by other authors58 and supplied by the producer.

From the data collected in Fig. 1A, B, S1 and S2^{\dagger} it can be concluded that increasing the oxidation time of the SWCNHs led to increasing N₂ adsorption, which consequently led to

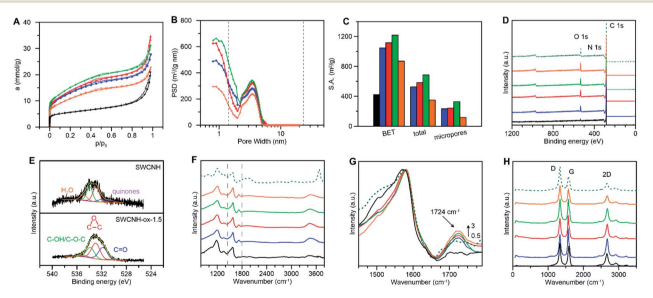


Fig. 1 (A) Low temperature N₂ adsorption–desorption isotherms, (B) pore size distributions, (C) pore surface areas, (D) XPS survey spectra, (E) high-resolution O 1s XPS spectra, (F and G) FTIR and (H) Raman spectra for SWCNH (black), SWCNH-ox-0.5 (blue), SWCNH-ox-1.0 (red), SWCNH-ox-1.5 (green), SWCNH-ox-3 (orange) and r-SWCNH-ox-1.5 (dotted dark green). For r-SWCNH-ox-1.5, practically the same adsorption isotherms and porosity parameters as for SWCNH-ox-1.5 are observed, and thus the adsorption and pore size distribution results for this sample are omitted for clarity.

Materials	$S_{\rm BET}$ (m ² g ⁻	¹) S_{total} (m ² g	$^{-1}$) $S_{\rm meso}$ (m ² g ⁻	$^{-1}$) S_{micro} (m ² g	$^{-1}$) $\nu_{\rm total}$ (cm ³ g ⁻	¹) $\nu_{\rm meso}$ (cm ³ g ⁻¹) $\nu_{\rm micro} \left({\rm cm}^3 {\rm g}^- \right)$	¹) D _{micro} (nm)) D _{meso} (nm)
SWCNH-ox-0.5	1047	527	294	233	0.33	0.25	0.08	0.69	3.4
SWCNH-ox-1	1116	584	342	242	0.37	0.30	0.07	0.83	3.3
SWCNH-ox-1.5	1219	687	358	328	0.42	0.31	0.11	0.90	3.4
SWCNH-ox-3	872	350	234	116	0.24	0.20	0.04	0.86	3.4

Table 1Structural parameters of internal nanospaces of SWCNHs obtained from the BET model and general adsorption integral equationimplemented with the kernel of local N2 adsorption isotherms (77 K) simulated in cylindrical carbon pores using GCMC^a

 a S_{BET} – the apparent BET surface area, equal to 421 m² g⁻¹ for pristine SWCNH, S_{total} – total internal surface area, S_{meso} – internal surface area of mesopores, S_{micro} – internal surface area of micropores, v_{total} – total internal pore volume, v_{meso} – internal volume of mesopores, v_{micro} – internal volume of micropores, v_{micro} – internal volume of micropores, v_{micro} – average diameter of micropores. For r-SWCNH-ox-1.5 (not shown), practically the same isotherms and porosity parameters as for SWCNH-ox-1.5 were observed.

increasing S_{BET} , S_{total} and total pore volumes (v_{total}) with the creation of micropores on the SWCNH surface (Table 1). This observation is schematically presented in Fig. 1C. The S_{BET} values are between 872 (SWCNH-ox-3) and 1219 m² g⁻¹ (SWCNH-ox-1.5) for the oxidized materials. The maximum values for all of these parameters were recorded for SWCNH-ox-1.5 (Table 1). Therefore, the reduction reaction of the SWCNH-ox-1.5, which was used in further studies as a comparison.

Oxidation treatment of SWCNHs causes gradual opening of the internal channels of the carbon nanotubes, as confirmed by the HR-TEM images collected in Fig. 2B. However, for oxidation time longer than 1.5 h, burn-off of tubules and creation of amorphous carbon around the walls were observed (Fig. 2C). These conclusions confirm the decrease in N₂ adsorption (Fig. 1A) and the consequent decrease of the S_{BET} and the v_{total} values that were observed for SWCNH-ox-3 (Table 1). However, for electrochemical application of CNs, the most commonly used parameter is S_{BET} , a more justified approach is the use of internal surfaces, the parameters connected with the presence of meso and micropores. The S_{meso} and the S_{micro} values for the oxidized SWCNHs were increased in the order SWCNH-ox-3 < SWCNH-ox-0.5 < SWCNH-ox-1 < SWCNH-ox-1.5 (Table 1). The sharp drop in the S_{meso} and the S_{micro} values with a long oxidation time (SWCNH-ox-3) is related to the adsorption of amorphous carbon on the SWCNH surface, as mentioned

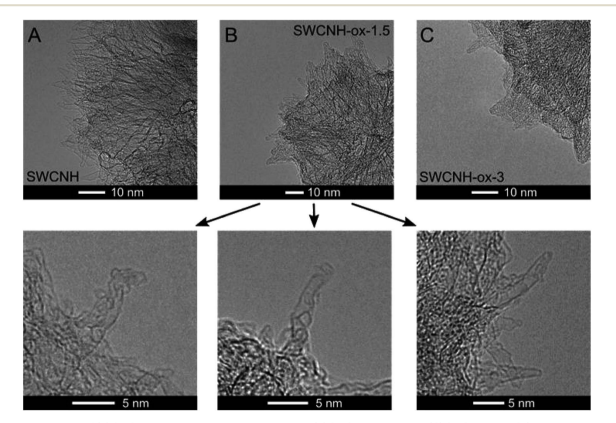


Fig. 2 HR-TEM images of (A) SWCNHs and selected oxidized samples, (B) SWCNH-ox-1.5 and (C) SWCNH-ox-3.0 (bottom panel shows the holes in the horns of the SWCNH-ox-1.5 sample).

previously, that was observed by HR-TEM studies (Fig. 2C). An analogous relationship for the SWCNH structures and oxidation time was observed for the volume of meso and micropores (Table 1). It must be emphasized that gradual treatment of the SWCNHs with HNO₃ shows minimal change in the average diameters of mesopores (D_{meso}) and slightly changes the diameters of micropores (D_{micro}) (Table 1). A longer oxidation process led to an increase in the average diameter of micropores from approximately 0.7 to 0.9 nm.

X-ray photoelectron spectroscopy (XPS) and Fourier transform infrared spectroscopy (FTIR) were performed to confirm the oxidation of the SWCNH surface. Quantitative and qualitative studies were conducted using XPS (Fig. 1D and E).

The survey spectra for the SWCNHs and the oxidized materials are compared in Fig. 1D, and the elemental quantification is shown in Fig. S3–S5.† The XPS spectra for the pristine SWCNHs contain one peak (of C 1s) and two peaks (of C 1s and O 1s) for SWCNH-ox (Fig. 1D). A small N 1s peak is also observed for the reduced sample (Fig. 1D).

