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Inside the different types of carbon black as nanomodifiers for screen-printed electrodes

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Abstract

The need to deliver high performant and miniaturised electrochemical sensors has boosted the use of carbon nanomaterials as smart modifier of working electrode surface. Among the carbon nanomaterials, the common and cost-effective carbon black has recently attracted the attention from the scientific community for its outstanding features as electrode nanomodifier for analyte detection. Herein, we report the structural and morphological characterisation of several types of carbon black, namely HP 160, HS20, MTN 990, N115, N220, N375, N660, PL6, Super P, and XE2B, by means of Raman spectroscopy, transmission electron microscopy (TEM), and scanning electron microscopy (SEM) analyses. Furthermore, the electrochemical characterisation of screen-printed electrodes modified with these types of carbon black was carried out by cyclic voltammetry and electrochemical impedance spectroscopy with ferro/ferricyanide as redox probe, highlighting the advantage to use carbon black as nanomodifier in respect to the bare electrode. Among several tested carbon black types, the lowest peak-to-peak separation and resistance to electron transfer values were achieved using screen-printed electrodes modified with CB N115, N375, HP 160, and PL6. The electrodes modified with these types of carbon black were successively tested in cyclic voltammetry towards epinephrine, benzoquinone, ascorbic acid, cysteine, catechol, and caffeic acid, observing a remarkable improvement of electrochemical performances in respect to the bare electrode, even when the amperometric mode was used. The results obtained demonstrated that several types of carbon black can remarkably improve the electrochemical performances of the sensors in terms of the decrease of applied potential or peak-to-peak separation, the increase of the peak intensity, and the decrease of the resistance of the electron transfer due to several key features, including nanodimensions, the onion-like carbon structure, -O-C=O presence, and the high number of defect sites.

Keywords: modified screen-printed electrodes, carbon-based nanomaterials, morphological characterisation, structural characterisation

1. Introduction

Recent advances of nanotechnology have boosted the use of nanomaterials as smart components in the design of electrochemical sensors thanks to their countless features, including high surface area, large number of defect sites, easiness of functionalisation, and fast electron transfer. Among carbon-based materials, the discovery of carbon nanotubes in 1991 by Ijima [1] laid out one of the main pillar in nanomaterial field, demonstrating how the carbon materials at nanoscale largely affect several characteristics of electrochemical sensors such as sensitivity, selectivity, tailorability, and robustness.

In 2004, Konstantin S. Novoselov and Andre K. Geim highlighted the features of a thick monocrystalline graphitic film, namely graphene, obtained from pyrolytic graphite by an easy exfoliation technique [2]. This thick monocrystalline graphitic film turned out in one of most promising nanomaterial in several fields including the electrochemical sensor one [3, 4]. As highlighted in our recent review focused on graphene-based screen-printed electrodes (SPEs), graphene could be seen as a *kaleidoscope* because it confers different features to the sensing tools, being used to tailor the electrochemical properties of printed electrodes as well as to act as a label and loading agent for biomolecules and inorganic nanomaterials [5]. **With the aim of employing** sustainable and cost-effective nanomaterials, since 2010 we have started to use carbon black as nanomodifier for printed sensors, taking into account that few applications were reported in literature until 2010 for analyte detection in solution [6, 7]. Carbon black (CB) is an amorphous carbon belonging to pure soot and can be produced by incomplete combustion of carbon e.g. pyrolysis of carbonaceous industrial waste material. The most relevant production method is the oil furnace process, involving the partial combustion of aromatic liquid hydrocarbons in a furnace, delivering fine black particles with diameters in the range of nm, which form fused aggregates at μm level. After 2010, some works have demonstrated the fascinating properties of CB as nanomaterial to **provide** high performing electrochemical sensors. Our group reported the attractive electrochemical properties of CB N220 to be used both in the ink/paste as well as in modifying

1 electrodes by drop-casting for the quantification of several analytes such as phenolic compounds,
2 NADH, thiols, ascorbic acid [8-11]. Its interesting features as nanomodifier for working electrode
3 surface have been successfully demonstrated when compared with the most famous multiwalled
4 carbon nanotubes [12, 13] or graphene [13, 14]. The enhancement in terms of electron transfer,
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6 sensitivity to the detected analyte, resistance to fouling, and decrease of applied potential for
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8 analyte quantification has been observed by using different types of CB, such as N220 [8-11],
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10 M1100 [12], 5358R [14], VXC72R [15], with several electrodes i.e. glassy carbon, carbon paste
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12 and screen-printed electrodes [16-20], demonstrating the robustness of this nanomaterial.
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14 Furthermore, additional interesting features of this nanomaterial rely on its use without any further
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16 chemical treatment (avoiding the use of strong oxidant as in the case of carbon nanotubes), the cost-
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18 effectiveness (around 1 € for 1 Kg), and the capability to obtain a suitable dispersion useful to
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20 modify the electrode by an easily scalable drop-casting method. Although the outstanding
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22 electroanalytical properties have been mentioned and demonstrated by several research groups for a
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24 wide range of analytes e.g. amino acids [21], organic acid and DNA [22], dopamine and catechol
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26 [23], there is still a lack in the understanding the key features for its electrochemical behaviour. To
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28 address this issue, herein we report an electrochemical study of several types of carbon black
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30 namely HP 160, HS20, MTN 990, N115, N375, N660, PL6, Super P, and XE2B by testing several
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32 species i.e. epinephrine, benzoquinone, ascorbic acid, cysteine, catechol, and caffeic acid in cyclic
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34 voltammetry. Moreover, a morphological and structural study has been combined with, to
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36 investigate the possible correlation between the CB structure and electrochemical behaviour of CB-
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38 based printed electrodes.
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53 2. Experimental section

54 2.1. Materials

55 Potassium ferrocyanide ($K_4Fe(CN)_6$), potassium ferricyanide ($K_3Fe(CN)_6$),
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57 hexaammineruthenium(III) chloride, sodium phosphate dibasic (Na_2HPO_4), epinephrine,
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benzoquinone, ascorbic acid, cysteine, catechol, and caffeic acid, dimethylformamide acid were purchased from Sigma-Aldrich (St. Louise, USA). CB N220, N115, N375, and N660 were obtained from Cabot Corporation (Ravenna, Italy). HP 160 and HS20 were purchased by Orion Engineered Carbons (Kingwood, USA). MTN 990, PL6, Super P, and XE2B were kindly supplied by Prof. Franco Cataldo.

2.2. Screen-printed electrode fabrication

SPEs were produced using a 245 DEK (Weymouth, UK) screen-printing machine. Flexible polyester film (Autostat HT5), purchased from Autotype Italia (Milan, Italy), was used as substrate to print the electrodes. Graphite-based ink (Electrodag 423 SS) from Acheson (Milan, Italy) was used to print both the working and auxiliary electrodes, while silver/silver chloride ink (Electrodag 6038 SS) to print the pseudo-reference electrode, and Argon Carbonflex 25.101S for the insulating layer to define the working electrode surface area. The resultant diameter of the working electrode was 0.3 cm with a geometric area equal to 0.07 cm².

2.3 Procedure of CB dispersion

The powder of each CB kind was dispersed in a mixture of dimethylformamide/water in ratio 1:1 v/v for a final concentration of 1 mg/ml. In detail, 10 mg of each CB were first dipped in 5 ml of dimethylformamide and then 5 ml of water were added. The dispersion was sonicated for 60 minutes at 59 kHz.

2.4. Procedure for SPE modification

SPEs were modified using drop-casting method by adding on the surface of working electrode 6 μL of the dispersion via three successive steps of 2 μL. The modified SPEs were tested after the evaporation of the solvent used for the dispersion.

2.5. Electrochemical study

Cyclic voltammetry (CV), amperometry and Electrochemical Impedance Spectroscopy (EIS) were performed by using the portable potentiostat PalmSens⁴ (Palm Instrument, The Netherlands) connected to a laptop and controlled by PSTrace software. CV and EIS measurements were carried out using a solution of 5 mM ferro/ferricyanide (1:1 v/v) in KCl 0.1 mM. In the case of EIS measurements, a frequency range between 1 Hz and 10 KHz, 10 frequencies per decade, at an open circuit potential, with a voltage amplitude of 0.01 V, was used while data fitting was provided by Z-View software (Scribner Associates Inc.). Cyclic voltammetry was also used as technique to test epinephrine, benzoquinone, ascorbic acid, cysteine, catechol, and caffeic acid at concentration of 2 mM in phosphate buffer 50 mM + KCl 100 mM, pH 7.4.

