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Doped Overoxidized Polypyrrole Microelectrodes as Sensors for the Detection of Dopamine Released from Cell Populations

Luigi Sasso, Arto Heiskanen, Francesco Diazzi, Maria Dimaki, Jaime Castillo-León, Marco Vergani, Ettore Landini, Roberto Raiteri, Giorgio Ferrari, Marco Carminati, Marco Sampietro, Winnie E. Svendsen and Jenny Emnéus

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A surface modification of interdigitated gold microelectrodes (IDEs) with a doped polypyrrole (PPy) film for detection of dopamine released from populations of differentiated PC12 cells is presented. A thin PPy layer was potentiostatically electropolymorized from an aqueous pyrrole solution onto electrode surfaces. The conducting polymer film was doped during electropolymerization by introducing counter ions in the monomer solution. Several counter ions were tested and the resulting electrode modifications were characterized electrochemically to find the optimal dopant that increases sensitivity in dopamine detection. Overoxidation of the PPy films was shown to contribute to a significant enhancement in sensitivity to dopamine. The changes caused by overoxidation in the electrochemical behavior and electrode morphology were investigated using cyclic voltammetry and SEM as well as AFM, respectively. The optimal dopant for dopamine detection was found to be polystyrenesulfonate anion (PSS). Rat pheochromocytoma (PC12) cells, a suitable model to study exocytotic dopamine release, were differentiated on IDEs functionalized with an overoxidized PSS-doped PPy film. The modified electrodes were used to amperometrically detect dopamine released by populations of cells upon triggering cellular exocytosis with an elevated K+ concentration. A comparison between the generated current on bare gold electrodes and gold electrodes modified with overoxidized doped PPy illustrates the clear advantage of the modification, yielding 2.6-fold signal amplification. The results also illustrate how to use cell population based dopamine exocytosis measurements to obtain biologically significant information that can be relevant in, for instance, the study of neural stem cell differentiation into dopaminergic neurons.

Introduction

Conducting polymers have gathered attention from the global scientific community since the discovery of the conductive properties of doped polyyacetylene in the late 70’s. Their use in analytical sensors has been moved forward mainly because of the flexibility they provide, offering both intrinsic properties and the ability for functionalization, rendering the polymer matrix sensitive to, e.g., pH, inorganic ions, organic molecules and gases. Particularly polypyrrole (PPy) has been attractive because of the properties of the pyrrole monomer, having good water solubility, as well as being readily oxidizable and commercially available. PPy has been widely used in electrochemical sensors because it facilitates biomolecule immobilization onto electrode surfaces, can be functionalized, is biocompatible, and provides protection against electrochemical fouling.

The complex matrix structure of an electropolymorized film facilitates the doping of PPy with specific ions and organic molecules, ranging from small ions to large organic molecules such as enzymes, by entrapping the dopants in the film during its deposition onto an electrode surface. Several studies have been presented in literature illustrating the effects of doping on the properties of a PPy film, which are dependent on the dopant and the electropolymerization conditions.

The catecholamine, dopamine, is produced in vivo in different parts of the peripheral and central nervous system (CNS) as well as in the medulla of adrenal glands. This molecule is involved in a wide range of physiological processes, including hormonal regulation, cardiovascular and renal functions, sleep, feelings of reward, memory and learning as well as control of motor functions. In the CNS, dopamine functions as a neurotransmitter, being released by presynaptic neurons upon arrival of an action potential and received by the G-protein coupled receptors of the post synaptic neurons, initiating a cascade of biochemical processes. Different neurological disorders, e.g., schizophrenia and Parkinson’s disease (PD), can be caused by abnormality in the dopaminergic activity in the CNS. Schizophrenia is related to increased activation of dopamine receptors. Although gradual loss of dopamine is a typical trait of old age, its complete depletion due to death of dopaminergic neurons in the CNS region substantia nigra causes impairment of motor functions, which is a characteristic symptom of PD, the second most common neurodegenerative disease related to old age.