Deconvolution of the high-resolution C 1s core level spectra for the pristine SWCNHs shows three main peaks centered at 284.7 eV (56.4% at. concentration of C; C=C sp²; graphitic carbon), 284.9 eV (13.6% at. concentration of C; C-C sp³) and 285.2 eV (8.2% at. concentration of C; high C-C sp³/C-H sp³), (Fig. S3[†]).⁵⁹⁻⁶¹ After the oxidation reaction, the C 1s XP spectrum exhibits additional energy regions that confirm the presence of oxygen-containing groups on the SWCNH surface and could be deconvoluted into six peaks at 285.7 eV (ether groups, C-O-C), 286.1 eV (hydroxyl groups, C-OH), 286.7 eV (epoxy groups, (CC) O), 287.3 eV (carbonyl groups, C=O), 289.2 eV (carboxyl groups, COOH, very weak signal), (Fig. S3[†]),^{62,63} and an additional peak centered at 284.2 eV related to the vacancy defects of the graphene lattice.⁶⁴ The level of oxidation of the SWCNHs was determined from the areas under the C 1s and O 1s peaks (Table S1[†]). As expected, the amount of functional groups (C-O-C, C-OH, (CC)>O, C=O and COOH) increased with increasing oxidation reaction time, and after reaching the reaction time of one and half hours, their amounts on the surface of the CN decrease (a progressive decrease in the C 1s/O 1s ratio, Table 2). However, it should be emphasized that this observation is correct in considering the total number of oxygen-containing groups (Table S1[†]). The amount of COOH groups on the SWCNH surface increased proportionally with the increase in the oxidation time (the value for 0.5 h was not determined; in the range of 1-3 h oxidation reaction the values are between 1.0%-2.3% at. concentration of C).

Deconvolution of the O 1s XPS peaks produced additional information on the nature of the surface-containing groups. However, for the SWCNHs, selected oxygen functionalities were found with the contribution of chemisorbed water. The calculated percentages of these groups are quite low, and they are shown in Fig. S4† for comparison purposes only. As shown in Fig. 1E, the O 1s core signal for the oxidized CNs could be deconvoluted into peaks at 530.5 eV (quinones),⁶⁵ 531.8 eV (C=O),⁶⁶ 532.9 eV ((CC)>O),⁶⁷ 533.8 eV (C-OH, C-O-C) and 535.0 eV (chemisorbed water).^{68,69} The percentage of these functional groups for all oxidized SWCNH materials is shown in Fig. S4.† After reduction of the SWCNH-ox-1.5 by hydrazine, the amount of oxygen-containing groups on the SWCNH surface decreases significantly, with the exception of the quinone groups, which proves that the groups containing O were successfully reduced.

This observation is confirmed by high-resolution N 1s XPS studies, which were decomposed into four identified components (Fig. S5[†]). For r-SWCNH-ox-1.5, we observed the appearance of N–H bonds at 399.8 eV, pyrrolic or amine moieties at 400.2 and 400.6 eV, and quaternary N at 401.8 eV.^{64,70,71} We observed the hydrophilization of the SWCNH surface by the creation of surface oxides. The ratio of C 1s/O 1s peaks decreases rapidly after a short oxidation process (after 0.5 h), from 15.110 to 4.143, and subsequently changes during a longer oxidation time, reaching a value of 2.603 for SWCNH-ox-3 (Table 2) and a value of 4.05 for r-SWCNH-ox-1.5.

This observation is also strongly supported by the decrease in the ζ potential values (from -6.65 to -29.65) determined at pH = 2 (Table 2). As confirmed by the XPS data (Table S1[†]) the presence of nitrogen-containing groups for the r-SWCNH-ox-1.5 sample is manifested by the positive ζ potential value (ζ potential = +2.45) caused by the basic nature of selected surface nitrogen groups.

The FTIR-HATR technique was used to determine the nature of the functional groups on the SWCNH surface. Fig. 1F and G show the FTIR spectra of the pristine, oxidized and reduced SWCNHs. Three clearly distinct bands appear for the pristine SWCNHs at 1573, 1356 and 1185 cm⁻¹, with small shoulders at 1059 cm⁻¹. Thus, the band at 1573 cm⁻¹ might be assigned to stretching vibrations of the C=C aromatic groups.⁷²⁻⁷⁴ The presence of the peaks at 1356 and 1185 cm⁻¹ are strongly affected by the defective structure of CNs and was also observed for ultradispersed diamond powders.⁷⁵ However, it should be emphasized that these peaks are found in the area of several overlapping peaks that occur in the range from 1400 to 900 cm⁻¹, therefore, the determination of their intensities is

Table 2 Selected parameters of the studied materials obtained from Raman spectroscopy, XPS and ζ potential values

Materials	C 1s/O 1s	$\nu_{\rm D} \left({\rm cm}^{-1} ight)$	$\nu_{\rm G} \left({\rm cm}^{-1} \right)$	$I_{\rm D}/I_{\rm G}$	ζ potential (mV)
SWCNH	15.110	1332	1580	1.20	n.a.
SWCNH-ox-0.5	4.143	1335	1575	1.28	-6.65
SWCNH-ox-1	3.087	1335	1571	1.37	-17.91
SWCNH-ox-1.5	3.172	1338	1575	1.50	-22.35
SWCNH-ox-3	2.603	1336	1574	1.68	-29.65
r-SWCNH-ox-1.5	4.050	1334	1577	1.66	+2.45

Paper

often ambiguous. In the literature, this phenomenon is often translated as the occurrence of 'dangling bonds' on the surface of CNs, which are sensitive to chemisorption of foreign elements.⁷⁵ These surface carbon atoms have free valances that can be saturated by chemisorption or that might interact or react with foreign elements (H_2O , H, N, O, *etc.*) to form bonded surface functional groups.⁷⁶ These absorption features still exist in the region of 1600 to 900 cm⁻¹ in the reduced form of the SWCNHs, but they are somewhat different. Therefore, this wavenumber region might be assigned to edge carbon atoms on the SWCNH surface, which are sensitive to chemical reactions (oxidation or reduction), that in successive stages have changed the chemical nature.

Successful oxidation of the SWCNH surface is strongly supported by the peak at 1724 cm^{-1} , which can be assigned to C=O stretching vibrations (Fig. 1G),75 and these vibrations are not observed for the pristine SWCNHs. The spectra of the oxidized SWCNH materials clearly indicate that a longer oxidation process of the CNs leads to an increase in the intensity of the C=O stretching vibrations (from 0.5 h to 3 h), which consequently proves the increase in the amount of oxygen-containing groups on the SWCNH surface. For the reduced form of the SWCNHs, the intensity of this peak decreased, but an additional shoulder is observed at 1716 cm⁻¹ (dashed line on Fig. 1G). This peak might be assigned to C=O stretching vibrations in ketones or aldehydes.⁷⁴ Based on the FTIR spectroscopic characterization of functionalized single-walled carbon nanotubes, the band detected at approximately 1570 cm⁻¹ was attributed to C=C stretching vibration of functionalized carbons.72,77 The increased intensity in the high frequency region between 3700 and 3200 cm⁻¹ could also indicate successful oxidation of the SWCNH surface.72 This band is the result of the O-H stretching vibration and can be observed in the spectra in Fig. 1F as a broad band with a maximum at 3440 cm⁻¹. Upon surface reduction with hydrazine, the spectrum for the r-SWCNH-ox-1.5 shows a strong and broad band in the 3500–3800 cm⁻¹ region.⁷⁵ The maximum falls near 3650 cm⁻¹, and the band might be assigned to N-H asymmetric stretching vibrations.74 Two additional broad peaks are observed. The first peak is found in the 2100-1830 cm⁻¹ region with two maxima at 1980 and 1860 cm⁻¹. These peaks might be assigned to C-NH₃⁺ and C=O stretching vibrations of anhydride or cyclic acid anhydride.74 In the 1050–900 cm^{-1} region with a maximum near 970 cm^{-1} , the peak observed might be assigned to C-C-C bending vibrations or C-O-C symmetric stretching vibrations.74

The Raman spectra of the pristine and the oxidized SWCNH samples registered at 532 nm are shown in Fig. 1H and S6† and are summarized in Table 2. The most distinctive signal at approximately 1580 cm⁻¹ is known as the G band, which corresponds to the in-plane optical mode of vibration for two adjacent sp² C atoms on an ideal hexagonal ring of graphite.⁷⁸⁻⁸⁰ The spectra are also dominated by the *D* band at 1332 cm⁻¹.^{81,82} The presence of the D band is due to defects in the carbon crystalline curved structure. Additional combined tones for the peaks are located at 2665 cm⁻¹ (2D) and 2919 cm⁻¹ (D + G) (Fig. S6†) that are observed for all investigated materials and correspond to the second-order Raman spectrum.⁸³ The D band

for all oxidized SWCNH materials and the reduced sample is shifted compared with that of the pristine SWCNHs (from 1332 to 1338 cm⁻¹). The origin and dispersive behavior of the D band can be explained by the double resonance Raman mechanism⁸⁴ that was also observed for nanographite.⁸⁵