2.6 Microstructural characterisation

Raman analyses were performed to investigate the structure and defectiveness of the CB samples. The Raman spectra were acquired by a Renishaw in Via Reflex Raman microscope (Renishaw plc, Wotton-under-Edge, UK) equipped with a 785 nm solid state laser source. The radiation was focused through a 20x objective and collected in backscattering configuration, using a laser power of 10 mW and an acquisition time of 50 s. To further reduce the power density, a defocusing level of 70% was selected. Each CB sample was analysed at three different positions and the obtained spectra were fitted and deconvoluted according to the 5 curves method proposed by Sadezky et al. [24] by means of the Raman software WiRE 5.1. The I_D/I_G ratio was calculated starting from the intensities of the D and G curves obtained by the deconvolution procedure.

X-ray photoelectron spectroscopy (XPS) studies were carried out with a PHI 5000 Versaprobe Scanning X-ray photoelectron spectrometer (monochromatic Al K-alpha X-ray source with 1486.6 eV energy) in order to investigate surface chemical composition. A spot size of 100 μm was used to collect the photoelectron signal for both the survey and the high-resolution (HR) spectra. All samples were analysed using a combined electron and argon ion gun neutralizer system to reduce

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the charging effect during the measurements. Spectra were analysed using Multipak 9.0 dedicated software. All core-level peak energies were referenced to C1s peak at 284.5 eV and the background contribution in HR scans has been subtracted by means of a Shirley function [25].

The nanoparticles size and shape were investigated with transmission electron microscopy. The powders were dispersed in ethanol and a drop of dispersion was put on a standard holey carbon copper grid. The measurements were carried out with a FEI TECNAI F20ST operating at 200 KV.

The deposition and distribution of the used CB dispersion on the surface of the SPE working electrode were analysed at field emission scanning electron microscopy (FEG-SEM, Leo Supra 35, UK).

3. Results and Discussion

3.1. Structural and Morphological characterisation

TEM analyses were conducted on the different CBs with the aim to evaluate their morphological and structural properties. In Figure 1 two magnifications for each type of CB are reported. All the samples exhibit an onion-like carbon structure [26, 27]. In details, XE2B powder is composed of hollow polyhedral carbon onions with few (4-8) layers (Fig. 1 L), observing that the layer distance fits that of graphene layers and particles size is in the 10-20 nm range. PL6 and Super P (Fig. 1 H and Fig. 1 I) powders show the same structural characteristics: they consist of multi-layered carbon onion (10-30 layers), as the predominant component, with an amorphous core and the particle size in the range of 20-50 nm. The not regular distance of the layers can be attributed to a reduced degree of graphitization [28]. Sample HP 160 (Fig. 1 A) presents of carbon onions with size in the range of 10-25 nm, composed of tenths of graphitic-like layers, with a very small inner amorphous core and with very irregular interplanar distances. Similar features can be found in samples N115 (Fig. 1 D), N220 (Fig. 1 E) and N375 (Fig. 1 F), and with carbon onions size in the range of 20-40 nm. Samples HS20 (Fig. 1 B), MNT 990 (Fig. 1 C) and N660 (Fig. 1 G) show the biggest particle size, with almost round carbon onion in the 50 nm range for HS20 and N660, and very big round

1 carbon onion in the range of 200-300 nm for MNT 990. They consist of by tenths of graphitic-like
2 layers with irregular interplanar distances. Except for the sample XE2B, all the samples show a
3 remarked content of defects given by the not complete graphitization of the carbon onions. The
4 correlation of these observations with Raman spectroscopy is of paramount importance to confirm
5 and to also establish the role of these defects in the different types of CB.
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11 (Figure 1)

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14 Indeed, in carbon-based electrodes the catalytic activity and electron transfer principally happens at
15 the surface defect sites [29], and Raman spectroscopy is an useful technique to evaluate the disorder
16 and imperfection of the carbon materials e.g. basal plane defects. The Raman spectra of the CB
17 samples were therefore collected and reported in Figure 2.
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24 (Figure 2)

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26 All the spectra are characterised by the presence of two main vibrational features around 1580 and
27 1350 cm^{-1} . The first band corresponds to the E_{2g} mode of graphite (G mode), due to the in-plane
28 vibration of the carbon-carbon bonds of the graphitic layers, while the second one (D mode),
29 assigned to the breathing of the 6-membered rings, only arises in the presence of defects which
30 disrupt the symmetry of the sp^2 systems [30, 31]. Three additional disorder-induced bands are
31 identified: the D' band around 1615 cm^{-1} , the D3 band around 1500 cm^{-1} , associated to the presence
32 of amorphous carbon, and the usually weak D4 band at about 1200 cm^{-1} . Their presence is pointed
33 out by the deconvolution procedure of the first order Raman spectra of the CB samples according to
34 the procedure optimized by Sadezky et al. [24]. Three broad and often unresolved bands appear
35 instead in the high Raman shift range and correspond to the second order modes of the D (2D, at
36 2675 cm^{-1}) and D' (2D', at 3180 cm^{-1}) bands and to a combination mode of the D and G bands
37 (D+G, at 2915 cm^{-1}) [32]. Despite the common vibrational pattern of the CB samples, some
38 differences in the relative intensity ratios and band widths are noticed. In particular, the I_D/I_G ratio is
39 a useful parameter to evaluate the level of order/disorder in the graphitic carbon [9] and its values
40 calculated for the CB samples, are listed in Table 1. As a general trend, the nanometric CB particles
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1 display higher I_D/I_G ratios with respect to the bigger ones, probably because of the combination of a
2 greater number of defects and an increased distortion of the graphitic-like domains at the surface of
3 the small nanoparticles. Indeed, quite similar ratios, i.e. around 1.3 were detected for most of the
4 samples, except for the N660 and MTN 990 powders, whose particle size is the largest, as
5 evidenced from the related TEM images, resulting in the lowest I_D/I_G ratios (around 1.2). Instead,
6 the highest I_D/I_G was measured for the XE2B, followed by the Super P and the PL6 samples. The
7 increase of the relative intensity of the defect-induced D band for these particles is not only related
8 to their small dimension, but also to their different structural features, as highlighted by a spectral
9 profile characterized by narrow D and G bands. Indeed, it is well known that an increase of the I_D/I_G
10 ratio can be correlated to a decrease of the size of the graphitic-like domains [33]. The sharpness of
11 the D and G bands witnesses, however, a globally increased degree of crystalline order for such
12 samples, consistently with a drop of the spectral intensity between the D and G bands. Such a
13 decrease is due to a reduced contribution in this region of the D3 disorder-induced mode whose
14 intensity was proposed to be indicative of the amorphous carbon content by previous studies on
15 carbonaceous materials [32]. The greater graphitic fraction of these samples is also confirmed by an
16 analogous narrowing of the band widths, observed in the second order region.

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39 (Table 1)

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41 To establish the effect of the elemental composition of different CB samples, XPS analysis was
42 carried out. Undeniably, XPS measurements seek to obtain information regarding the chemical state
43 of each CB sample involved in this study, since electrochemical performances are deeply related
44 with materials surface chemistry. From survey spectra (Figure S1) we have obtained information
45 about elemental composition and relative atomic concentration of each present species (Table 2).
46 The O concentration varies from 9.9 at.% for sample HP160 to 0.7 for Super P sample: in between
47 the average O concentration is almost 2 at.%. Then, we focused our attention on C1s photoelectron
48 peaks, to evaluate the chemical shifts due to the C-O bonds. After a deconvolution procedure
49 applied on all the C1s HR spectra (by using a pseudo-Voigt function for each peak), we obtained
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six components for all the samples apart from HP160, who needed an extra component (C-C sp¹ at 283.5 eV) [34] to reproduce the experimental curve. The chemical shift found are: C-C sp² (284.5 eV), C-C sp³ (285.2 eV), C-O-C/H (286.0 eV), O-C-O (287.4 eV), -O-C=O- (289.2 eV) and $\pi-\pi^*$ shake up satellite (>291 eV) [35]. All the chemical shift percentage are reported in Table 3.

Finally, in order to investigate the deposition and the distribution of the used CB types on the surface of the working electrode, SEM analysis was carried out for all SPEs. From the comparison among the SEM micrographs reported in Figure 3, it is evident that, in all cases, a continuous and uniform layer of CB particles was obtained, which completely covered the underneath randomly orientated micrometric graphite flakes, evidenced in the SEM micrograph of bare SPE (Fig. 3 A, B). Indeed, the bare SPE presented graphite flakes (Fig. 3 A), characterised by the presence of numerous small particles consisting in the cross-linking agents present in the original ink (Fig. 3 B) [36]. Thus, these experimental evidences confirmed the obtainment of a good dispersion and the efficacy of the used drop-casting technique. However, in all cases it is possible to observe the presence of CB agglomerates, as expected: due to its very high specific surface area, CB primary particles spontaneously tend to generate nanostructured aggregates which can form agglomerates, leading to distinct and characteristic porous structures based on nanometric spherical particles [37]. From the reported SEM micrographs, it is evident that the samples HP 160, N115, N220, N375, PL6, Super P and XE2B appear comparable, due to the similarity of the used CB powders, composed of nanometric particles (with an average diameter size of 20-40 nm) which tend to aggregate due to their very high specific surface area, in very good agreement with TEM results (Fig. 1). On the other hand, the samples HS20, MTN 990 and N660 were characterised by bigger, more heterogeneous and more rounded particles, as evident from the higher magnifications micrographs (*insets* of Fig. 3 D, E and I, respectively), particularly in the case of MTN 990 modified SPE, being MTN 990 powder composed of heterogeneous particles with different diameter sizes (ranging between 100 nm and 300 nm). In these latter cases, the bigger dimension of the used particles led to the formation of a less uniform layer, characterised by diffuse voids (Fig. 3

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E). Accordingly, the CB layers resulted more compact and uniform in the case of smaller CB particles modified SPEs (Fig. 3 C, F, H, L, M, N), as expected.