Because of the tight correlation between dopamine and brain functions, detection of dopamine release from cells in vitro and in vivo has been a fundamental topic in brain research. Dopamine and other catecholamines can be detected electrochemically upon oxidation on an electrode surface, resulting in numerous studies that have revealed, for instance, the fundamental mechanism of neurotransmitter release in the process known as exocytosis.
Inside neurons, neurotransmitters are sequestered in vesicles that upon stimulation are fused with the plasma membrane in a Ca\(^{2+}\)-influx mediated process, resulting in the release of the vesicular contents into the extracellular environment. Electrochemical investigations have demonstrated that neurotransmitters are released as small packages upon fusion of one vesicle at a time,\(^{20}\) known as quantal release. Typically, micromanipulated carbon fiber microelectrodes (CFMEs) have been applied in electrochemical detection of released dopamine from single cells.\(^{20}\) However, during recent years, metal electrode systems based on lithographic microsensor technology have gained increasing attention.\(^{21}\) Aside from suitability for single cell detection, microfabricated metal electrodes have the advantage of facilitating monitoring of dopamine release from populations of adherently growing cells. This approach gives possibility for fast statistically significant screening of cell populations with respect to their dopaminergic properties. It can effectively answer biologically relevant questions in the field of neural stem cell research where the ability of chosen differentiation conditions need to be evaluated in terms of an increase in the number of dopaminergic neurons in a cell population.\(^{22,23}\)

Dopamine detection at a bare metal electrode surface can be hampered by electrode fouling due to polymerization of dopamine on the electrode surface\(^{24}\) during electrooxidation and by the presence of negatively charged interferents, such as ascorbate.\(^{25}\) To overcome the influence of these factors, different electrode modifications, e.g., thiol self-assembled monolayers (SAMs)\(^{26}\) and overoxidized PPy,\(^{25}\) have been utilized. In this paper, we describe PPy modification of interdigitated gold electrode modifications, e.g., thiol self-assembled monolayers (SAMs)\(^{26}\) and overoxidized PPy,\(^{25}\) have been utilized. In this paper, we describe PPy modification of interdigitated gold IDEs, which retains its conductive metallic properties. The effect of counter ion doping on the morphological film created by electropolymerization can be seen in the SEM and AFM images shown in Fig. 2. In the case of both small and large counter ion doping, overoxidation of PPy results in flattening of the film (compare Fig. 2A,B with 2C,D and Fig. 2E,F with 2G,H). The AFM images show that the flattening resulting from the overoxidation also reduces the roughness of the film, therefore affecting its porosity. Hence, even if the overoxidation yields a loss of conductivity in the film, the thinning of this film reduces the electron transfer distance between the electrochemically active analyte and the underlying gold electrode, which retains its conductive metallic properties.

Results and discussion

All the electrodes used in this work were fabricated onto a silicon-based microchip in gold using standard cleanroom technology. Each set of interdigitated electrodes used as working electrode (WE) was coupled with its own counter (CE) and reference electrode (RE) present on the microchip, as shown in Fig. 1A. This individual addressability of the WEs allowed selective formation of PPy on each WE using a specific counter ion.

Polypyrrole electropolymerization and influence of overoxidation

In potentiostatic polymerization, the formation of PPy is initiated by the creation of pyrrole radical-cation intermediates and further on dimmers and oligomers that are adsorbed at the electrode surface, initiating the formation of polymer chains in different nucleation points.\(^{27}\) Allowing this process to proceed for a certain time, PPy films with varying morphological and electrochemical properties are created on the electrode depending on the time poised at the applied potential and the used counter ion.

Subjecting PPy films to several potential cycles in the presence of oxygen, reaching sufficiently high overpotentials (up to 1 V), results in irreversible overoxidation of the polymer film, blocking the electron transfer in the nitrogen-containing aromatic rings of the backbone and forming oxygen-containing functionalities such as carbonyl\(^{28}\) and carboxylic groups\(^{29}\). Although these functionalities reduce the conductivity of the PPy film, the resulting electronegativity attracts cations to the surface of the film, therefore rendering the electrode more sensitive to, for instance, dopamine, as well as more perselective, eliminating anionic interferents, e.g. ascorbate and dopamine metabolite dihydroxyphenylacetic acid.\(^{30}\)