The ratio of the intensities of the D and G bands (I_D/I_G) is used as a convenient measure of crystalline order and could confirm the functionalization process of the pristine SWCNHs (Table 2). The effectiveness of the functionalization of CNs increased, and the heights of the "disorder" deconvoluted peaks (D', D) also increased as the G peak decreased.⁸⁶ Based on this relationship, the Raman spectrum for the pristine SWCNHs shows the lowest D/G intensity ratio $(I_D/I_G = 1.2)$, and the highest I_D/I_G value was determined for SWCNH-ox-3 (I_D/I_G = 1.68). Oxidation with strong acids causes the creation of pores with different dimensionality. Therefore, the hybridization of the C atoms changes from sp^2 to sp^3 , which is recognized as formation of the defect in the CN structure.87 The larger D band intensity is also caused by the presence of oxygen functional groups on the SWCNH surface that are recognized as a defect in the structure of carbonaceous material. It must also be noted that the G band in the SWCNH-ox spectra exhibits an apparent shoulder, which corresponds to the D' band, a second-order band, at $\sim 1604 \text{ cm}^{-1}$. This phenomenon is commonly associated with disorder in graphite and could be associated with oxidation treatment of CNs (Fig. S6[†]).^{86,88}

2.2. Electrochemical performance of SWCNH electrodes

To explore the potential application of these materials in energy storage devices, the samples were used as supercapacitor electrodes and characterized using cyclic voltammetry (CV) in 0.1 mol dm⁻³ aqueous solution. Fig. 3A shows a comparison of the electrochemical properties of the glassy carbon electrode (GCE) modified with SWCNH, SWCNH-ox-1 and r-SWCNH-ox-1 layers. Modification of the electrode was performed by the drop-casting method using an ethanol dispersion of SWCNH, SWCNH-ox-1 or r-SWCNH-ox-1 with the same concentration of CNs (please see Experimental section, where modification of the GCE is described in the details).

The potential was cycled between -0.2 and +0.6 V vs. Ag/AgCl without any noticeable change in the shape of the CVs. The electrodes modified with SWCNH and r-SWCNH-ox-1 exhibit CV curves with the typical capacitance behaviour and almost pseudorectangular anodic and cathodic profiles. The symmetry of both CVs indicates that the capacitance of the electrodes is reversible in the anodic and cathodic directions.

A small departure from the ideal rectangular shape is observed for the oxidized SWCNH materials (Fig. 3). The broad and poorly defined peaks Ox and Red were recorded on CVs in anodic and cathodic cycles, respectively (Fig. 3A, red line). The difference between the observed capacitances of the nonmodified SWCNH electrodes and their oxygen-containing derivatives could be due to the more microporous structure of SWCNH-ox and the pseudocapacitive faradaic reactions (reversible oxidation/reduction) of oxygen functionalities (mainly C–OH, C=O and COOH) generated during chemical

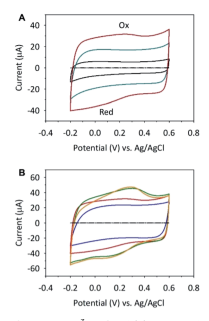


Fig. 3 CVs in 0.1 mol dm⁻³ NaCl of (A) nonmodified GCE (black dashed line) and covered with SWCNH (black line). SWCNH-ox-1 (red line) or r-SWCNH-ox-1 (blue line); and (B) nonmodified GCE (black dashed line) and SWCNH-ox prepared with different oxidation time: SWCNH-ox-0.5 (blue line), SWCNH-ox-1 (red line), SWCNH-ox-1.5 (green line), and SWCNH-ox-3 (orange line). The sweep rate was 10 mV s^{-1}

oxidation.89,90 The characteristic weak shoulders in both cycles is characteristic of the oxidized CNs due to the low redox kinetics of the surface functional groups compared with the charging kinetics of the electrochemical double layer.89,91 This effect also depends on the small fraction of oxygen-containing groups relative to the whole surface of the CNs. Moreover, all CVs are stable for these films, which proves their electrochemical stability in the studied potential range.89,91

A previous study showed that an increase in surface oxides on the CNs,92 which are used as an electrode material, results in increasing intensities of both redox processes (Fig. 3B). The amount of oxygen-containing groups on the oxidized SWCNH surface depends on the oxidation time of this nanostructure (please see the spectroscopic results in Fig. 1G). A previous study also showed that surface oxides prefer proton adsorption in the micropores of the carbon materials.89,91 Therefore, the specific adsorption results in local changes to the charge density in the double-layer capacitance. The pseudocapacitive effect is observed for all SWCNH-ox (Fig. 3B).

The specific capacitances (C_s) of the SWCNH and their oxidized and reduced forms (SWCNH-ox and r-SWCNH-ox, respectively) were determined using the CV method based on the following eqn (1):

$$C_{\rm s} = \frac{\int_{E_2}^{E_1} i(E) dE}{2vm(E_1 - E_2)}$$
(1)

where E_1 and E_2 are the respective initial and final potentials (V), $\int_{E_2}^{E_1} i(E) dE$ is the integrated current in the potential window, ν is the sweep rate (V s⁻¹), and *m* is the mass of the active material. The calculated C_s are summarized in Table 3. A clear correlation

exists between the concentration of the oxygen functionalities in the SWCNH structure and the C_s values (Fig. 3B). The first interesting observation is an enhancement of the C_s of the SWCNH materials, from 65 F g^{-1} found for the pristine SWCNH to 180 F g^{-1} for the oxidized SWCNHs (SWCNH-ox-1.5 in acidic media, H_2SO_4). This value is in good agreement with a previous report on the oxidized SWCNHs exhibiting higher capacitance than their pristine form due to modification of the micropore surface area and micropore volume due to hole-opening of the SWCNHs.^{26,29,31,44}

Generally, increasing the S_{BET} value of materials leads to an increasing overall capacitance due to a higher electrode/ electrolyte contact area (Table 3). However, it should be noted that the number of pores and also their size (micro and mesopores, and outer surfaces of the particles) affect the capacity of the layers (S_{micro} and S_{meso}) (Table 1). These parameters determine the possibility of ion mobility in the electrode layer and, consequently, the formation of electric double-layer capacitance.13,93 In addition, the method of measuring the electrode surface has an influence on the capacitance of porous carbons,92 considering the Cs relative to the morphology of the carbon material and surface chemistry (e.g., wettability and type of oxygen-containing group). As presented above, the capacitance of materials is a highly complex process, and therefore, for each newly obtained or modified CN, the structure and surface properties must be analyzed in detail in relation to many parameters.