(Figure 3)

3.2. Electrochemical characterisation using ferro/ferricyanide as redox probe

In order to evaluate the electrochemical performances of CB as nanomaterial for electrode fabrication, the most common approach relies on the modification of well-known characterised electrode materials by drop casting. To explore the response of CB-modified electrode surface in function of coverage effect, which can affect the electrochemical performances, the response of ferro/ferricyanide was evaluated after having modified the working electrode surface with 2, 4, 6 and 8 μL of different CB dispersions at concentration of 1 mg/mL. As depicted by Figure 4, for almost all CB-modified SPEs at increasing of CB amount a decrease of peak to peak separation was observed, in combination with the increase of peak intensity, due to the higher amount of CB cast on the working electrode surface. Taking into account the electrochemical performances as well as the time of modified electrode preparation, 6 μL (means 6 μg of CB) was selected as optimal volume of dispersion at 1 mg/mL to cast on the working electrode surface, in agreement with our previous work using only CB N220 [11]. After having selected the amount of CB, cyclic voltammetric measurements were carried out to obtain deeper information about the electrochemical properties of the different types of CB used as electrode modifier, except for CB N220 being previously largely characterised in our previous studies [8-11].

(Figure 4)

Figure 5 shows the voltammograms obtained in ferro/ferricyanide 5 mM + KCl 0.1 M, by using SPEs modified with the selected CBs. Different behaviours were observed depending on the CB used to modify SPEs by drop-casting method. SPE modified with MTN 990 gave the largest peak

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to peak separation (i.e. 470 mV) even when compared with bare SPE, and this behaviour is ascribed to its structure, which is not at the nanodimension level. A peak to peak separation comprised between 160 and 180 mV was observed with SPEs modified with CB N660, N375, Super P. The smallest peak to peak separations were observed in the case of SPEs modified with N115 and HP 160, being equal to 120 ± 7 and 129 ± 6 mV, respectively.

To quantify the effect of CB types on the electron transfer, the heterogeneous rate constant (k^o) for the ferro/ferricyanide redox probe was calculated based on Nicholson's theory [38], using $D_O = D_R = 5.9 \times 10^{-5}$ cm²/s and an electrochemical transfer coefficient $\alpha = 0.5$, taking into account that the I_{ap}/I_{cp} value is very close to the unity for bare and CB modified SPEs [39]. The heterogeneous rate constants, reported in Table 4, highlighted the fastest electron transfer in the case of SPEs modified with N115 and HP 160, giving a value of $(6.3 \pm 0.6) \times 10^{-3}$ and $(5.4 \pm 0.9) \times 10^{-3}$ cm/s, respectively, which are comparable to the one obtained using CB N220 as modifier (4.5×10^{-3} cm/s) using the same solvent for the dispersion and the same batch of CB [13]. In contrast, the lowest value of heterogeneous rate constant was obtained using SPEs modified with HS20 ($1.2 \pm 0.3) \times 10^{-3}$ cm/s.

Analysing the intensity of the anodic and cathodic peaks, the lowest intensities were observed using SPEs modified with CB Super P, N660, N375, HS20, MTN 990 with the anodic peak current equal to 73 ± 1 , 75 ± 6 , 73 ± 3 , 73 ± 1 , and 42 ± 7 μ A, respectively. In contrast, higher peak intensities were found using SPEs modified with HP 160, N115, PL6 and XE2B (i.e. 91 ± 6 , 86 ± 1 , 89 ± 6 , and 88 ± 2 μ A).

The enhancement of the intensity of the anodic and cathodic peaks in the case of SPEs modified with these latter CB samples is ascribed to the higher electroactive surface area calculated using Randles-Sevcik equation with the diffusion coefficient equal to $D = 5.9 \times 10^{-5}$ cm²/s [40]. Indeed, these CB-SPEs were characterised by electroactive surface areas equal to 3.8 ± 0.2 , 3.6 ± 0.2 , 3.7 ± 0.3 , and 3.7 ± 0.2 mm², respectively, higher values when compared with the other ones reported in Table 4 for the remaining CB-SPEs. In addition, as expected, the electroactive areas resulted higher than the one reported for the bare electrode, 1.6 ± 0.2 mm² calculated in our previous work [40].

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6 Furthermore, the voltammetric peak intensity for each CB-SPE was evaluated as a function of the
7 scan rate and plotted versus the square root of the scan rate ($v^{0.5}$). Figure S2 showed linear
8 behaviour for bare SPE and for each modified CB-SPE, indicating that the process is fully
9 controlled by the diffusion of the electroactive species. The following slopes ($\text{mA}/(\text{mVs}^{-1})^{0.5}$) were
10 obtained: 3.6 ± 0.2 ($R^2 = 0.983$), 13.6 ± 0.4 ($R^2 = 0.992$), 10.3 ± 0.4 ($R^2 = 0.998$), 3.6 ± 0.2 ($R^2 =$
11 0.937), 12.7 ± 0.3 ($R^2 = 0.996$), 14.9 ± 0.3 ($R^2 = 0.993$), 11.0 ± 0.4 ($R^2 = 0.990$), 12.8 ± 0.4 ($R^2 =$
12 0.981), 9.0 ± 0.4 ($R^2 = 0.987$), 10.6 ± 0.6 ($R^2 = 0.976$), respectively for bare SPE, HP 160, HS20,
13 MTN 990, N115, N375, N660, PL6, Super P, and XE2B. The highest slope was observed using HP
14 160, confirming its better electrochemical properties.
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28 Additionally, we have analysed the variation of $\log I_p$ versus $\log v$ using ferro/ferricyanide as redox
29 probe, obtaining the following gradients 0.52 ± 0.03 , 0.51 ± 0.03 , 0.52 ± 0.02 , 0.52 ± 0.03 , $0.50 \pm$
30 0.02 , 0.52 ± 0.03 , 0.54 ± 0.03 , 0.50 ± 0.03 , 0.48 ± 0.03 , respectively for HP 160, HS20, N115,
31 N220, N375, N660, PL6, Super P, and XE2B (Figure S3). The values obtained, close to 0.5,
32 suggested no thin-layer effects, ensuring the absence of the redox probe entrapped within the
33 network of the film. Therefore, a purely diffusional (semi-infinite) electrode response was observed.
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35 While in the case of bare SPE and SPE modified with MTN 990, a different gradient equal $0.37 \pm$
36 0.02 and 0.37 ± 0.03 was found, respectively, demonstrating a dissimilar behaviour in respect to
37 nanomodified electrode. This behaviour is probably ascribed to the presence of insulating zones in
38 the printed working electrode surface, being the graphite ink composed of carbonaceous particles
39 mixed with insulating polymers. In the case of SPE modified with MTN 990, the presence of MTN
40 990 allowing for a less uniform layer, characterised by diffuse voids avoiding the coverage of
41 insulating area.
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To deeply investigate the electrochemistry of the different CBs, Electrochemical Impedance Spectroscopy (EIS) was used, obtaining information on the impedance changes of the electrode surface by varying the CB modifier (Figure 6). Indeed, R_{ct} value gives the difficulty of electron transfer of ferro/ferricyanide redox probe between the solution and the electrode, and its measurement can highlight the capability of CB to improve the electron transfer to the redox probe. Using this technique, the higher R_{ct} values were observed in the case of SPEs modified with MTN 990, Super P and HS20 with R_{ct} equal to $4 \pm 2 \text{ M}\Omega$, $1.3 \pm 0.5 \text{ M}\Omega$, and $1.2 \pm 0.2 \text{ M}\Omega$, respectively, while the lowest R_{ct} using SPEs modified with HP 160, N115 and PL6 (51 ± 8 , 51 ± 6 and $55 \pm 7 \Omega$), confirming the behaviour of cyclic voltammetric experiments.

(Figure 6)

3.3. Reproducibility and storage stability of CB dispersion

One of the main advantageous key feature of CB is the facile preparation of a stable dispersion assisted by just one-step of sonication (60 minutes at 59 kHz) without any further chemical and physical treatment. The easy way to prepare a stable dispersion allows for a reproducible dispersion as well as reproducible CB-modified electrodes. The reproducibility of CB dispersion was investigated by modifying SPE with three different dispersions of each CB samples and by testing them with ferro/ferricyanide as redox probe. The results reported in Figure S4 demonstrated similar electrochemical response towards the redox probe investigated within the experimental error, highlighting the reproducibility of the different CBs dispersion, even when diverse CB samples were used. The good reproducibility proved the robustness of the dispersion procedure preparation as well as the process of CB-modified electrode production, establishing a well-known route for CB-modified electrode fabrication.