When evaluating the influence of counter ion doping on the morphology of electropolymerized PPy, two cases have to be considered: one where the dopant counter ion is a relatively small anion, as in the case of Cl\(^{-}\), SO\(_4^{2-}\) and PO\(_4^{3-}\), and one where the dopant molecule is large enough to have an effect on the final polymer film morphology and thickness, as for the large anions PSS\(^{-}\) and DS\(^{-}\). The nominal thickness of the obtained PPy films can be estimated based on Faraday’s law of electrolysis by integrating the amperometric current-time traces for the polymerization experiments, assuming 100% current efficiency, an average polymer density of 1.5 g/cm\(^3\) and the involvement of 2 electrons in the polymerization reaction.\(^{31}\) The calculated average thickness of the obtained PPy film in the presence of small counter-ions was 44 nm ± 1 nm (s.e.m.; n=24), while in the case of large counter-ions the estimated thickness was 82 nm ± 5 nm (s.e.m.; n=12). A larger variation in thickness was found for PPy films polymerized in the presence of large dopant counter-ions.

The effect of overoxidation on the morphology of the polymer film created by electropolymerization can be seen in the SEM and AFM images shown in Fig. 2. In the case of both small and large counter ion doping, overoxidation of PPy results in flattening of the film (compare Fig. 2A,B with 2C,D and Fig. 2E,F with 2G,H). The AFM images show that the flattening resulting from the overoxidation also reduces the roughness of the film, therefore affecting its porosity. Hence, even if the overoxidation yields a loss of conductivity in the film, the thinning of this film reduces the electron transfer distance between the electrochemically active analyte and the underlying gold electrode, which retains its conductive metallic properties.

Fig. 2 clearly illustrates the differences between the morphologies when using small and large counter ions. Small dopant ions make the resulting PPy film tightly packed onto the surface of the electrode (Fig. 2A,B), whereas large dopant ions render the PPy film relatively rough with a more amorphous surface (Fig. 2E,F). This visual investigation of the PPy surfaces can be related to differences in the calculated thickness of the PPy films doped with small and large counter ions.

A characteristic increase in sensitivity to dopamine due to overoxidation of PPy can be seen in Fig. 3, which shows cyclic voltammograms of a PPy-modified gold IDE set before and after overoxidation. Before overoxidation, the neutral state PPy modification results in voltammetric peaks corresponding to dopamine oxidation and reduction hidden under currents generated by redox processes of the polymer and non-faradic processes caused by an increased double layer capacitance, making the anodic current rise with an almost ohmic behavior. After overoxidation of the PPy film, dopamine electron transfer occurs directly to the underlying gold IDEs. The effect of the double layer capacitance is drastically decreased and the faradic processes of the neutral state PPy are eliminated, which facilitates...
observed response in comparison with the ones recorded using bare gold IDEs. This lack of difference between signals from a non-doped IDE and a small counter ion doped PPy film can be explained by the overoxidation process, which, as mentioned above, introduces electronegative functionalities in the film. These can repel small negative counter ions entrapped in the polymer matrix, whereas large counter ions remain entrapped in the polymer film, further enhancing the attraction to positively charged analytes, such as dopamine.

The dopant counter ion that contributed to the highest electrode response and hence to the greatest sensitivity to dopamine in all experiments was PSS; clear peaks could be observed in cyclic voltammograms down to dopamine concentrations in the nM range (see Fig. 4B). The observed increase in response to dopamine is related to the formation of carbonyl- and carboxylic acid functionalities. The effect of these functionalities can be seen in an analogous manner as has been shown in the case of thiol SAMs as a modification to improve dopamine electrochemistry (thiol SAMs having a partially deprotonated carboxylic tail group, e.g., thiocetic acid and mercaptopropionic acid). It is believed that the presence of carbonyl- and carboxylic acid functionalities orient the electroactive hydroxyl groups of dopamine molecules toward the electrode surface, in this way enhancing the oxidation of dopamine.

In the case of overoxidized PPy, an additional effect can be proposed to further explain the observed current amplification during oxidation of dopamine. Despite the fact that the overoxidized PPy film becomes morphologically flattened in comparison with the initial conducting PPy film, it is a 3-dimensional polymer network comprising carbonyl- and carboxylic acid functionalities as well as retained counter ions (PSS) with dispersed negative charges. This structure can more effectively accommodate the diffusing dopamine in the vicinity of the electrode surface, increasing the fraction of oxidized dopamine.

