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Pt-grown carbon nanofibers for detection of hydrogen peroxide†

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Removal of left-over catalyst particles from carbon nanomaterials is a significant scientific and technological problem. Here, we present the physical and electrochemical study of application-specific carbon nanofibers grown from Pt-catalyst layers. The use of Pt catalyst removes the requirement for any cleaning procedure as the remaining catalyst particles have a specific role in the end-application. Despite the relatively small amount of Pt in the samples ($7.0\pm0.2\%$), they show electrochemical features closely resembling those of polycrystalline Pt. In O₂-containing environment, the material shows two separate linear ranges for hydrogen peroxide reduction: $1-100~\mu\text{M}$ and $100-1000~\mu\text{M}$ with sensitivities of $0.432~\mu\text{A}~\mu\text{M}^{-1}~\text{cm}^{-2}$ and $0.257~\mu\text{A}~\mu\text{M}^{-1}~\text{cm}^{-2}$, respectively, with a $0.21~\mu\text{M}$ limit of detection. In deaerated solution, there is only one linear range with sensitivity $0.244~\mu\text{A}~\mu\text{M}^{-1}~\text{cm}^{-2}$ and $0.22~\mu\text{M}$ limit of detection. We suggest that the high sensitivity between $1~\mu\text{M}$ and $100~\mu\text{M}$ in solutions where O_2 is present is due to oxygen reduction reaction occurring on the CNFs producing a small additional cathodic contribution to the measured current. This has important implications when Pt-containing sensors are utilized to detect hydrogen peroxide reduction in biological, O_2 -containing environment.

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1. Introduction

Hydrogen peroxide (H_2O_2) is a versatile chemical, its use ranging from industrial and medical to household applications. One particular branch of research related to H_2O_2 is the study of oxygen reduction reaction (ORR), which can be, for example, the cathodic half-cell reaction in polymer electrolyte membrane fuel cells. ORR is also of fundamental importance in corrosion studies. It has been proposed that oxygen can be reduced to water either via direct four electron reduction or via two two-electron steps where H_2O_2 acts as an intermediate. However, some controversy still remains and it has been stated that studying H_2O_2 oxidation and reduction could bring more insight into ORR as well.

As one of the reactive oxygen species, it is considered one of the radicals linked to aging.³ On the other hand, despite its harmful effects, H₂O₂ is also acting in carefully monitored signaling pathways in cells.⁴ Moreover, the function of

Discovery of carbon nanotubes (CNTs) in the early 90's (ref. 5) started widespread interest in the research of various carbon nanomaterials. Carbon is relatively inexpensive and abundant material with large number of allotropes and tunable properties which makes it an ideal option for development of sensor applications. Moreover, carbon materials often have functional groups that allow immobilization of enzymes via covalent bonds making them good platforms for enzymatic biosensors. Different carbon materials can also be combined to give new hybrid materials with enhanced properties. For example, CNTs grown from tetrahedral amorphous carbon (ta-C) have been shown to induce selectivity between oxidation of dopamine and ascorbic acid, which would otherwise give overlapping signals in the voltammograms. In addition to CNTs, carbon nanofibers (CNFs) can also be grown from ta-C. Growing the CNFs from another carbon material, here ta-C, offers an additional carbon source and changes the fiber morphology from bamboo-like to stacked platelet-like structure.8

Ni is often utilized as catalyst metal for the growth of CNFs. However, some of the catalyst remains at the tips of the fibers and cannot be completely removed for example by treating the samples with concentrated acid. Moreover, when aiming to develop applications for possible future *in vivo* use in

enzymatic biosensors utilizing certain oxidases is based on electrochemical detection of H_2O_2 which is one of the products from oxidation of the substrate molecule. Thus, detecting H_2O_2 accurately and reliably is important in the biomedical field.

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electrochemical sensors, nickel should be avoided as it is a common contact allergen. 10 We have previously presented the use of application-specific catalyst layers in fabrication of Ptcontaining carbon nanofibers (Pt-CNFs).11 From 10 nm thick Pt layer with underlaying ta-C thin film it is possible to grow CNFs with height up to several hundreds of nanometers. Pt is present in these fibers as a larger particle at the tip and as smaller particles along the length of the fibers. Even though part of the Ti used as adhesion layer was distributed as fine particles in the fibers, it did not visibly affect the electrochemical response. The advantages of this novel hybrid carbon nanomaterial include the well-ordered structure and control over the location where fibers are grown, which is important from the device fabrication point of view. The preliminary results showed that this novel hybrid carbon nanomaterial could be utilized for H₂O₂ detection. Here, we proceed to the in depth structural and electrochemical characterization of the Pt-CNFs and especially asses the roles of dissolved oxygen and chloride ions on the electrochemical performance. To verify the crucial role of Pt nanoparticles in the electrochemical behavior of the CNFs, carbon nanofibers grown from nanodiamonds were utilized as a metal-free reference.