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Almost all CB samples gave a homogeneous dispersion by using just one-step of sonication, except XE2B, Super P, and PL6, which required further sonication time. The homogeneity of the dispersions using different CB samples was evaluated in the same day of preparation and after 1 week, demonstrating good storage stability for almost all CB samples tested, except in the case of XE2B, Super P, and PL6, for which a poor uniformity of the layer cast on the filter paper was observed (Figure S5). The stability of CB dispersions has been also evaluated by drop-casting the dispersion with two-week as life-time, maintained at room temperature without any additional sonication step. In Figure S4, the response of CB-modified SPEs prepared using a dispersion with 1 hour of life-time (continuous black line) or two weeks as life-time (dashed black line) sheds light a similar behavior, demonstrating the good storage stability of almost all CB dispersions.

3.4. Electrochemical behaviour of epinephrine, benzoquinone, ascorbic acid, cysteine, catechol and caffeic acid

To get a further insight into CB types that gave the best performances using ferro/ferricyanide probe (i.e. HP 160, N115, N375 and PL6), the modified SPEs were challenged towards some interesting compounds for biological and environmental applications namely epinephrine, benzoquinone, ascorbic acid, cysteine, catechol and caffeic acid. Figure 7 sheds light the improvement of all modified SPEs when compared with bare SPE (black dashed line), demonstrating the electrocatalytic properties of the different CBs tested.

For epinephrine, caffeic acid, benzoquinone, and catechol a decrease of peak to peak separation was observed together with an increase of the peak intensity. In the case of the epinephrine and caffeic acid a smaller peak to peak separation combined with higher anodic and cathodic peak intensity was observed for SPEs modified with HP 160 and N115. For instance, the cyclic voltammeteries of epinephrine using HP 160 and N115 gave a peak to peak separation equal to 422 ± 6 and 419 ± 1 mV and the anodic peak intensity equal to 111 ± 4 and $101 \pm 7 \mu\text{A}$, respectively. In the case of benzoquinone and catechol the best electrochemical performances were observed using HP 160 and

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N375 as nanomodifier in terms of peak to peak separation (i.e. 109 ± 1 and 220 ± 20 in case of benzoquinone) and peak intensities (i.e. 36 ± 2 and $28 \pm 2 \mu\text{A}$ in case of benzoquinone), respectively. For the irreversible redox reaction, as the case of ascorbic acid and cysteine, the presence of CB to modify working electrode surface largely decreased the applied potential allowing for the detection of the analyte at a lower applied potential as well as an improvement of sensitivity. In the case of ascorbic acid, the presence of the HP 160 and N115 gave the best resolution of the peaks and the lowest applied potential, while in the case of cysteine we observed a similar behaviour for all CBs tested. The achieved results demonstrated that CB positively affects the detection of the analytes decreasing the peak to peak separation in the case of (semi)reversible reaction or decreasing the applied potential, demonstrating the electrocatalytic properties of CB. In addition, in all cases we observed an increase of the sensitivity due its high surface.

(Figure 7)

3.5. Electroanalytical characterisation

SPEs modified with the most performant CBs, namely HP 160 and N115, were tested in amperometric mode toward ascorbic acid and cysteine, analytes chosen for their importance in biological field as well as for their features in electrochemistry (e.g. fouling problem, high applied potential). In addition, to compare the behaviour with the well-known N220, the amperometric analyses were performed by using also SPEs modified with this latter CB. For the amperometric detection, applied potentials equal to +50 mV and +400 mV were chosen for ascorbic acid and cysteine detection, respectively, taking into account the previous cyclic voltammetries. As depicted in Figure 8, for both ascorbic acid and cysteine detection, the presence of CB largely improves the sensitivity of the measurements in respect to the bare SPE (black line), overcoming the fouling problem, indeed successive additions of the analyte gave almost the same signal intensity. In the case of ascorbic acid detection (Fig. 8 A), higher sensitivity was obtained by using SPEs modified with N220 ($0.021 \pm 0.001 \mu\text{A}/\mu\text{M}$) when compared to the ones obtained using SPEs modified with

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HP 160 and N115 (i.e. $0.016 \pm 0.002 \mu\text{A}/\mu\text{M}$ and $0.0196 \pm 0.0003 \mu\text{A}/\mu\text{M}$). Limit of detection (calculated as $S/N = 3$) was found equal to $0.1 \mu\text{M}$, $0.4 \mu\text{M}$ and $0.3 \mu\text{M}$ for the SPEs modified respectively with N220, HP 160 and N115.

As for ascorbic acid detection, in the case of cysteine measurement, SPEs modified with CB N220 resulted in higher sensitivity, i.e. $(0.0217 \pm 0.0009) \mu\text{A}/\mu\text{M}$, compared with sensitivities obtained by using HP 160 and N115 modified SPEs, respectively equal to $(0.017 \pm 0.001) \mu\text{A}/\mu\text{M}$ and $(0.0161 \pm 0.0008) \mu\text{A}/\mu\text{M}$. The calculated LOD was $0.3 \mu\text{M}$, $1 \mu\text{M}$ and $0.6 \mu\text{M}$ for the SPEs modified respectively with N220, HP 160 and N115, confirming the better performances of N220.

(Figure 8)

3.6. Discussion

The morphological and structural characterisation combined with electrochemical studies with ferro/ferricyanide as redox probe, highlighted that the nanodimension of CB positively affects the electrochemical behaviour of the sensor when it is used as working electrode modifier. Undeniably, CB characterised by large particles gave only a smaller improvement of the electrochemical performances of electrodes: in the case micromaterial MTN 990, modified electrode with MTN 990 gave similar performances in respect to the bare electrode, confirming the advantage of the nanodimension over the microdimension. Additionally, the case of HS20 and N660, with almost round carbon onion in the 50 nm range, a higher peak to peak separation and resistance to electron transfer (R_{CT}) with lower peak intensity were observed, in respect to electrodes modified with CBs characterised by smaller nanoparticles. The results achieved demonstrated that the structure affects the electrochemical response: among the different CB samples, the ones with carbon onions size in the range of 10-25 nm, composed of tenths of graphitic-like layers with a very small inner amorphous core and very irregular interplanar distances, namely HP 160, N115, N220, and N375, provided the best electrochemical performances. As reported in literature [29], the defect sites play a crucial role in electrocatalysis. CB samples are characterised by high number of defect sites with

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I_D/I_G equal to 1.19 or higher, but the defect sites are not the only key features: XE2B has the highest I_D/I_G with smaller particles size in the 10-20 nm range, with hollow polyhedral carbon onions with few (4-8) layers, but it is not the most performant CB among the CB tested. Another relevant issue regards the presence of surface functional groups e.g. C=O, O=C-O, which highly affect the electrochemical response as demonstrated using inner-sphere probe i.e. ferro/ferricyanide. Otherwise, using no-sensitive surface electrochemical redox system i.e. hexaammineruthenium(III) chloride (outer-sphere probe) [67], the peak to peak separation is similar among the several CB types, having similar electronic structure (Table S1). Additionally, XPS analysis gave information related to the effect of C/O group on the electrochemical behavior. HP 160, which is one the most promising CB, is characterised by the higher content in O atoms (9.9 at.%), while in the remnant samples the average O content is around 2.2 at.%. Moreover, it is the only one which possesses a chemical shift in the C1s photoelectron peak, due to sp^1 hybridization (19.99%), together with sp^2 (52.34%) and sp^3 (9.07%), which are also present in the other samples. The more intense C-O bond is the one due to O-C-O chemical shift at 1287.4 eV (9.26%), while an average value for the other samples is 4.7%. Among the C/O groups, the CB with higher content of -O-C=O (i.e. N115 3.99 % and N 220 4.07 %) is characterised by better electrochemical performances, demonstrating that O content and, in detail the -O-C=O amount, can give an improvement because of particular electrochemical electrocatalytic interaction with these specific surface oxygenated groups/species.