Even for SWCNHs oxidized via the heat-treatment method,^{26,27,29,31,44} for which the literature is guite scarce, we find many discrepancies and no simple relationship between S_{BET} and C_s. For example, oxidation of the SWCNHs via heattreatment in an air atmosphere at different temperatures in the range of 400 to 580 °C results in an increasing specific surface area, from 400 to 1420 m² g⁻¹ (limit values for the pristine SWCNHs and the oxidized SWCNHs, respectively), and consequently leads to increasing C_s , from 66 to 114 F g⁻¹.³¹ Using the same type of modification (at 550 °C), other authors have produced materials with the following parameters: S_{BET} of 1690 m² g⁻¹ and C_s of 100 F g⁻¹.²⁶ However, Sano *et al.* used heat treatment of pristine SWCNHs in an air atmosphere at the same temperature (550 °C), and they obtained an oxidized material with a much lower S_{BET} value of 964 m² g⁻¹, although the C_{s} was quite high 365 F g^{-1} (at sweep rates of 10 mV s^{-1}). As noted

Table 3 Specific capacitance of SWCNH, SWCNH-ox-1.5 and r-SWCNH-ox-1.5 in aqueous solutions of different supporting electrolytes

	Specific capacitance ^{a} (F g ^{-1})					
Materials	NaCl	Na ₂ SO ₄	H_2SO_4	(Et) ₄ NCl		
SWCNH	32	52	65	28		
SWCNH-ox-1.5	103	125	180	52		
r-SWCNH-ox-1.5	95	102	134	96		

^{*a*} Based on eqn (1); $\Delta E = 300$ mV; m = 0.018 g; $\nu = 10$ mV s⁻¹; concentration of supporting electrolyte was 0.1 mol dm⁻³.

Paper

many times, no simple correlation is found between the S_{BET} and C_{s} of the CNs because the values presented above show that the same method of oxygenation and nearly the same experimental conditions can lead to different electrochemical properties of the materials. Although there is a general tendency that more efficient oxidation of CNs leads to an increase of S_{BET} and consequently an increase in the C_{s} of SWCNHs, the resulting C_{s} values differ quite significantly. It seems that a key to increasing C_{s} , as Kaneko reports, is the type and the size of the pores created in the SWCNHs during the oxidation treatment, known as 'tip opening' and/or 'sidewall opening'.^{29,31}

For the first time, we used acid treatment for oxidation of SWCNHs and hole-creation in their structure. Because the pore size of an electrode material should be suitable for the ion size of the electrolyte, a systematic study of the effects of SWCNH porosity on the ion-electrode interactions was performed.

Additionally, the surface area of porous carbons are measured by gas adsorption, which can penetrate the open pores that are available for the molecular size of the adsorbate. Electrolyte penetration into pores is more restricted,⁹² and thus a correlation between the electrolyte accessibility and carbon structure and their surface properties was studied.

Herein, the effect of the supporting electrolyte on the capacitive properties of the GCE modified with SWCNH (Fig. 4, panel (I)), SWCNH-ox-1.5 (Fig. 4, panel (II)) or r-SWCNH-ox-1.5 (Fig. 4, panel (III)) films was studied. This investigation was performed in an aqueous solution containing different supporting electrolytes of H_2SO_4 , Na_2SO_4 , NaCl or $(Et)_4NCl$. The C_s values were determined using eqn (1) and are summarized in Table 3.

There are two effects responsible for the influence of the supporting electrolyte on C_s , the degree of the film penetration of the counter ions and the structure of the double layer on the CNO surface and electrolyte solution.²⁴ A few parameters might have an influence on these effects: the hydrated radius of the ions (r_h), the chemistry of the CN surface, the number of pores

in the CNs and their size. According to Stokes' law, the migration velocity is inversely proportional to the effective $r_{\rm h}$.⁹⁴ The Stokes radius (r_s) , r_h and standard molar enthalpy of hydration $(\Delta_{hvd}H^{\circ})$ for the ions of the supporting electrolytes are summarized in Table S2.[†] However, the size of $r_{\rm h}$ is almost the same, and therefore, the supporting electrolyte affected the electrochemical properties of the films (Table 3). The SWCNHox might interact more strongly with cations than with anions. Additionally, the interaction of water molecules with the oxidized surface of CNs can influence the wettability of the carbon surface in aqueous solution and affect their electrochemical properties. The observed value of the C_s of r-SWCNHox-1.5 is higher than for unmodified nanostructures in all supporting electrolytes (Fig. 4, panel (III)), but the tendency of the supporting electrolyte to influence the electrochemical properties remains as in the oxidized form of SWCNHs. However, it should be emphasized, that reversible oxidation and reduction of oxygen functionalities generated during chemical oxidation are not visible on CVs of r-SWCNH-ox-1.5. Therefore, it may be assumed, that the chemical reduction of SWCNH-ox-1.5 with the use of hydrazine led to the reduction of carboxylic groups, creating quinone or alcoholic groups. The higher values of C_s of the reduced nanostructures (r-SWCNH-ox-1.5) compared to SWCNHs may also result from incomplete reduction of SWCNH-ox-1.5. It can be assumed that the surface of CNs has not been completely reduced and is still hydrophilic in nature (Table 3).

To investigate the rate performance of the CNs, the electrodes were cycled between -0.2 and 0.6 V (*vs.* Ag/AgCl) at different potential sweep rates. Fig. 5 and S7† show the CV curves of the SWCNH (Fig. 5) and SWCNH-ox-1.5 (Fig. S7†) layers on the GCE surface in an aqueous solution of 0.1 mol dm⁻³ in the different supporting electrolytes recorded at different sweep rates.

The CV curves were obtained for each nanomaterial at 5, 10, 15, 30, 50, 75 and 100 mV s⁻¹, showing an increase in the

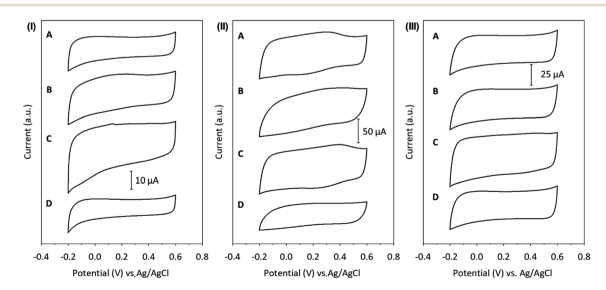


Fig. 4 CVs of (I) SWCNH, (II) SWCNH-ox-1.5 and (III) r-SWCNH-ox-1.5 films in 0.1 M (A) NaCl, (B) Na₂SO₄, (C) H₂SO₄, and (D) (Et)₄NCl. The sweep rate was 10 mV s⁻¹.



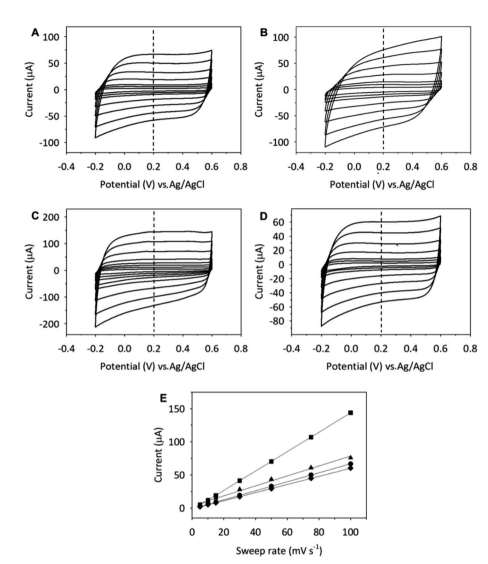


Fig. 5 CVs of SWCNH in different supporting electrolytes (0.1 M): (A) NaCl, (B) Na₂SO₄, (C) H₂SO₄, and (D) (Et)₄NCl at different sweep rates: 5, 10, 15, 30, 50, 75, and 100 mV s⁻¹. (E) Dependence of the current of SWCNH at 0.2 V (vs. Ag/AgCl) for (\bigcirc) NaCl, (\blacktriangle) Na₂SO₄, (\blacksquare) H₂SO₄, and (\diamondsuit) (Et)₄NCl.

current peak with increasing v. A linear relationship between the capacitive current (*I*) and the sweep rate (v) is observed for all forms of CNs (Fig. 5 and S7†). In this case, the capacitive current is expressed by the following equation:

$$I = C_{\rm s} vm \tag{2}$$

where the mass of active material deposited on the electrode surface was 0.018 g and other values are as explained above.