2. **Experimental**

2.1. Sample preparation

Samples were prepared on boron doped p-type Si(100) wafers with 0.005Ω cm resistivity (Siegert, Germany). The wafer was first coated with a 20 nm Ti adhesion layer with direct current magnetron sputtering. Next, 7 nm ta-C was deposited on top of the Ti layer by pulsed filtered cathodic vacuum arc deposition in base pressure below 1.3×10^{-4} Pa. Before CNF growth 10 nm Pt catalyst layer was deposited also with cathodic arc deposition. More detailed description of the deposition procedures can be found in our earlier publication12 with the difference that here instead of using a gradient function for the deposition, first only carbon was sputtered from one cathode and then only Pt from another cathode. Finally, samples were placed in the cold-wall plasma-enhanced chemical vapor deposition reactor (Aixtron, Black Magic, Germany) and CNF were grown as described in detail in our previous publication.11 In brief, NH3/C2H2 gas mixture (125/30 sccm) was injected to the reactor at growth temperature of 750 °C for 30 minutes. In addition to using Pt as catalyst, nanofibers were also grown from carboxyl functionalized zeta-negative nanodiamonds (NDs, Carbodeon uDiamonds, Carbodeon, Vantaa, Finland) in order to obtain a metal-free reference material for the Pt-CNFs. The concentration of the original waterbased ND solution was 5 wt% and it was diluted into 0.05 wt% in ethanol. The NDs were deposited on the silicon chips by spray coating with pressurized air (3.5 bar) as the carrying gas. The distance between the spray gun and samples was 10 cm. Scanning over the samples was repeated 10 times. CNF growth process was the same as for the Pt-CNF samples except for the growth time which was 30 min and temperature which was 600 $^{\circ}$ C.

In addition to the ND-CNFs also 10 nm Pt thin film samples and spherical polycrystalline Pt electrode were utilized for comparison in electrochemical experiments.

For electrochemical experiments (cyclic voltammetry and amperometry) a circular area (varying geometric areas) was defined with PTFE tape. For rotating disk electrode experiments a special holder with radius of 3 mm was utilized.

2.2. Physical characterization

HRTEM samples were prepared by focused ion beam (FIB) milling using polymer ink as filler material. 70-80 nm Pt and Pt-C (FIB deposited) were used to protect the CNFs during sample fabrication. Double aberration-corrected microscope (JEOL 2100, Japan) equipped with an X-ray detector was utilized for high-resolution TEM imaging. Digital recording was performed with a Gatan 4× 4 UltraScan 4000 CCD camera.

SEM images were obtained with Zeiss Sigma VP. Sample were mounted on holders (planar and cross-sectional) with double-sided carbon tape and Cu tape. No additional coating was used.

XPS measurements were performed with Kratos AXIS Ultra Xray photoelectron spectroscope (Kratos Analytical, Manchester, UK) using monochromated Al Kα X-ray irradiation at 100 W. Samples were pre-evacuated in sample transfer chamber overnight. Both survey-spectra and high-resolution spectra of C 1s, O 1s and Pt 4f was collected from three locations per sample. Cellulose filter paper (Whatman) was used as an internal in situ ref. 13 Analysis depth was expected to be <10 nm and area 400 imes800 µm². CasaXPS software was used for both extracting the elemental atomic percentages from the survey spectra and fitting the C 1s and Pt 4f high-resolution regimes (Shirleybackground with assumed Gaussian line shapes). The reported cellulose in situ reference was used for charge correcting the binding energies, where 286.7 eV was assigned for carbon atoms bonded to single oxygen atom.14

2.3. Electrochemical experiments

All electrochemical experiments were performed with either Gamry Reference 600 or 600 + potentiostat (Gamry Instruments) with Ag/AgCl/KCl sat'd reference electrode (Radiometer Analytical) and Pt wire counter electrode. To address the difference in pH some of the results are presented against RHE. The conversion was done by utilizing Nernst equation taking into account solution pH. Rotating disk electrode (RDE) experiments were conducted by using a Radiometer Analytical rotating motor (Model EDI101). Samples used in RDE were examined in SEM before and after the experiments and despite some minor agglomeration the fibers retained their vertical alignment. Amperometric experiments were performed by stepping the potential from 0 V to -0.15 V vs. Ag/AgCl. Between measurements the solution was mixed either with magnetic stirrer or bubbling with N2. The magnetic stirrer was off during the measurement.