4. Conclusions

Herein, we reported a comparison among SPEs modified with several types of CB to investigate the influence of the morphological and structural characteristic of the investigated CBs on the electrochemical behaviour. The nanodimension of CB allows for the development of highly sensitive sensor when compared with the bare one, with the advantage to be a cost-effective nanomaterial, to do not require any treatment before the use, and to easily furnish a stable dispersion needed for a reproducible and easy scalable drop-casting procedure. Several CBs were

1 identified as best candidates for the modification of the electrode surface such as HP 160, N115,
2 N220, N375 and PL6, demonstrating low peak to peak separation in the case of (semi)reversible
3 species and low applied potential in the case of irreversible compounds, always associated to an
4 increase of sensitivity. Because the well-established electrochemical properties combined with
5 advantageous features reported above, several CBs deserve to be used as nanomodifier of
6 electrochemical sensors. The reasons for these outstanding electrochemical properties are ascribed
7 to nanodimensions onion-like carbon structure (carbon onions size in the range of 10-25 nm), high
8 number of defect sites (I_D/I_G ratios equal to 1.3 or higher), higher content in O atoms (especially –
9 O-C=O groups). However, these characteristics are not the only ones able to furnish the high
10 performant electrochemical behaviour of CB-modified electrodes, opening further questions about
11 still undiscovered reasons.
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38 Figure Captions

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43 **Figure 1.** TEM images of CB HP 160 (A), HS20 (B), MTN 990 (C), N115 (D), N220 (E), N375 (F)
44 N660 (G), PL 6 (H), Super P (I), XE2B (L). (scale: 20 or 50 nm, 5 nm or 10 nm for the inset).
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51 **Figure 2.** Normalized average Raman spectra of the CB powders.
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55 **Figure 3.** SEM micrographs of the bare SPE (A-B) and of the SPEs modified with HP 160 (C),
56 HS20 (D), MTN 990 (E), N115 (F), N220 (G), N375 (H), N660 (I), PL6 (L), Super P (M), XE2B
57 (N) (insets: higher magnification micrographs).
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2 **Figure 4.** Value for anodic peak current (black dots) and peak to peak separation (red dots) for SPE
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4 modified with 2 μL , 4 μL , 6 μL and 8 μL of CB dispersion HP 160 (A), HS20 (B), MTN 990 (C),
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7 N115 (D) N220 (E), N375 (F), N660 (G), PL6 (H), Super P (I), XE2B (L).
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14 **Figure 5.** Cyclic voltammetry measurements performed in $[\text{Fe}(\text{CN})_6]^{3-/4-}$ 5 mM + KCl 0.1 M,
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16 obtained with bare SPEs (dashed black line) and SPEs modified with HP 160 (green line), HS20
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18 (dashed green line), MTN 990 (dashed red line), N115 (grey line), N375 (black line), N660 (red
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20 line), PL6 (cyan line), Super P (dashed blue line) and XE2B (blue line).
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29 **Figure 6.** Nyquist plot of EIS measurement obtained with bare SPEs (dashed black line) and
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31 modified SPEs with HS20 (dashed green line), MTN 990 (dashed red line), Super P (dashed blue
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33 line), and N660 (red line). Inset: Nyquist plot at high frequencies for EIS measurement obtained
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35 with SPEs modified with HP 160 (green line), N115 (grey line), N375 (black line), PL6 (cyan line),
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37 XE2B (blue line). Measurements performed in $[\text{Fe}(\text{CN})_6]^{3-/4-}$ 5 mM + KCl 0.1 M at open circuit
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48 **Figure 7:** Cyclic voltammograms obtained in phosphate buffer 0.05 M + KCl 0.1 M, pH 7.4, in
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50 presence of 2 mM of epinephrine (A) benzoquinone (B), ascorbic acid (C), cysteine (D), catechol
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52 (E) and caffeic acid (F). All recorded at a scan rate of 50 mV/s using bare SPEs (dashed line) and
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54 SPEs modified with HP 160 (green line), N115 (yellow line), N375 (black line), and PL6 (cyan
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Figure 8: Amperograms obtained in phosphate buffer 0.05 M + KCl 0.1 M, pH 7.4, after successive addition (25 μ M) of ascorbic acid (A) and cysteine (B), for bare SPE (black line) and modified with N220 (green line), N115 (blue line) and HP 160 (red line).

Table 1. Average D and G band positions and I_D/I_G ratio obtained by the deconvolution of the first order Raman spectra of the CB powders.

Table 2. XPS relative atomic concentration (at.%), obtained from HR spectra.

Table 3. XPS C1s chemical shifts assignments, obtained from deconvolution procedures (not reported).

Table 4. Electrochemical and structural features of several CB types.