Given the existence of a linear correlation between the current peak and the scan rate, the $C_{\rm s}$ can be calculated from the slope of the linear fit using eqn (2).⁹⁵ The $C_{\rm s}$ values are summarized in Table S3,† which compares these values obtained by both methods (eqn (1) and (2)). The values of $C_{\rm s}$ obtained by integration of the *I*–*E* curves are slightly higher than those calculated from the slope of the *I*– ν plots. Generally, the $C_{\rm s}$ of the SWCNH, SWCNH-ox-1.5 and r-SWCNH-ox-1.5 electrodes

calculated by these two methods in the presence of the inorganic electrolytes are highly similar.

Frequently, the presence of functional groups on the CN's surface can contribute to capacitor instability of these electrodes.⁹² Therefore, we checked the electrochemical stability of the oxidized SWCNH electrodes in two supporting electrolytes, an inert and an acidic solution. Fig. 6 shows the CVs for up to 1000 cycles of the SWCNH-ox-1.5 electrode. As shown in Fig. 6, the $C_{\rm s}$ of the SWCNH-ox-1.5 electrode showed a slightly different electrochemical stability in the two supporting electrolytes. The $C_{\rm s}$ decreases rapidly during the first 100 cycles but remained nearly constant thereafter up to 1000 cycles in H₂SO₄ solution.

In the case of NaCl solution, the $C_{\rm s}$ for this electrode decreases gradually throughout the cycling potential window. The value of $C_{\rm s}$ decreased by approximately 12% (from 230 to 202 F g⁻¹) for NaCl solution and approximately 5% (from 233 to 221 F g⁻¹) for H₂SO₄ solution after 1000 cycles. The SWCNH-ox-

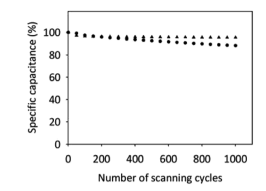


Fig. 6 Cyclic life test of SWCNH-ox-1.5 in 0.1 mol dm⁻³ NaCl (\bullet) and H₂SO₄ (\blacktriangle) measured by CVs at +0.2 V (vs. Ag/AgCl). The sweep rate was 50 mV s⁻¹.

1.5 electrode in H_2SO_4 solution was retained approximately 95% of the initial capacitive current after 1000 cycles. The study confirmed long-term cycling stability of the SWCNH-ox-1.5 electrode in acidic solution.

3 Experimental

3.1. Modification of SWCNHs

An amount of 0.5 g of SWCNH (NEC, Minato, Tokyo, Japan) was oxidized using 250 mL of 65% HNO3 (Chempur, Piekary Śląskie, Poland) at 80 °C for different times (0.5, 1, 1.5 or 3 h). The samples were labeled as SWCNH for the initial material and SWCNH-ox-y, where 'ox' denotes the oxidation and 'y' oxidation time, respectively. After the oxidation reaction, the mixture was cooled and diluted to 1000 mL, filtered under vacuum on a 0.45 µm nylon membrane (Whatman) and washed with hot distilled water until the conductivity of the mixture was no larger than 10 µS. The obtained material was collected, dispersed in distilled water, centrifuged and freeze-dried. The SWCNH-ox-1.5 was reduced. To the reduction reaction, 50 mg of SWCNH-ox-1.5 was added to 100 mL of 5% hydrazine aqueous solution. The mixture was refluxed for two hours, and in the last step, the mixture was centrifuged, washed with water until low conductivity was reached, and freeze-dried. The material was labeled as r-SWCNH-ox-1.5.

3.2. Methods

Nitrogen adsorption–desorption isotherms (77 K) were measured using an ASAP 2010 adsorption apparatus (Micromeritics, Norcross, USA). We computed the BET apparent surface areas (S_{BET}) from the classical Brunauer–Emmett–Teller⁵⁷ adsorption isotherm equation. Porosity parameters were calculated from of the kernel of local N₂ isotherms (77 K) simulated in infinitely long structureless graphene cylinders and the in-house grand canonical Monte Carlo (GCMC) simulation code. We used a single-site (12, 6) Lennard-Jones potential model ($\sigma_{\rm ff} = 0.3615$ nm and $\varepsilon_{\rm ff}/k_{\rm B} = 101.5$ K) for computing N₂–N₂ interactions. The cylindrical sections of SWNHs were modeled by a structureless graphene layers. The solid-fluid potential was computed from:

$$U_{\rm sf}(r,R) = \pi^2 \rho_{\rm s} \varepsilon_{\rm sf} \sigma_{\rm sf}^2 \Biggl\{ \frac{63}{32} \frac{F[-4.5; -4.5; 1.0; \beta^2]}{[R^*(1-\beta^2)]^{10}} - 3 \frac{F[-1.5; -1.5; 1.0; \beta^2]}{[R^*(1-\beta^2)]^4} \Biggr\}$$
(3)

where $\varepsilon_{\rm sf}/k_{\rm B} = 53.22$ K, $\sigma_{\rm sf} = 0.3494$ nm, $\beta = r/R$, $R^* = R/\sigma_{\rm sf} (\sigma_{\rm sf} = 0.3494$ nm), *r* is the radial coordinate of N₂ molecule calculated from the pore center, *R* denotes the pore radii, $\rho_{\rm s} = 38.2$ nm⁻² is the surface number density of carbon atoms in graphene, and *F* [*a*;*b*;*c*;*d*] is the hypergeometric function.^{58,59}

For GCMC simulations of N₂ adsorption in graphene cylinders, we used a periodic unit with a length of $10\sigma_{\rm ff}$. The periodic boundary conditions and the minimum image convention was used for computing N₂-N₂ interactions in the longitudinal direction. Interactions between N₂ molecules were truncated at $5\sigma_{\rm ff}$ without applying the long-range correction. The relationship between pressure and chemical potential for N₂ at 77 K was computed from canonical Monte Carlo simulations and the Widom particle insertion method.⁹⁶ At each point along the N₂ adsorption isotherm, the GCMC simulations were equilibrated using 4×10^7 Monte Carlo steps. The ensemble averages were computed from additional 2×10^7 Monte Carlo steps. In all GCMC simulations, we used an equal probability of 1/3 for the insertion, deletion and displacement perturbation step.

The absolute value of N_2 adsorption per surface area of graphene wall, S_{cvl} , was computed from:

$$n_{\rm abs}(D;h) = N(h)/S_{\rm cyl} \tag{4}$$

where $h = p/p_0$ denotes the relative pressure, D = 2R, $S_{cyl} = 2\pi RH$, *H* is the length of the graphene cylinder, and *N*(*h*) is the ensemble average of the number of N₂ molecules simulated at the selected value of *h*.

The pore size distribution function f(D) was computed from the general adsorption integral equation:

$$N_{\rm abs}(h) = \int_{D_{\rm min}}^{D_{\rm max}} n_{\rm abs}(D;h) f(D) \mathrm{d}D \tag{5}$$

where $N_{abs}(h)$ is the absolute value of N₂ adsorption inside graphene cylinder, and $D_{min} = 0.8$ nm and $D_{max} = 20$ nm are the minimum and maximum diameters of the graphene cylinder in our kernel, respectively.

The experimental value of the absolute N_2 adsorption inside SWNHs was estimated from the following equation:

$$N_{\rm abs}(h) = N_{\rm opened}(h) - N_{\rm closed}(h)$$
(6)

where $N_{\text{opened}}(h)$ and $N_{\text{closed}}(h)$ are the experimentally measured absolute values of N₂ adsorption on closed and opened SWNHs, respectively. To compute f(D) from eqn (5), we simulated 142 N₂ (77 K) adsorption isotherms in graphene cylinders with pore dimeters uniformly distributed between D_{\min} and D_{\max} (selected isotherms are collected on Fig. S1†). The Tikhonov regularization method with Sobolev seminorm implemented in the adsorption stochastic algorithm (ASA) was used in stable inversion of eqn (5).⁹⁷ The micropore surface area and micropore volume were computed by integration of the total surface and total volume to $D_{\text{max}} = 2.0 \text{ nm}$ (upper limit of the size of micropores following to IUPAC classification). Finally, the bimodal pore size distribution of the internal nanospaces of SWNH f(D) was deconvoluted using two Gaussian peaks to estimate the average size of the micro and mesopores.