Unless stated otherwise, the solution were purged with N2 before starting the experiment and gas blanket was used during the measurements. The electrochemical characterization was performed by scanning in 0.15 M H₂SO₄ (diluted from 95–97% H₂SO₄, Merck Millipore) at 50 mV s⁻¹. Scanning was also repeated in phosphate buffered saline (PBS, pH 7.4) containing

NaCl (137 mM), KCl (2.7 mM), Na₂HPO₄ (10 mM) and KH₂PO₄ (1.8 mM) as well as phosphate buffer without chlorides (PB_{Na}, pH 7.2) containing H₂NaPO₄ (30 mM) and HNa₂PO₄ (70 mM). Samples were also characterized in 1 mM hexaamminer-uthenium(III) chloride (Ru(NH₃)₆Cl₃, Aldrich, 98%) in 1 M KCl. H₂O₂ stock solutions were prepared from 30% H₂O₂ (Merck

For comparison, phosphate buffer (PB_K, pH 7.2) was also prepared using potassium phosphates instead of sodium phosphates. To assess the effect of pH and adding phosphates a solution containing both 0.1 M $\rm H_2SO_4$ and 0.1 M $\rm KH_2PO_4$ was prepared (pH 1.3).

Results and discussion

KGaA, Germany) by diluting in deionized water.

3.1. Characterization

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The SEM micrographs in Fig. 1 show that the CNFs form a thick forest of fibers with average length of approximately 500 nm. From the top view image (Fig. 1B) it can be seen that the fibers form bundles with some tens of nanometers between them. The catalyst particles are also clearly visible in the SEM micrographs. Especially the top view illustrates the size of the particles varies considerably, possibly partly due to the bundling of the fibers. HRTEM micrographs in Fig. 1C and D show the fibers and the catalyst particles in more detail. Together with the SEM images (Fig. 1A), the HRTEM micrograph in Fig. 1C highlights that the length of the fibers varies from approximately 60 nm to 600 nm. This is somewhat shorter than presented in our

previous publication¹¹ where the lengths of the fibers were observed to span from few hundreds of nanometers to even up to micrometer scale. This is explained by the shorter growth period (30 min vs. 60 min) for the sample batch used here as well as the stochastic growth process. It is visible from Fig. 1D that some of the catalyst Pt particles at the tips of the fibers are in fact single crystals. However, as presented in the electrochemical characterization section below, the Pt–CNF show polycrystalline-like behavior when cycled for example in H₂SO₄, indicating that the particles are not terminated in any preferential orientation. Such lack of preferential orientation might be a consequence of the oxidation and reduction cycles applied during the cleaning procedure.

The XPS spectra in Fig. 1E further verifies the presence of Pt through the Pt 4f doublet. Moreover, it was estimated from XPS data that the amount of Pt in the samples was $7.0\pm0.2\%$. Fitting the Pt 4f peaks (ESI Fig. S1†) showed that approximately 75% of the Pt in the samples is in metallic form and the rest as different Pt oxides. Some estimates on the nature of the oxides is provided in the ESI.† Interestingly the amount of Ti detected by the XPS was 7.2% which is considerably higher than for amorphous carbon films with intrinsic Pt-gradient, where all the Ti utilized as adhesion layer was buried under the thin film. We have previously shown that when the Pt catalyst layer is 10 nm in addition to the irregularly shaped Pt particles at the tips the grown fibers have finely distributed Ti along their lengths. However, most of the Ti adhesion layer is still intact which has been suggested to be a consequence of thick enough

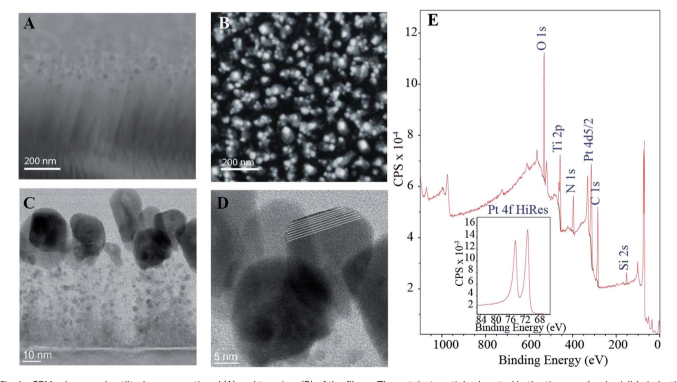


Fig. 1 SEM micrographs: tilted cross-sectional (A) and top view (B) of the fibers. The catalyst particles located in the tips are clearly visible in both images. The HRTEM micrographs (C and D) show the fibers and the catalyst particles in more detail. As can be seen from the middle particle in D, some of the catalyst particles are single crystals even though the samples show mostly polycrystalline behavior (the white lines in D have been added to help guide the eye). (E) XPS wide spectrum for Pt-CNFs with the inset showing the high resolution spectrum for the Pt 4f doublet.