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Figures

Figure 1

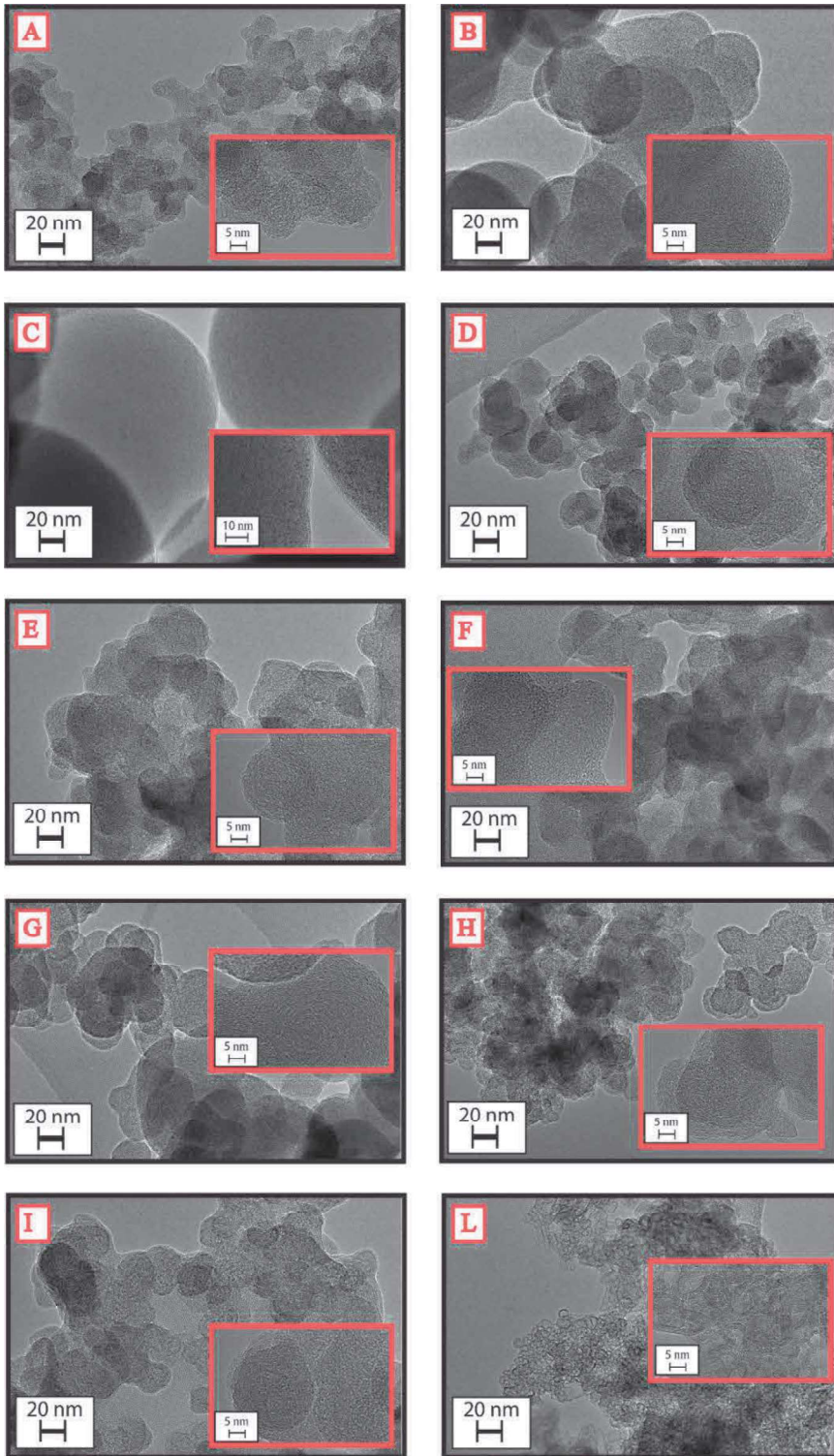


Figure 2

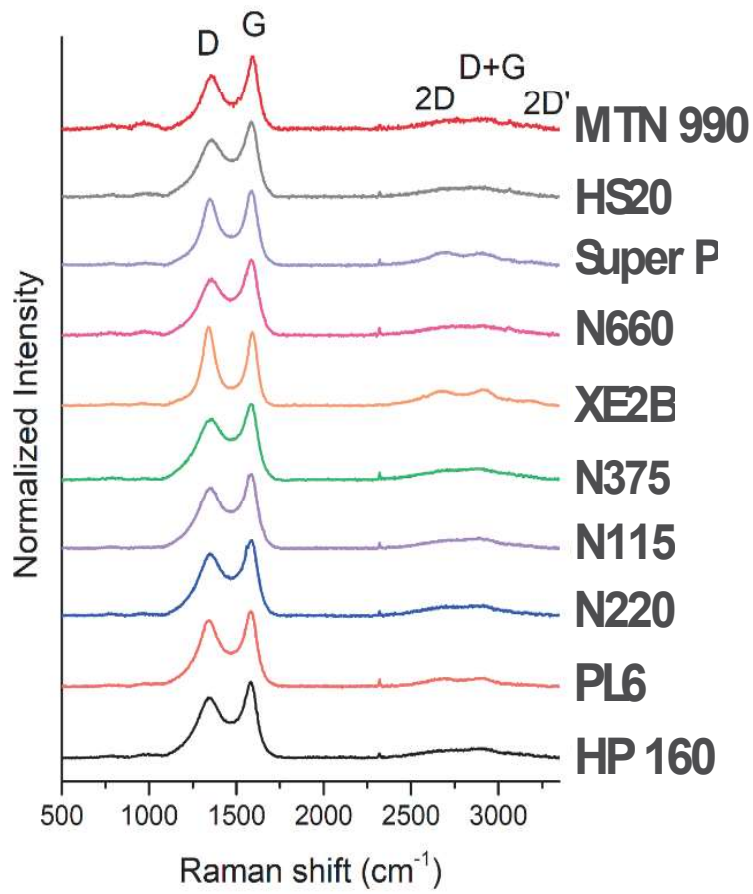


Figure 3

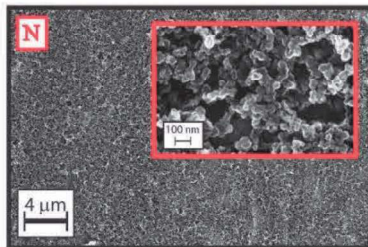
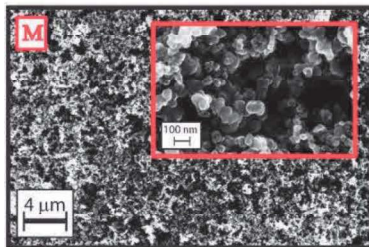
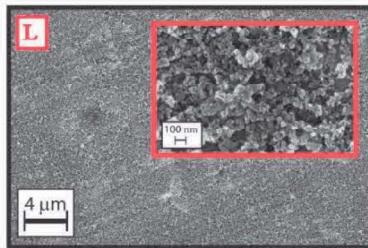
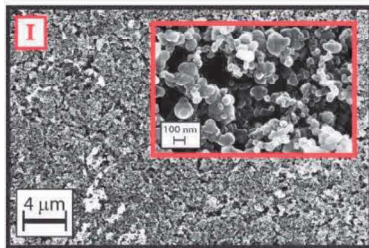
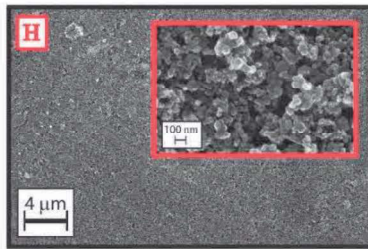
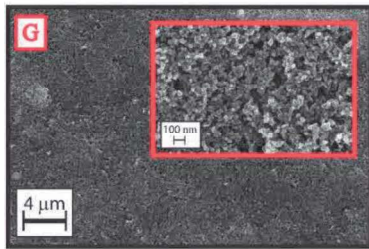
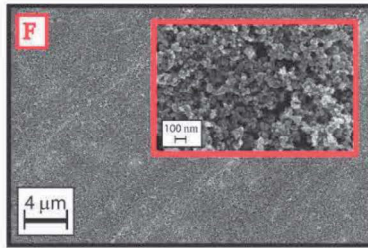
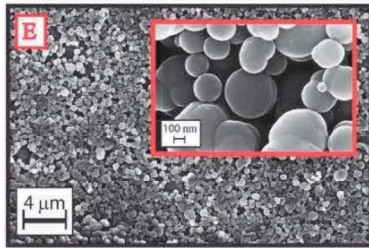
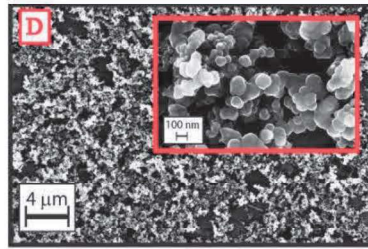
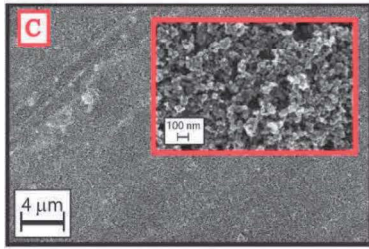
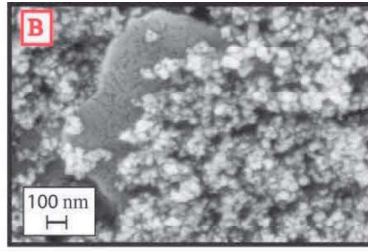
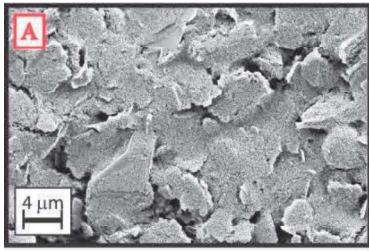