For X-ray photoelectron spectroscopy (XPS), an UHV analytical system (Prevac, Rogow, Poland) equipped with a nonmonochromatic Al K (1486.7 eV) radiation source (VG Scienta SAX 100) and a monochromator (VG Scienta XM 780) was used. The Al K source was operated at 12 kV and 30 mA. A lowresolution survey scan (0–1200 eV) was performed for all samples. The pristine SWCNH and SWCNH-ox powder samples were placed on naturally oxidized Si wafers (ITME, Warsaw, Poland) that were precoated with approximately 50 nm of Au (purity 99.999%) using molecular beam epitaxy (MBE, Prevac). The measurements were performed at room temperature. The collected data were described using Casa XPS software (Casa Software Ltd., Teignmouth, USA).

The materials were also characterized using high-resolution transmission electron microscopy (HR-TEM). The images were collected using a transmission electron microscope F20X-TWIN (FEI-Tecnai, Hillsboro, USA) operated at 200 kV. The drop of sample dispersed in methanol (99.8% p.a.) was placed on a Cu grid coated with an ultrathin amorphous carbon film and dried under ambient conditions.

The ζ -potential (25 °C, Particulate Systems, NanoPlus HD, Micromeritics, Norcross, USA) was measured using solutions with a concentration of carbon materials of 0.01 mg mL⁻¹ and an ionic strength of 0.1 M. Each solution was prepared using HCl and KCl (analytical grade, Chempur, Piekary Śląskie, Poland) to adjust the ionic strength. After 24 h, the pH value was measured again, and the ζ -potential was measured.

The infrared (IR) spectra were recorded using a NICOLET IN10 MX infrared microscope (Thermo Scientific, Waltham, Massachusetts, USA). The microscope was operated primarily in reflectance mode, and the mercury–cadmium–telluride (MCT) detector was cooled with liquid N₂. A zinc selenide (ZnSe) crystal ($\theta = 45^\circ$, $n_p = 2.4$) was used in the FTIR-HATR (Fourier transform infrared spectroscopy-horizontal attenuated total reflectance) technique. The solution of the analyzed samples was placed in HATR cells, and a beam of infrared radiation entering a crystal underwent multiple internal reflections. The resultant radiation was measured and plotted as a function of wavelength. The spectral resolution was 4 cm⁻¹, and 256 scans were averaged to obtain a single spectrum.

Raman spectra were recorded using a SENTERRA (Bruker Optics, Billerica, MA, USA) micro-Raman system. Spectra were investigated in the range of 60–4500 cm⁻¹. A green laser (532 nm) was used as the excitation source, and the beam was focused on samples through a $20 \times$ microscope objective. Spectra were collected with laser powers at 2 mW to avoid sample heating and damage. Each sample was measured at least three times with a total integration time of 60 s (2 × 30 s). The obtained spectra were averaged and normalized to the G band intensity.

Voltammetric experiments were performed using an AUTO-LAB (Utrecht, The Netherlands) computerized electrochemistry

system equipped with a PGSTAT 12 potentiostat. The AUTOLAB system was controlled with Nova 2.0 software from the same manufacturer. A three-electrode cell was selected. A glassy carbon disk electrode (GCE, Bioanalytical Systems Inc. West Lafayette, USA) with a diameter of 2 mm was used as the working electrode. Before each experiment, the surface of the GCE electrode was polished using extra fine carborundum paper (Buehler) followed by 0.3 μ m Al₂O₃ and 0.25 μ m diamond polishing compounds (Metadi II, Buehler). The GCE electrode was sonicated in water to remove traces of Al₂O₃ from the metal surface, washed with water, and dried. The film on the GCE was prepared as follows. An amount of 4 mg of active material (SWCNH, oxidized or reduced form) was suspended in ethanol containing conductive carbon paint (v/v; 10:1) and sonicated for 15 minutes. An amount of 10 µL of the SWCNH dispersion was transferred to the GCE surface, and the solvent was evaporated under an Ar atmosphere. The counter electrode was a platinum flag with an area of approximately 0.5 cm². A silver wire immersed in 0.1 mol L^{-1} AgCl and separated from the working electrode by a ceramic tip (Bioanalytical Systems Inc. West Lafayette, USA) served as the reference electrode. All experiments were performed in water purified through a Millipore apparatus. Oxygen was removed from the solution by purging with argon. The studied solutions were not buffered.

4 Conclusions

Nitric acid treatment for oxidation of SWCNHs leading to hole creation in their structure was used for the first time. The obtained materials were characterized using several physicochemical methods and experimental and theoretical studies. The N₂ adsorption technique, which was supported by GCMC simulations, led to the porosity parameters. XPS, HR-TEM, FTIR and Raman spectroscopy and ζ-potential proved the successful oxidation of the SWCNH structures and their chemical composition. Electrochemical measurements were performed, supplying the correlation between the structural properties of oxidized CNs and the specific capacitance of the SWCNH electrodes. The most important conclusion from our study is that use of simple functionalization of CNs under controlled conditions leads to an almost 3-fold increase of their specific capacitance. Thus, nitric acid treatment is a simple method for enhancement of the specific capacitance of SWCNHs and facilitates application of this material for supercapacitor electrodes. Additionally, the resulting SWCNH electrode has excellent electrochemical stability, indicating the possibility of potential future applications.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank the National Science Center (NSC), Poland for financial support *via* grant number OPUS 13 UMO-2017/25/ B/ST5/00975.