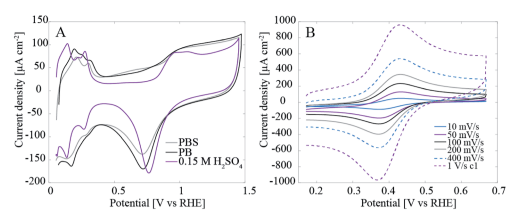


Fig. 2 (A) Voltammograms for Pt–CNFs at 50 mV s $^{-1}$ in PBS, in phosphate buffer and in 0.15 M H $_2$ SO $_4$. (B) Electrochemical characterization of the Pt–CNFs in 1 mM Ru(NH $_3$) $_6$ Cl $_3$ in 1 M KCl at different scan rates. To take the pH difference into account the graphs in both A and B are presented in RHE scale even though the original measurements were performed against Ag/AgCl/KCl sat'd. Conversion was done by utilizing Nernst equation taking into account solution pH.

catalyst layer protecting the ta-C film as well as a prerequisite for the proper fiber growth.

3.2. Electrochemical characterization and H₂O₂ detection

Fig. 2A shows that especially in $\rm H_2SO_4$ the Pt features including oxide formation and reduction as well as hydrogen adsorption and desorption are clearly visible. The hydrogen region below 0.4 V vs. RHE has been denoted as the fingerprint for clean polycrystalline Pt and since especially the smaller peak at 0.2 V vs. RHE is only visible on very clean surfaces it can be said that despite the dirty initial conditions Pt can be cleaned efficiently by potential cycling. The dirty conditions refer here to the inevitable presence of carbon impurities in the solution – originating from the samples themselves – that could contaminate Pt. On the other hand, in the buffer solutions all the Pt features are still visible but clearly the cleaning effect of potential cycling in this media is less efficient.

Samples were also characterized with 1 mM hexaamminer-uthenium($_{\rm III}$) chloride (Ru(NH₃)₆Cl₃, Aldrich, 98%) in 1 M KCl. Pt–CNFs showed reversible electron transfer kinetics: $\Delta E_{\rm p}$ increased from 59 mV for 10 mV s⁻¹ to 64 mV for 1 V s⁻¹ (Fig. 2B). As the electron transfer was practically reversible, the electrochemically active surface area was estimated from the data by using Randles–Sevcik equation:

$$I_{\rm P} = (2.69 \times 10^5) n^{2/3} A D^{1/2} v^{1/2} C_0, \tag{1}$$

where $I_{\rm p}$ is the anodic peak current, n is the number of transferred electrons (here n=1), A is the electrochemically active surface area, D is the diffusion coefficient (for $[{\rm Ru}({\rm NH_3})_6]^{2^+}$: 7.7 \times 10⁻⁶ cm² s⁻¹), ¹⁶ C_0 is the bulk concentration of ${\rm Ru}({\rm NH_3})_6{\rm Cl_3}$. The surface area was calculated for each scan rate with eqn (1) and the average was 0.062 \pm 0.003 cm² which is 88% of the geometric area of the sample (0.071 cm²). The electrochemically active surface area was similarly estimated for the ND-grown nanofibers (ND-CNFs) resulting in approximately 92% of the geometric area (0.031 cm²). The use of Randles–Sevcik equation was justified also in this case as the ND-CNFs were also

characterized with 1 mM Ru(NH₃)₆Cl₃ and they showed reversible behavior (ESI, Fig. S2†). Thus, it cannot be stated that the electrochemical activity in this case would purely arise from the Pt particles. As the actual area for nanomaterials is typically larger than the geometric area this result indicates that there are certain regions that are possibly inaccessible for the electrolyte or the probe molecules. We propose that this might be due to, for example, stacking of the nanofibers. The area was also estimated from CV data in 0.15 M H₂SO₄ by integrating the area below the hydrogen desorption features using the doublelayer region current as baseline and dividing this by charge density (210 μC cm⁻²). The calculated area was 0.11 cm² and it corresponds only to Pt available for hydrogen adsorption. It is not surprising that electroactive area from adsorption processes is higher than that measured from a diffusional process since it is well known that the overlay of diffusional fronts from different microroughnesses tends to decrease the area measured in this way.18 Since the electrochemical methods for active surface area measurements give contradictory results, geometric area was utilized when calculating current densities in this work.

We have previously presented initial results that showed that the Pt-CNFs are suitable for amperometric detection of H₂O₂ in O₂-containing environment.¹¹ Fig. 3A shows that when potential is stepped from initial value of 0 V to -0.15 V vs. Ag/AgCl and when H₂O₂ concentration is increased by intervals of 10 μM between 10 μM and 100 μM and by intervals of 100 μM between 100 μM and 1000 μM (increase in volume was the same throughout the whole region) in non-deaerated PBS there are in fact two separate linear ranges: first from 1 μM to 100 μM with sensitivity of 0.432 $\mu A \mu M^{-1} cm^{-2}$ and second from 100 μM to 1000 μ M with sensitivity of 0.257 μ A μ M⁻¹ cm⁻². The limit of detection was 0.21 μ M (S/N = 3). The limit of detection presented here is considerably larger than what we presented for the Pt-CNFs in our previous publication (0.2 nM). On the other hand, the sensitivities for both linear ranges are slightly higher but still in the same range as the one published previously (0.148 μ A μ M⁻¹ cm⁻²). The inconsistency between these results