Aside from simply orienting the hydroxyl group of dopamine molecules toward the electrode surface or accommodating more dopamine molecules in the vicinity of the electrode surface, the 3-dimensional structure with dispersed negative charges can also decrease the freedom of movement of the tail of dopamine molecules with the protonated positively charged amino group. This can be seen as a combined steric and electrostatic hindrance, limiting the possibility of intramolecular cyclization, which is the prerequisite for polymerization of dopamine on an electrode surface. Hence, overoxidized PPy can serve as a protection against electrode fouling that otherwise could decrease current response. However, since steric hindrance in the packing of the PPy matrix may leave gaps between the individual PPy chains on the electrode surface, there is, nevertheless, a possibility for electrode fouling, which can be observed during continued potential cycling (results not shown).

**Exocytosis measurements**

Since the optimization experiments showed that a PSS-doped overoxidized PPy film is, among all the options tested in this work, the best electrode modification in terms of sensitivity, this specific modification was chosen for the detection of cellular dopamine release. PC12 cells were chosen because of their wide use as models to study neuronal functions, such as exocytosis. This cell line was created several decades ago from a rat adrenal pheochromocytoma with the aim of creating a cell model for neurochemical and neurobiological studies. These cells are known for their reliable release of dopamine, and have been used for implantation into neuronal tissue. 100,000 cells/cm² was chosen as the cell density for seeding PC12 cells onto the electrode chip (seen in Fig. 1B) in order to facilitate differentiation for several days without reaching too high a confluency. This precaution is significant in order to allow proper formation of the characteristic axon-like outgrowths. As shown in Fig. 5A, the PPy electrode modification did not hinder proper cell differentiation. Since PC12 cells require an adhesion factor on a growth substrate, the PPy-modified electrodes were coated using physisorbed laminin, which has been shown to facilitate good differentiation of different cell types, including neurons and PC12 cells. A PC12 cell population differentiating on one set of IDEs can be seen in Fig. 5B. The PC12 cells extended the axon-like outgrowths on the IDEs, seemingly preferring the modified electrode surface to the rest of the chip (see Fig. 5C). Cells on the laminin-modified electrodes showed the typical cell adhesion, differentiation morphology and confluency expected for PC12 cells. This electrode configuration is optimal for accommodating the axon-like outgrowths when monitoring release of neurotransmitters from a whole population of differentiating cells. The function of population-based exocytosis monitoring from growing non-differentiated PC12 cells using 30-µm disk electrodes has been
demonstrated previously.\textsuperscript{41} However, such an electrode structure and dimension is limited to primarily address the cell bodies and not the axon-like outgrowths. Hence, disk electrodes are mainly suitable for recordings from growing PC12 cells, which have round cell bodies. Even when the diameter of disk electrodes is increased to 100 µm,\textsuperscript{42} a population of differentiating PC12 cells cannot be accommodated well on them due to the elongated morphology of the cells’ outgrowths.

Fig. 6 depicts amperometric current-time traces generated on IDEs during a measurement, as well as the baseline for a control measurement where no cells were present on the IDE. As expected, the amplitude of the recorded current generated by dopamine oxidation on an IDE modified with overoxidized PPy-doped PPy (PPy/PSS) was significantly higher than that obtained on a bare gold IDE. In detection of single vesicle exocytotic events, various parameters can be extracted from recorded current-time traces (spikes), such as the integrated area of the current spikes (the charge Q yielding the amount of molecules released during each event), the maximum oxidation current, half-width \(t_{1/2}\), and the rise-time.\textsuperscript{43} In measurements of dopamine release from a cell population, the recorded current-time traces are composed of a large number of superimposed single vesicle events. In this case, the most significant parameter is the total charge, which, in an analogous manner as in the case of single vesicle measurements, is proportional to the detected amount. The average generated charge on bare gold IDEs having an average cell population of 217 ± 5 cells (s.e.m.; \(n=10\)) (based on visual counting under microscope) was 8.7 ± 0.6 nC (s.e.m.; \(n=5\)), while on PPy/PSS modified IDEs, having cell populations of the same dimension as on bare gold IDEs, the average generated charge was 23 ± 2 nC (s.e.m.; \(n=5\)).