References

Paper

- 1 Y. Liu, H. Zhang, P. K. Behara, X. Wang, D. Zhu, S. Ding, S. P. Ganesh, M. Dupuis, G. Wu and M. T. Swihart, *ACS Appl. Mater. Interfaces*, 2018, **10**, 42417–42426.
- 2 A. B. Papandrew, R. A. Elgammal, M. Tian, W. D. Tennyson, C. M. Rouleau, A. A. Puretzky, G. M. Veith, D. B. Geohegan and T. A. Zawodzinski, *J. Power Sources*, 2017, 337, 145–151.
- 3 D. Mirabile Gattia, M. V. Antisari, L. Giorgi, R. Marazzi, E. Piscopiello, A. Montone, S. Bellitto, S. Licoccia and E. Traversa, *J. Power Sources*, 2009, **194**, 243–251.
- 4 G. G. Wallace, J. Chen, D. Li, S. E. Moulton and J. M. Razal, *J. Mater. Chem.*, 2010, **20**, 3553.
- 5 S. L. Candelaria, Y. Shao, W. Zhou, X. Li, J. Xiao, J.-G. Zhang, Y. Wang, J. Liu, J. Li and G. Cao, *Nano Energy*, 2012, **1**, 195– 220.
- 6 W. Xu, Z. Wang, Z. Guo, Y. Liu, N. Zhou, B. Niu, Z. Shi and H. Zhang, *J. Power Sources*, 2013, 232, 193–198.
- 7 S. He, H. Hou and W. Chen, *J. Power Sources*, 2015, **280**, 678–686.
- 8 Z. Zhou and X.-F. Wu, J. Power Sources, 2014, 262, 44-49.
- 9 X. Li and B. Wei, Nano Energy, 2013, 2, 159–173.
- 10 E. Frackowiak and F. Béguin, Carbon, 2001, 39, 937-950.
- 11 S. R. S. Prabaharan, R. Vimala and Z. Zainal, *J. Power Sources*, 2006, **161**, 730–736.
- 12 S. He and W. Chen, J. Power Sources, 2014, 262, 391-400.
- 13 E. Frackowiak and F. Beguin, Carbon, 2002, 40, 1775-1787.
- 14 G. Centi and S. Perathoner, *Catal. Today*, 2010, **150**, 151–162.
- 15 Z. Zhao, S. Liu, J. Zhu, J. Xu, L. Li, Z. Huang, C. Zhang and T. Liu, ACS Appl. Mater. Interfaces, 2018, 10, 19871–19880.
- 16 Y. Yang, S. Jin, Z. Zhang, Z. Du, H. Liu, J. Yang, H. Xu and H. Ji, *ACS Appl. Mater. Interfaces*, 2017, **9**, 14180–14186.
- 17 J. Cao, C. J. Jafta, J. Gong, Q. Ran, X. Lin, R. Félix, R. G. Wilks, M. Bär, J. Yuan, M. Ballauff and Y. Lu, ACS Appl. Mater. Interfaces, 2016, 8, 29628–29636.
- 18 Y. S. Kim, J. Y. Oh, J. H. Kim, M. H. Shin, Y. C. Jeong, S. J. Sung, J. Park, S. J. Yang and C. R. Park, ACS Appl. Mater. Interfaces, 2017, 9, 17552–17564.
- 19 K. H. An, W. S. Kim, Y. S. Park, Y. C. Choi, S. M. Lee, D. C. Chung, D. J. Bae, S. C. Lim and Y. H. Lee, *Adv. Mater.*, 2001, 13, 497–500.
- 20 R. Ryoo, S. H. Joo, M. Kruk and M. Jaroniec, *Adv. Mater.*, 2001, 13, 677–681.
- M. Kruk, K. M. Kohlhaas, B. Dufour, E. B. Celer, M. Jaroniec,
 K. Matyjaszewski, R. S. Ruoff and T. Kowalewski, *Microporous Mesoporous Mater.*, 2007, 102, 178–187.
- 22 O. Barbieri, M. Hahn, A. Herzog and R. Kötz, *Carbon*, 2005, **43**, 1303–1310.
- 23 V. V. N. Obreja, *Physica E Low Dimens. Syst. Nanostruct.*, 2008, **40**, 2596–2605.
- 24 M. E. Plonska-Brzezinska, A. Lapinski, A. Z. Wilczewska,
 A. T. Dubis, A. Villalta-Cerdas, K. Winkler and
 L. Echegoyen, *Carbon*, 2011, 49, 5079–5089.
- 25 T. Azami, D. Kasuya, R. Yuge, M. Yudasaka, S. Iijima, T. Yoshitake and Y. Kubo, *J. Phys. Chem. C*, 2008, **112**, 1330–1334.

- 26 R. Yuge, T. Manako, K. Nakahara, M. Yasui, S. Iwasa and T. Yoshitake, *Carbon*, 2012, **50**, 5569–5573.
- 27 Y. Nan, B. Li, X. Song and N. Sano, *Carbon*, 2019, **142**, 150–155.
- 28 E. Bekyarova, K. Kaneko, M. Yudasaka, D. Kasuya, S. Iijima, A. Huidobro and F. Rodriguez-Reinoso, *J. Phys. Chem. B*, 2003, **107**, 4479–4484.
- 29 C.-M. Yang, Y.-J. Kim, J. Miyawaki, Y. A. Kim, M. Yudasaka, S. Iijima and K. Kaneko, *J. Phys. Chem. C*, 2015, **119**, 2935– 2940.
- 30 W. Wu, Y. Zhao, C. Wu and L. Guan, *RSC Adv.*, 2014, 4, 28636–28639.
- 31 C.-M. Yang, Y.-J. Kim, M. Endo, H. Kanoh, M. Yudasaka, S. Iijima and K. Kaneko, *J. Am. Chem. Soc.*, 2007, **129**, 20–21.
- 32 E. Bekyarova, K. Kaneko, D. Kasuya, K. Murata, M. Yudasaka and S. Iijima, *Langmuir*, 2002, **18**, 4138–4141.
- 33 S. M. Unni, R. Illathvalappil, S. N. Bhange, H. Puthenpediakkal and S. Kurungot, ACS Appl. Mater. Interfaces, 2015, 7, 24256–24264.
- 34 S. M. Unni, S. Ramadas, R. Illathvalappil, S. N. Bhange and S. Kurungot, *J. Mater. Chem. A*, 2015, **3**, 4361–4367.
- 35 E. Sánchez-Tirado, A. González-Cortés, M. Yudasaka, S. Iijima, F. Langa, P. Yáñez-Sedeño and J. M. Pingarrón, *J. Electroanal. Chem.*, 2017, **793**, 197–202.
- 36 M. Zhang, M. Yudasaka, K. Ajima, J. Miyawaki and S. Iijima, ACS Nano, 2007, 1, 265–272.
- 37 C.-M. Yang, H. Noguchi, K. Murata, M. Yudasaka, A. Hashimoto, S. Iijima and K. Kaneko, *Adv. Mater.*, 2005, 17, 866–870.
- 38 E. Aryee, A. K. Dalai and J. Adjaye, *Top. Catal.*, 2014, 57, 796– 805.
- 39 F. Agresti, S. Barison, A. Famengo, C. Pagura, L. Fedele, S. Rossi, S. Bobbo, M. Rancan and M. Fabrizio, *J. Colloid Interface Sci.*, 2018, **514**, 528–533.
- 40 S. Zhu, X.-E. Zhao, J. You, G. Xu and H. Wang, *Analyst*, 2015, **140**, 6398–6403.
- 41 M. V. Bracamonte, M. Melchionna, A. Giuliani, L. Nasi, C. Tavagnacco, M. Prato and P. Fornasiero, *Sens. Actuators, B*, 2017, 239, 923–932.
- 42 F. Valentini, E. Ciambella, A. Boaretto, G. Rizzitelli, M. Carbone, V. Conte, F. Cataldo, V. Russo, C. S. Casari, D. F. Chillura-Martino, E. Caponetti, M. Bonchio, F. Giacalone, Z. Syrgiannis and M. Prato, *Electroanalysis*, 2016, 28, 2489–2499.
- 43 F. Valentini, E. Ciambella, V. Conte, L. Sabatini,
 N. Ditaranto, F. Cataldo, G. Palleschi, M. Bonchio,
 F. Giacalone, Z. Syrgiannis and M. Prato, *Biosens. Bioelectron.*, 2014, 59, 94–98.
- 44 H. J. Jung, Y.-J. Kim, J. H. Han, M. Yudasaka, S. Iijima,
 H. Kanoh, Y. A. Kim, K. Kaneko and C.-M. Yang, *J. Phys. Chem. C*, 2013, 117, 25877–25883.
- 45 Y. Zhao, J. Li, Y. Ding and L. Guan, *Chem. Commun.*, 2011, 47, 7416–7418.
- 46 Y. Zhao, J. Li, Y. Ding and L. Guan, *RSC Adv.*, 2011, 1, 852–856.
- 47 H. Lai, J. Li, Z. Chen and Z. Huang, ACS Appl. Mater. Interfaces, 2012, 4, 2325–2328.

- 48 R. Yuge, N. Tamura, T. Manako, K. Nakano and K. Nakahara, *J. Power Sources*, 2014, **266**, 471–474.
- 49 Q.-F. Lü, S. Wang, J. Zhou, F.-F. Duan, H. Yang and R. Liu, *ChemistrySelect*, 2019, 4, 7270–7277.
- 50 H. Hwang, C. H. Kim, J.-H. Wee, J. H. Han and C.-M. Yang, *Appl. Surf. Sci.*, 2019, **489**, 708–716.
- 51 L. Lv, C. Cui, C. Liang, W. Quan, S. Wang and Z. Guo, *Food Control*, 2016, **60**, 296–301.
- 52 G. Zhang, P. He, W. Feng, S. Ding, J. Chen, L. Li, H. He, S. Zhang and F. Dong, *J. Electroanal. Chem.*, 2016, **760**, 24–31.
- 53 L. Yang, X. Ran, L. Cai, Y. Li, H. Zhao and C.-P. Li, *Biosens. Bioelectron.*, 2016, **83**, 347–352.
- 54 E. Antolini, Appl. Catal., B, 2009, 88, 1-24.