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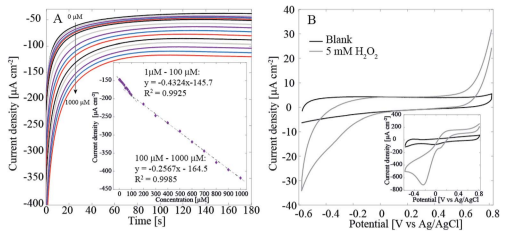


Fig. 3 (A) Amperometric detection of H_2O_2 with the Pt–CNF samples in non-deaerated PBS. The potential was stepped from 0 V to -0.15 V vs. Ag/AgCl for 180 s. The concentration was first increased to 1 μ M and then to 5 μ M. Between 10 μ M and 100 μ M the concentration was increase by steps of 10 μ M and above 100 μ M by steps of 100 μ M. The data points were obtained by averaging the current between 0.5 s and 1 s. (B) Cyclic voltammograms recorded at 50 mV s⁻¹ for ND–CNFs in blank and H_2O_2 -containing PBS (deaerated). Inset shows the same measurement for Pt–CNFs.

could, at least to some extent, arise from the variation between and within sample batches. The growth process of the fibers depends on such parameters as temperature, time period and catalyst layer thickness. In addition, the condition of the growth chamber, *i.e.* if it has been cleaned recently, can have an effect on the fibers. Finally, the growth process is highly stochastic

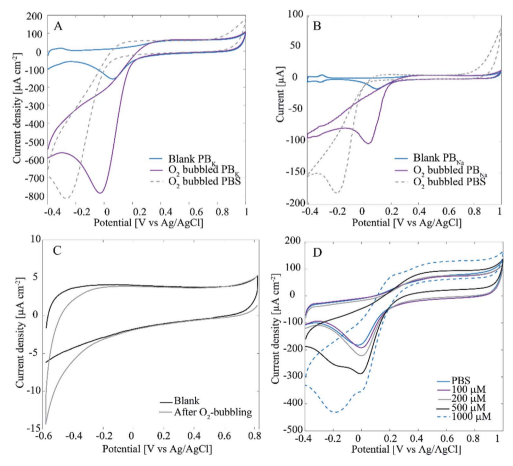


Fig. 4 (A) Voltammograms for Pt–CNFs in blank (N_2 -saturated) PB_K and O₂-saturated PB_K and PBS. (B) Voltammograms for polycrystalline Pt in blank (N_2 -saturated) PB_{Na} and O₂-saturated PB_{Na} and PBS. (C) Voltammograms for ND–CNFs in blank (N_2 -saturated) PBS. (D) Voltammograms for Pt–CNFs in blank and H₂O₂-containing PBS (non-deaerated). Scan rate in (A–D) was 50 mV s⁻¹.

PBS

1.07

Table 1 Resistivities and pH values for H_2SO_4 , phosphate buffers (PB_{Na} and PB_K), mixed solution of H_2SO_4 and KH_2PO_4 , and PBS

which affects for example the length and diameter of the fibers (Fig. 1A and C).

To verify that the main current here arises from the Pt catalyst particles, we measured the reduction of $\rm H_2O_2$ with the Pt-free ND-CNFs for comparison. Fig. 3B shows that even with as high concentration as 5 mM of $\rm H_2O_2$ the current density for its reduction is very small (less than 10%) compared to the current density obtained in with Pt-CNFs (Fig. 3B inset). This supports the assumption that it is mainly Pt particles, and not the CNFs, that catalyse the reaction.

Interestingly, when measuring H_2O_2 in deaerated (O_2 -free) solutions there is only one linear range from 5 μ M to 1000 μ M with sensitivity of 0.244 μ A μ M⁻¹ cm⁻² and 0.22 μ M limit of

detection (results not shown here). Since the only difference between the two experiments was the presence of O_2 in the cell, it is assumed that it must contribute to this behavior. In fact, it is well known that Pt is an efficient catalyst for ORR. Another important phenomenon is the effect of Cl⁻ adsorption on ORR and H₂O₂ reactions of Pt. To test these phenomena with the Pt-CNFs, voltammograms were recorded in O2-saturated PBS containing high concentration of Cl⁻ and chloride-free phosphate buffers (PB_K, PB_{Na}) as well as 0.15 M H₂SO₄ (Fig. 4A shows results only for PBS and PBK since PBNa and 0.15 M H2SO4 were very similar to PB_K). It has been previously shown that chlorides strongly inhibit the oxygen reduction reaction.19 For all solutions the current magnitude in the cathodic end is increased considerably indicating the presence of O2 in the solution. However, there is a clear difference for the onset and peak potentials for oxygen reduction between PBS and the other solutions: for example in PB_K the current starts to decrease at 0.2 V vs. Ag/AgCl reaching the peak at 0 V whereas for PBS ORR onset is shifted to approximately 0 V and the peak to -0.2 V. As said, the main difference in composition especially between the two phosphate buffers and PBS is the absence/presence of chlorides. PBS contains large amounts of chlorides (approximately 140 mM) which explains the shifting of the peak to -0.2 V vs. Ag/AgCl compared to the blank, N₂-saturated