Figure 4

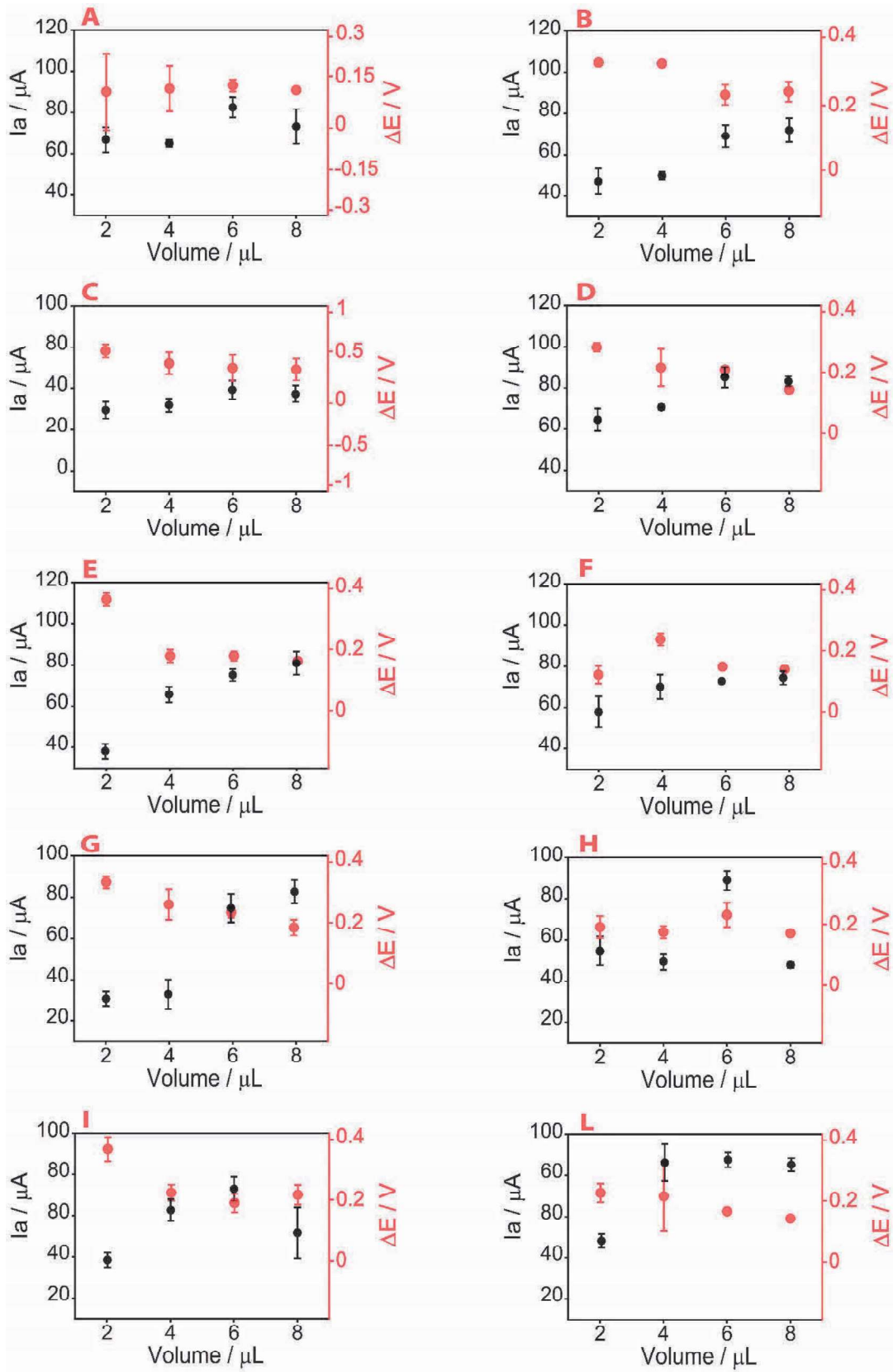


Figure 5

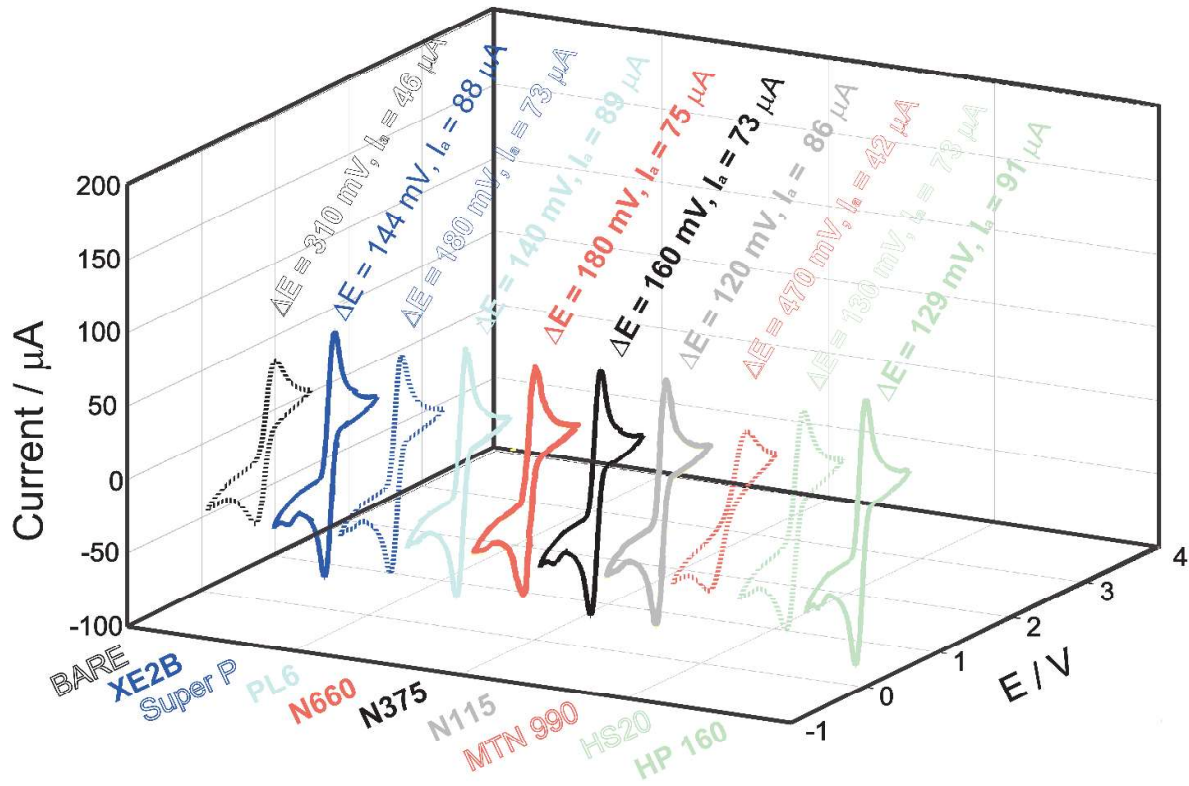


Figure 6

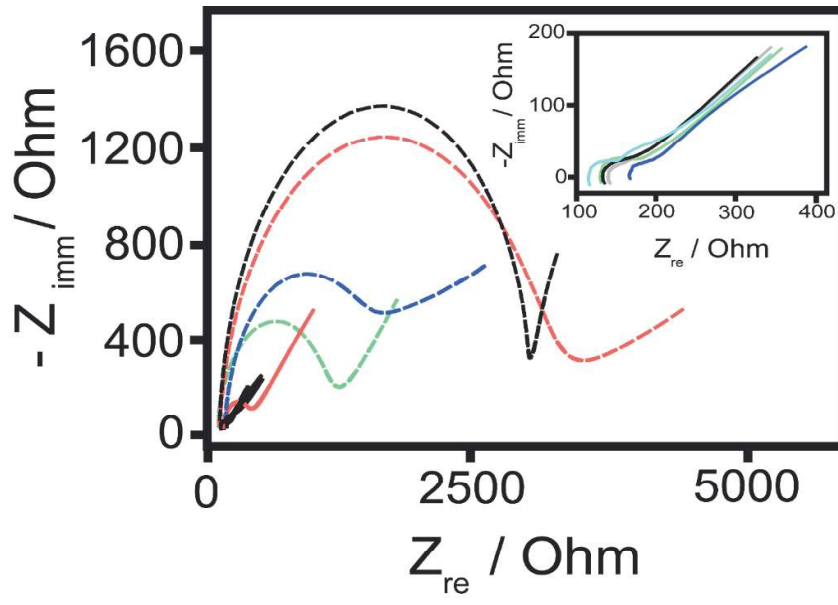


Figure 7

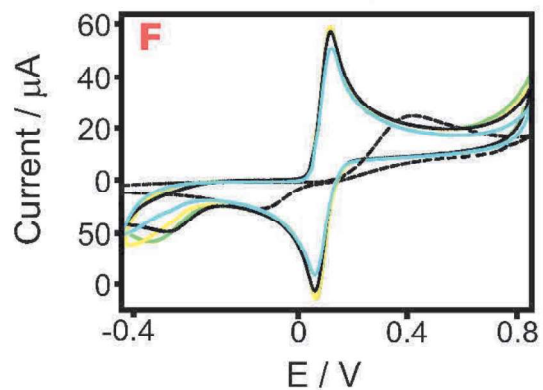
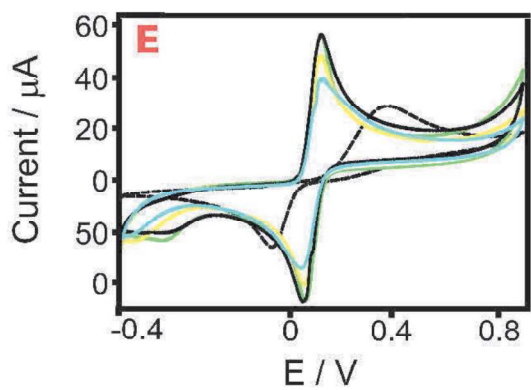
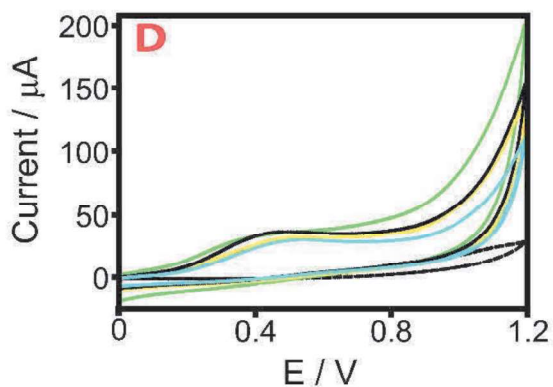
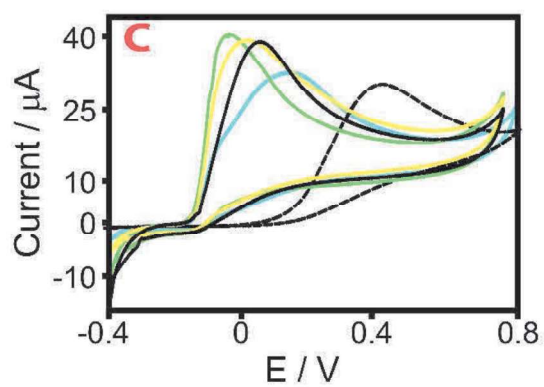
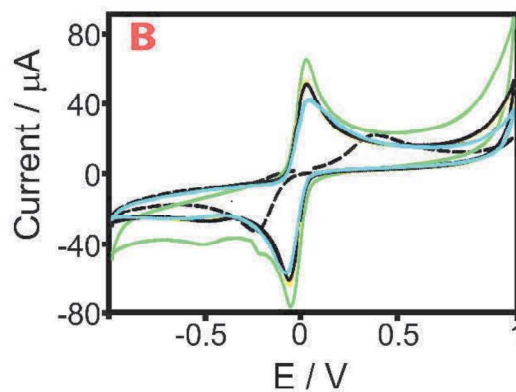
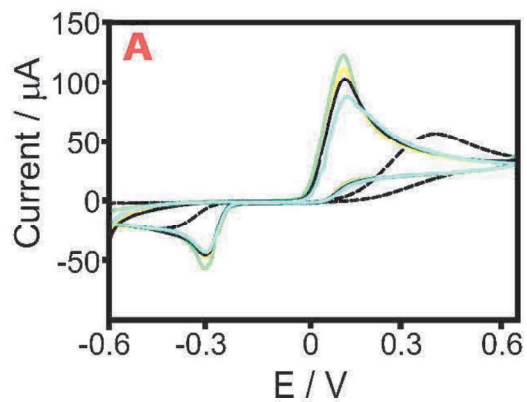
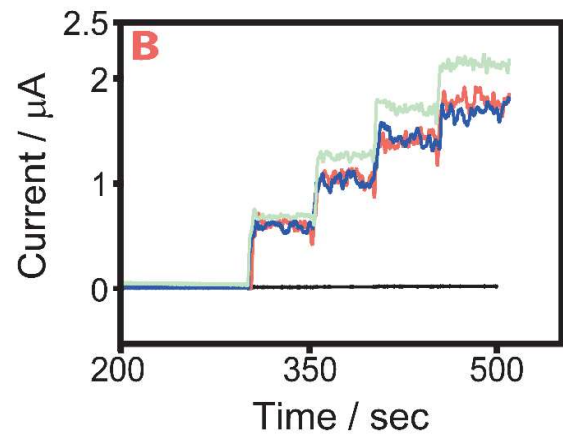
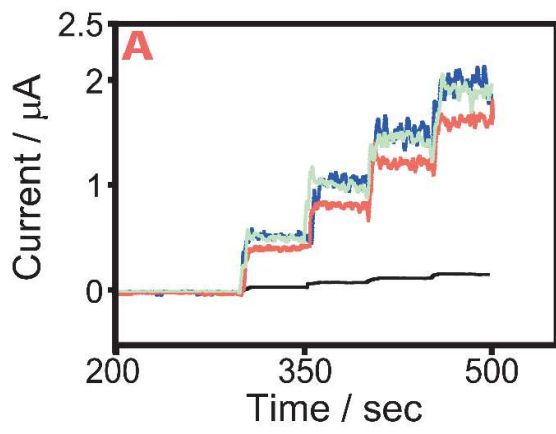