RSC Advances

- 55 L. Zhang, N. Zheng, A. Gao, C. Zhu, Z. Wang, Y. Wang, Z. Shi and Y. Liu, *J. Power Sources*, 2012, **220**, 449–454.
- 56 C. Zhu, D. Liu, Z. Chen, L. Li and T. You, *J. Colloid Interface Sci.*, 2018, **511**, 77–83.
- 57 S. Brunauer, P. H. Emmett and E. Teller, *J. Am. Chem. Soc.*, 1938, **60**, 309–319.
- 58 K. Murata, K. Kaneko, F. Kokai, K. Takahashi, M. Yudasaka and S. Iijima, *Chem. Phys. Lett.*, 2000, **331**, 14–20.
- 59 D. Wei, Y. Liu, Y. Wang, H. Zhang, L. Huang and G. Yu, *Nano Lett.*, 2009, **9**, 1752–1758.
- 60 A. Dimiev, D. V. Kosynkin, A. Sinitskii, A. Slesarev, Z. Sun and J. M. Tour, *Science*, 2011, 331, 1168–1172.
- 61 R. Blume, P. R. Kidambi, B. C. Bayer, R. S. Weatherup, Z.-J. Wang, G. Weinberg, M.-G. Willinger, M. Greiner, S. Hofmann, A. Knop-Gericke and R. Schlögl, *Phys. Chem. Chem. Phys.*, 2014, **16**, 25989–26003.
- 62 M. Koinuma, H. Tateishi, K. Hatakeyama, S. Miyamoto,
 C. Ogata, A. Funatsu, T. Taniguchi and Y. Matsumoto, *Chem. Lett.*, 2013, 42, 924–926.
- 63 Y.-C. Chiang, Y.-J. Chen and C.-Y. Wu, *Materials*, 2017, **10**, 1296.
- 64 M. K. Rabchinskii, S. A. Ryzhkov, D. A. Kirilenko, N. V. Ulin, M. V. Baidakova, V. V. Shnitov, S. I. Pavlov, R. G. Chumakov, D. Yu. Stolyarova, N. A. Besedina, A. V. Shvidchenko, D. V. Potorochin, F. Roth, D. A. Smirnov, M. V. Gudkov, M. Brzhezinskaya, O. I. Lebedev, V. P. Melnikov and P. N. Brunkov, *Sci. Rep.*, 2020, **10**, 6902–6914.
- 65 A. Barinov, O. B. Malcioğlu, S. Fabris, T. Sun, L. Gregoratti, M. Dalmiglio and M. Kiskinova, *J. Phys. Chem. C*, 2009, 113, 9009–9013.
- 66 A. Fujimoto, Y. Yamada, M. Koinuma and S. Sato, *Anal. Chem.*, 2016, **88**, 6110–6114.
- 67 B. Lesiak, L. Kövér, J. Tóth, J. Zemek, P. Jiricek, A. Kromka and N. Rangam, *Appl. Surf. Sci.*, 2018, **452**, 223–231.
- 68 C. Petit, M. Seredych and T. J. Bandosz, *J. Mater. Chem.*, 2009, 19, 9176.
- 69 E. Desimoni, G. I. Casella, A. M. Salvi, T. R. I. Cataldi and A. Morone, *Carbon*, 1992, **30**, 527–531.
- 70 N. K. Gupta, B. Peng, G. L. Haller, E. E. Ember and J. A. Lercher, ACS Catal., 2016, 6, 5843–5855.
- 71 R. Arrigo, M. E. Schuster, Z. Xie, Y. Yi, G. Wowsnick, L. L. Sun, K. E. Hermann, M. Friedrich, P. Kast,

M. Hävecker, A. Knop-Gericke and R. Schlögl, *ACS Catal.*, 2015, 5, 2740–2753.

- 72 J. Zhang, H. Zou, Q. Qing, Y. Yang, Q. Li, Z. Liu, X. Guo and Z. Du, *J. Phys. Chem. B*, 2003, **107**, 3712–3718.
- 73 J. Kastner, T. Pichler, H. Kuzmany, S. Curran, W. Blau,
 D. N. Weldon, M. Delamesiere, S. Draper and
 H. Zandbergen, *Chem. Phys. Lett.*, 1994, 221, 53–58.
- 74 T. Petit and L. Puskar, *Diamond Relat. Mater.*, 2018, **89**, 52–66.
- 75 T. Jiang and K. Xu, Carbon, 1995, 33, 1663-1671.
- 76 R. Sappok and H. P. Boehm, Carbon, 1968, 6, 283-295.
- 77 M. X. Pulikkathara, O. V. Kuznetsov and V. N. Khabashesku, *Chem. Mater.*, 2008, **20**, 2685–2695.
- 78 F. Tuinstra and J. L. Koenig, *J. Chem. Phys.*, 1970, **53**, 1126–1130.
- 79 R. Al-Jishi and G. Dresselhaus, *Phys. Rev. B*, 1982, **26**, 4514-4522.
- 80 R. J. Nemanich and S. A. Solin, *Phys. Rev. B*, 1979, **20**, 392-401.
- 81 R. Tsu, J. H. González and I. C. Hernández, *Solid State Commun.*, 1978, 27, 507–510.
- 82 V. Mochalin, S. Osswald and Y. Gogotsi, *Chem. Mater.*, 2009, **21**, 273–279.
- 83 E. D. Obraztsova, M. Fujii, S. Hayashi, V. L. Kuznetsov, Yu. V. Butenko and A. L. Chuvilin, *Carbon*, 1998, **36**, 821–826.
- 84 C. Thomsen and S. Reich, *Phys. Rev. Lett.*, 2000, **85**, 5214–5217.
- 85 L. G. Cançado, K. Takai, T. Enoki, M. Endo, Y. A. Kim, H. Mizusaki, A. Jorio, L. N. Coelho, R. Magalhães-Paniago and M. A. Pimenta, *Appl. Phys. Lett.*, 2006, 88, 163106.
- 86 D. Torres, J. L. Pinilla, R. Moliner and I. Suelves, *Carbon*, 2015, **81**, 405–417.
- 87 M. E. Plonska-Brzezinska, A. T. Dubis, A. Lapinski,
 A. Villalta-Cerdas and L. Echegoyen, *ChemPhysChem*, 2011, 12, 2659–2668.
- 88 L. M. Malard, M. A. Pimenta, G. Dresselhaus and M. S. Dresselhaus, *Phys. Rep.*, 2009, **473**, 51–87.
- 89 M. A. McArthur, N. Hordy, S. Coulombe and S. Omanovic, *Electrochim. Acta*, 2015, 162, 245–253.
- 90 D. Torres, S. Pérez-Rodríguez, D. Sebastián, J. L. Pinilla, M. J. Lázaro and I. Suelves, *Appl. Surf. Sci.*, 2020, **509**, 144774.
- 91 S. K. Park, Q. Mahmood and H. S. Park, *Nanoscale*, 2013, 5, 12304.
- 92 A. G. Pandolfo and A. F. Hollenkamp, J. Power Sources, 2006, 157, 11–27.
- 93 C. Portet, G. Yushin and Y. Gogotsi, *Carbon*, 2007, **45**, 2511–2518.
- 94 K. D. Collins and M. W. Washabaugh, *Q. Rev. Biophys.*, 1985, 18, 323–422.
- 95 M. Nasibi, M. A. Golozar and G. Rashed, *J. Power Sources*, 2012, **206**, 108–110.
- 96 B. Widom, J. Chem. Phys., 1963, 39, 2808-2812.
- 97 P. Kowalczyk, A. P. Terzyk, P. A. Gauden, R. Leboda,
 E. Szmechtig-Gauden, G. Rychlicki, Z. Ryu and H. Rong, *Carbon*, 2003, 41, 1113–1125.