Table 2 Summary of some previous works from the last 10 years where Pt has been utilized in some form for sensors detecting H_2O_2 reduction. The studies have been divided into those where H_2O_2 has been detected in phosphate buffer and those where PBS has been used as the electrolyte (chloride-containing solution). In some studies the sensitivity was given in respect to current and not currents density. This is indicated by /A in the table

7.42

Material	$E_{ m Ag/AgCl}$	$_{\left(\mu M\right) }^{LOD}$	Sensitivity $(\mu A \mu M^{-1} cm^{-2})$	Response time (s)	Linear range (µM)	Reference
Phosphate buffer						
Pt-CNT nanocomposites	-0.1	1.5	_	5	5-25 000	26
Au/Pt NP-coated CNT/silica nanocables	-0.1	0.3	_	3.5	0.5-1670	27
PtNPs on graphene hydbrid nanosheet	0	0.08	_	4	1-500	28
Au-Pt nanowires	0.045	1.5	0.1292	7	20-8380	29
Pt NP/MWCNT clusters on	-0.4	1.23	0.003359/A		5-2000	30
screen-printed Au nanofilm					1780-16 800	
Ultralow Pt-loading Au nanoflowers	0.09	0.006	0.003184/A		0.025-3.45	31
C			0.000316/A		5.91-94.3	
Pt/porous graphene on GC	-0.1	0.5	0.3411	3	1-1477	32
Au@C@Pt nanocomposite on GC	0	0.13	0.1447	10	9-1860	33
-			0.0801		1860-7110	
Pd core-PtNDs/PDDA-rGO on GC	0.018	0.027	0.6728	5	5-500	34
Au-Pt core@shell NPs	-0.6	0.06	_	3	0.1-1	35
Fe@Pt core-shell NPs	-0.355	0.75	0.2190	1	2.5-41 605	36
SiO ₂ nanorods/APTMS/AuPt	-0.155	2.6	0.0467	5	5-72 000	37
SiO ₂ nanorods/APTMS/Pt	-0.275	0.7	0.0608	12	5-5100	
			0.0127		5100-134000	
PtNPs	-0.035	4.2	0.1103	3	5-58 000	
AuNPs	-0.175	5	0.0014	4	5-43 000	
			0.0028		43 000-77 000	
Phosphate buffered saline						
Pt-Au NP/titania nanotube array	-0.2	10	0.00292/A		10-80	38
Pt-Au NPs/reduced graphene sheets	0.045	0.31	0.006168/A		1-1780	24
a-C/Pt	-0.2	0.32	0.32	0.05	50-1000	12
Pt-CNFs	-0.15	0.21	0.432		1-100	This work
			0.257		100-1000	

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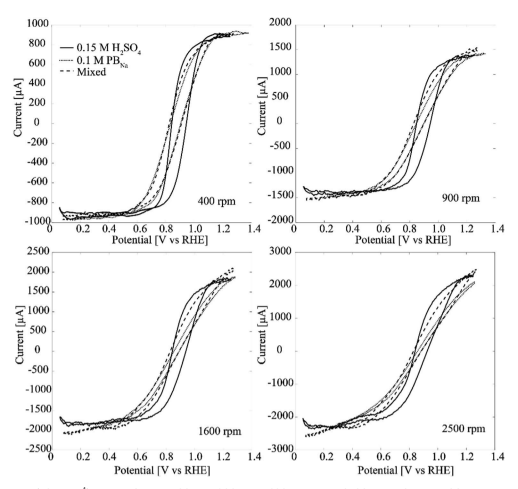


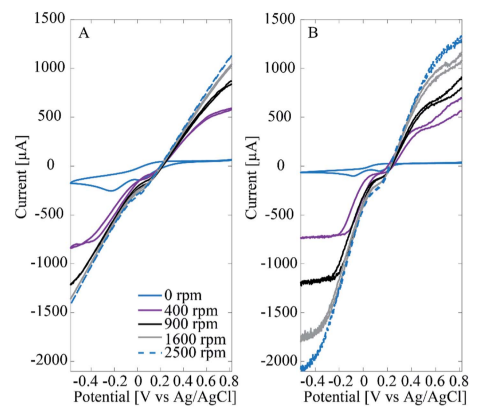
Fig. 5 RDE experiments (50 mV s⁻¹) with Pt–CNFs at 400 rpm, 900 rpm, 1600 rpm and at 2500 rpm in 0.15 M H_2SO_4 , PB_{Na} and mixed solution of phosphate buffer and H_2SO_4 with 5 mM H_2O_2 . To take the pH difference into account the graphs are presented in RHE scale even though the original measurements were performed against Ag/AgCl/KCl sat'd. Conversion was done by utilizing Nernst equation taking into account solution pH. In addition the ohmic drop was compensated for the PB_{Na} by running a post iR correction ($R_u = 100 \ \Omega$) in Echem Analyst software.