Figure 8



Tables

Table 1

	$G (cm^{-1})$	$D (cm^{-1})$	I_D / I_G
HP 160	1577 ± 0.1	1346 ± 2	1.28 ± 0.2
HS20	1582 ± 0.9	1358 ± 0.2	1.32 ± 0.02
MTN 990	1587 ± 2	1357 ± 0.9	1.19 ± 0.08
N115	1578 ± 2	1348 ± 0.5	1.33 ± 0.02
N220	1580 ± 0.8	1349 ± 1	1.28 ± 0.06
N375	1577 ± 0.6	1348 ± 1	1.27 ± 0.01
N660	1577 ± 0.2	1355 ± 0.9	1.22 ± 0.1
PL6	1575 ± 5	1342 ± 3	1.41 ± 0.04
Super P	1577 ± 0.7	1347 ± 0.5	1.41 ± 0.03
XE2B	1584 ± 2	1342 ± 1	1.57 ± 0.06

Table 2

Sample	C1s	O1s
HP 160	90.1	9.9
PL6	97.3	2.7
N220	98.3	1.7
N115	98.4	1.6
N375	96.7	3.3
XE2B	97.8	2.2
N660	98.1	1.9
Super P	99.3	0.7
HS20	97.0	3.0
MTN 990	97.6	2.4

Carbon Black	k^0 (cm/s)	Surface Area (BET) (m^2/g)	Electroactive Area (mm^2)	Particle diameter (nm)	I_D/I_G	Analytes
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Table 3

Sample	C1s components									
	283.5 eV C-C sp ¹	284.5 eV C-C sp ²	285.2 eV C-C sp ³	286.0 eV C-O-C/H	287.4 eV O-C-O	289.2 eV -O-C=O-	291.0 eV π - π^* shake up			
HP 160	19.99%	52.34%	9.07%	3.64%	9.26%	2.19%	3.50%			
HS20	-	67.65%	12.95%	7.99%	4.22%	2.95%	4.24%			
MTN 990	-	67.90%	16.01%	6.11%	3.58%	1.83%	4.58%			
N115	-	63.73%	12.47%	8.74%	4.64%	3.99%	6.43%			
N220	-	65.51%	8.87%	9.12%	5.53%	4.07%	6.91%			
N375	-	62.59%	14.49%	8.80%	5.55%	2.94%	5.63%			
N660	-	67.25%	11.84%	8.17%	4.27%	2.88%	5.59%			
PL6	-	64.00%	11.52%	9.33%	5.23%	3.69%	6.23%			
Super P	-	67.97%	13.71%	5.78%	4.41%	2.71%	6.41%			
XE2B	-	59.36%	19.23%	5.44%	5.14%	3.54%	7.30%			

Table 4

N220	1.59×10^{-2} [9], 4.5×10^{-3} [13], 1.4×10^{-2} [40] ² , 1.85×10^{-2} [41], 3.6×10^{-3} [42] ³	124 [10, 43], 120 [9]	3.2 [40], 9.3 [44], 23 [41], 2.3 [42], 3.1 [this work]	17.95-32.5 [9], 19-29 [43] ¹ , 19-25 [44], 19-29 [45] ¹ , 17.95-32.5 [46], 25 [16]	0.96 [9], 0.97 [44], 1.28 [this work]	Cat [43], NADH [9, 10, 13, 42], BQ [10], EP [10], H ₂ O ₂ [10, 44, 49], Heme protein [40], Hg [50], BPA [51, 52], As [53], AA [44], Cys [13], Phosphate [54], Glucose [44], HQ [44], Free Chlorine [55]
VXC72R	9.26×10^{-2} [16] ⁴ , 9.77×10^{-3} [48] ⁵			52 [17], 20-25 [56], 32.1 [16]	2.64 [16, 17], 1.01 [56], 0.99 [58]	DA [16, 17, 58, 59], APAP [16, 17, 47, 59, 60], AA [17], UA [17], LEV [47], Mesotrione [48], Estriol [56], AM [57], NI [57], EP [58], IP [60], PP [60], CAF [60], CAT [61]
N375	3×10^{-3} [this work]	90 [9], 105 [62,63]	14 [63], 3.1 ± 0.1 [this work]	36 [62,63]	1.27 [this work]	NADH [9], Sulfonamides [62], Nitrite [63], BQ [63], HQ [63]
N773		30 [9, 62, 63]	8 [63]	124 [62,63]		NADH [9], Sulfonamides [62], Nitrite [63], BQ [63], HQ [63]
N115	6.3×10^{-3} [this work]	143 [62, 63]	19 [63], 3.6 ± 0.2 [this work]	27 [62,63]	1.33 [this work]	NADH [9], Sulfonamides [62], Nitrite [63], BQ [63], HQ [63], AA, EP, CYS, BQ, CA, CAT [This work]
Printex XE 2	8.1×10^{-3} [4]	950 [4]		30 [5]		DA [5], AA [5,4], Pb [5], Cd [5], Cu [4]
M 430				25-37 [20]		FM [20], APAP [20], Phenolic compound [64]
ECP-600ID		1270 [65]		34 [65]		Paraoxon [65]
M 1100				14 [12]		Nicotine [12]
5358R	1.7×10^{-3} [14] ⁶			29 [14]	0.91 [14]	
BP4750	5.74×10^{-3} [16] ⁴			26 [16]	2.36 [16]	DA [16], APAP [16]
E2000	6.23×10^{-4} [16] ⁴			27 [16]	3.65 [16]	DA [16], APAP [16]
Super P Li			50.84 [66] ⁶			
HP 160	5.4×10^{-3} [this work]		3.8 ± 0.2 [this work]		1.28 [this work]	AA, EP, CYS, BQ, CA, CAT [This work]
HS20	1.2×10^{-3} [this work]		3.0 ± 0.2 [this work]		1.32 [this work]	
MTN 990	—		1.7 ± 0.1 [this work]		1.19 [this work]	
N660	2.6×10^{-3} [this work]		3.2 ± 0.2 [this work]		1.22 [this work]	
PL6	4.8×10^{-3} [this work]		3.7 ± 0.3 [this work]		1.41 [this work]	
Super P	2.2×10^{-3} [this work]		3.1 ± 0.1 [this work]		1.41 [this work]	
XE2B	4.3×10^{-3} [this work]		3.7 ± 0.2 [this work]		1.57 [this work]	

¹Given by the company, ² Using $D_0 = D_R = 5.9 \times 10^{-6} \text{ cm}^2/\text{s}$, ³ Using $D_0 = D_R = 7.6 \times 10^{-6} \text{ cm}^2/\text{s}$, ⁴ Using rutheniumhexaamine, ⁵ Calculated by impedance measurements,

⁶ Calculated by chronocoulometry. AA = Ascorbic Acid, AM = Amoxicillin, APAP = acetaminophen, BQ = Benzoquinone, BPA = Bisphenol A, CA = Caffeic acid, CAF = Caffeine, CAT Catechol, CYS = Cysteine, DA = Dopamine, FA = Folic Acid, EP = Epinephrine, FM = Ferrocenemethanol, IP = Isoproterenol, LEV = Levofloxacin, NI = Nimesulide, PP = Propranolol, TRP Tryptophan, TYR = Tyrosine.

Supplementary Materials

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