solution. For clarity, Fig. 4A shows only the blank for PB_K as it was nearly identical with that of PBS. In absence of chlorides $(PB_{Na},\ PB_K\ and\ H_2SO_4)$ oxygen reduction coincides with Pt oxides reduction as expected.

Fig. 4A also shows that in PBS the current magnitude is increased also at the anodic end of the curve when O_2 is present. For the other solutions, the anodic end were comparable to the respective blanks. This is suggested to arise from incomplete ORR, which does not proceed to H2O in chloride-containing solution but actually produces additional H₂O₂ into the solution, which is then oxidized at anodic potentials above 0.8 V vs. Ag/AgCl. Similar experiment was also tested with a spherical polycrystalline Pt electrode under well-controlled conditions and the results obtained (Fig. 4B) were identical to those of Pt-CNFs. The proposed production of H₂O₂ was further tested with polycrystalline Pt by reversing the potential at 0.2 V vs. Ag/AgCl to avoid the ORR. There was no extra anodic current compared to the blank curve which supports the suggestion that H2O2 is produced during the ORR (results not showed here). Finally, the experiment was also repeated with ND-CNFs (Fig. 4C). Oxygen is reduced at the cathodic end but the current (density) magnitude from ORR on carbon can be considered negligible and thus the

main contribution indeed arises from Pt in the Pt-grown nanofibers. The small cathodic current arising from the ORR at the CNFs themselves and not the Pt catalyst particles provides an additional contribution to the measured currents for small $\rm H_2O_2$ concentrations. This additional contribution becomes negligible as soon as the concentration of $\rm H_2O_2$ becomes high enough so that the current arising from this process starts to dominate the total current. It is to be noted that the amperometric experiments were conducted at -0.15 V νs . Ag/AgCl which is in the potential range for ORR. Thus, this coupling of two electrochemical processes in $\rm O_2$ -containing solutions explains why there are two linear ranges in the non-degassed PBS and not in the $\rm N_2$ -purged solution. This is a fact that must be taken into account when results from the literature (see Table 2) are considered.

When cycling the potential of the samples in non-deaerated PBS containing large amounts ($<500 \,\mu\text{M}$) of H_2O_2 , an additional peak appeared approximately at -0.2 V νs . Ag/AgCl in the voltammogram (Fig. 4D). Similar peak appeared also in N_2 -purged solution but only with even higher concentration of H_2O_2 (1000 μM , see ESI Fig. S3†). It has been stated by Katsounaros $et~al.^2$ that at high enough anodic potentials where OH has been adsorbed on Pt, O_2 is produced in a non-electrochemical step:



RDE experiments (50 mV s⁻¹) with Pt-CNFs at 400 rpm, 900 rpm, 1600 rpm and at 2500 rpm in PBS with (A) Pt-CNF and (B) 10 nm Pt thin film.

$$2Pt(OH) + H_2O_2 \rightarrow 2Pt(H_2O) + O_2$$
 (2)

Thus, even in initially degassed solutions when cycling the potential in a large enough window, O2 originating from added H_2O_2 can be expected to be present. This means that in addition to hydrogen peroxide redox reactions oxygen reduction is also likely to occur. Moreover, there might be some threshold concentration for O2 since the peak only appears with addition of H₂O₂ in non-degassed solutions where O₂ can be expected to be present initially. This is further supported by the peak only appearing with addition of larger amount of H2O2 in N2-purged PBS (Fig. S3†). It should be noted that the peak does not appear if the upper potential limit is not high enough (>0.5 V vs. Ag/ AgCl, results not shown here) which further supports the proposed origin (OH has to be stable on Pt for producing O_2).

H₂O₂ oxidation and reduction were also studied by using RDE. It is shown in Fig. 5 that compared to 0.15 M H₂SO₄ oxidation and reduction of H₂O₂ are kinetically hindered in 0.1 M phosphate buffer as higher (lower) potentials are needed for reaching the limiting currents at both anodic and cathodic ends. The solution was prepared without chlorides to avoid the inhibitory effect. One explanation for the apparently slower kinetics could be that the resistivitities for the buffer solutions are higher compared to H_2SO_4 : 1.48 Ω m (PB_{Na}) and 1.18 (PB_K) vs. 0.58 Ω m, respectively. In order to compensate this difference in resistivity a post iR correction was run for the PB data presented in Fig. 5 in Echem Analyst software for the ohmic drop (original graphs without post iR correction are presented in the ESI as Fig. S4†). When adding KH_2PO_4 into H_2SO_4 the resistivity slightly increased to 0.67 Ω m, which is likely a consequence of the increased pH. Fig. 5 shows that especially the H₂O₂ reduction is clearly hindered both in PB (iR corrected data) and the mixed solution compared to pure H₂SO₄. The effect is somewhat larger than what would be expected from the relatively small change in resistivity. On the other hand, hydrogen peroxide is reduced according to eqn (3):

$$H_2O_2 + 2H^+ + 2e^- \leftrightarrow 2H_2O$$
 (3)

which indicates that acidic environments can promote the reaction. Thus, the considerably higher pH of PB could partly explain the slower kinetics. However, as the pH for H₂SO₄ + KH₂PO₄ mixed solution was 1.26 compared to 0.95 for H₂SO₄. we assume that such small change is pH would be unlikely to have such a large effect. Resistivity and pH values for different solutions utilized in this study are shown in Table 1.

Since the addition of KH₂PO₄ causes a significant effect in the RDE curve especially in the cathodic end of the transition region, we suggest that this could be related to adsorption of phosphate anions. It has been shown that phosphate species adsorb on polycrystalline as well as Pt (111) and Pt (100) surfaces, with the exact nature of the adsorbed species changing with pH and electrode potential.20-22

Fig. 6 shows results from RDE experiments in deaerated PBS with 5 mM H₂O₂ for both Pt-CNF and 10 nm Pt thin film **RSC Advances**

samples. The inhibiting effect of Cl⁻ is clear. In both Fig. 6A and B there is a prewave between 0.0 V and 0.1 V vs. Ag/AgCl. We propose that at larger overpotentials Cl- is desorbed and the H₂O₂ reduction is further enhanced. With the Pt thin films especially at lower rotation rates the limiting current is eventually reached, particularly in the cathodic end. However, with Pt-CNFs there is similar hindrance than seen in phosphate buffer and the mixed solution both before and after the prewave.

The main concern related to biosensors based on H₂O₂ detection that arises from the results presented here is that in biological environment it is not possible to completely avoid chlorides. This has been previously stated also by Hall et al.23 when they studied the inhibitory effect of Cl⁻ on H₂O₂ oxidation on Pt. Table 2 summarizes some works from the last 10 years where Pt has been used in some form in sensors for H₂O₂ reduction. Majority of these studies has been in fact conducted in phosphate buffer without any chlorides. Here it has been shown that chlorides have an effect in the whole water window of Pt which should be taken into account when designing sensors based on H₂O₂ detection for biomedical purposes. For amperometric sensors operating at potentials where ORR is expected, special care should be taken since the production of H₂O₂ from chloride-inhibited reduction of O₂ will likely affect the obtained response. O2 has been previously mentioned as cathodic interfering species for H2O2 detection on Pt24 and on ZnO²⁵ in biological environment due to ORR. However, to our knowledge, the effect of additional H₂O₂ from ORR in chloridecontaining environment has not been addresses before. Moreover, in regard to enzymatic biosensors, consumption of oxygen can seriously affect the enzymatic production of H2O2 and consequently the sensor operation since O2 is a necessary cosubstrate for various oxidases common in biosensors.

Conclusions

Here we have presented in depth physical and electrochemical characterization of carbon nanofibers grown from Pt-catalyst. Despite some of the catalyst particles at the tips of the fibers being single crystals, the Pt-CNFs show electrochemical characteristics that resemble those of polycrystalline Pt. The Pt-CNFs are suitable for H₂O₂ detection: in O₂ containing solutions there are two separate linear ranges, 1-100 µM and 100-1000 μM with sensitivities of 0.432 μA μM^{-1} cm⁻² and 0.257 μA μM^{-1} cm⁻², respectively. The limit of detection was 0.21 μM. In deaerated solutions they show wide linear range between 5 µM and 1000 μ M for H₂O₂ with sensitivity of 0.244 μ A μ M⁻¹ cm⁻² (similar to the second linear range in O2-containing solution) and 0.22 µM limit of detection. We suggest that these observations are the consequence of ORR occurring on the surfaces of CNFs in O2-containing solutions, which produces a small additional cathodic current that contributes to the total measured current, the effect being significant especially at low concentrations of added H₂O₂. Also, when chlorides are present in the measurement medium, their inhibiting effect on the ORR on Pt nanoparticles will heavily affect the measurements at the higher (>500 μM) region. Thus, when designing H₂O₂ sensors for biological applications the effect of chlorides and dissolved

oxygen on the measured response should be taken into account as both are difficult to avoid in the above stated applications.

Conflicts of interest

There are no conflicts to declare.

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