The different quantities of charge generated on bare gold IDEs and PPy/PSS modified IDEs upon dopamine detection do not, however, mean that the cells residing on bare gold electrodes and PPy/PSS modified electrodes released different amounts of dopamine. Instead, the PPy/PSS modified IDEs show an increased sensitivity to dopamine, indicating 2.6-fold amplification. This is in accordance with the value previously published for PPy modified CFMEs.\textsuperscript{30} It is significant to point out that measurements conducted on bare electrodes and PPy/PSS modified electrodes are not directly comparable since, for cell populations of the same dimension, different levels of charge are being recorded. On the other hand, the increase in detection sensitivity makes the electrode modification highly suitable for monitoring dopamine release under conditions, where very small changes need to be detected, such as in vivo monitoring of the dynamics of dopamine release in the rat brain\textsuperscript{30} and characterization of conditions leading to differentiation of neural stem cells to dopaminergic neurons.\textsuperscript{22,23}

The results presented in this paper highlight the novelty of overoxidized PPy as an electrode modification for detection of cellular dopamine release. The modification is readily prepared, resulting in a reproducible electrode behavior (for details of the effect of the prerequisite electrode cleaning, see ESIF), and improves the capability to monitor small amounts of released molecules. Furthermore, the modification is stable under cell culture conditions for long-term experiments and shows no adverse effects on the differentiated cells.

Although PSS as the dopant clearly provided the highest amplification in dopamine detection, the results obtained in this work when using DS as the dopant also indicated amplification. This is in accordance with the previously published findings.\textsuperscript{25} However, the dimension of DS ion is very critical. Due to its size, it has an increased possibility to leak out from the overoxidized PPy film during long-term experiments, which can cause harmful effects on the cultured/differentiated cells. This was indicated as decreased and irreproducible cellular dopamine release (results not shown). Hence, for a reliable application of overoxidized PPy as an electrode modification for monitoring cellular dopamine release, PSS is considered the optimal dopant among those tested in this work during PPy electropolymerization.

### Experimental

#### Chemicals

- Potassium chloride (BioXtra), sodium sulfate (ReagentPlus), monobasic potassium phosphate (cell culture tested), sodium dodecyl sulfate (DS, Mw 288.4/g/mol (molecular biology grade), poly(sodium 4-styrenesulfonate) (PSS) (average Mw ~70,000 Da, 30% wt. in water), hydrochloric acid (BioReagent), magnesium chloride hexahydrate (BioReagent), calcium chloride dihydrate (BioReagent), sodium chloride (BioReagent), glucose (BioXtra), 4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES) (1M solution; BioReagent), 2-(3,4-dihydroxyphenyl)ethylamine hydrochloride (Dopamine), 3-(3,4-dihydroxyphenyl)-L-alanine (L-DOPA), sodium hydroxide (BioXtra), pyrrole (Reagent), hydrogen peroxide (30% in water), potassium hydroxide (semiconductor grade), polyethyleneimine (PEI) (branched, average Mw ~25,000), nerve growth factor-β from rat (NGF), laminin from Engelbreth-Holm Swarm murine sarcoma basement membrane, horse serum (HS), glutaaraldehyde (25% in H2O, specially purified for use as an electron microscopy fixative), and water (cell culture tested) were purchased from Sigma-Aldrich Corporation (St. Louis, MO, USA).
- Dulbecco’s Modified Eagle Medium (DMEM)/Ham’s Nutrient Mixture F12 with GlutaMAX (DMEM/F12), Trypsin-EDTA (0.05%) and penicillin/streptomycin (P/S) were purchased from Life Technologies Ltd (Paisley, UK). Foetal bovine serum (FBS) and cell culture tested phosphate buffered saline (PBS) were purchased from Biowest S.A.S. (Nuaillé, France) and Lonza Group Ltd (Basel, Switzerland), respectively. All solutions used for electropolymerization and electrochemical characterization were prepared in ultra pure water (resistivity 18.2 MΩ·cm) from a Milli-Q® water purification system (Millipore Corporation, Billerica, MA, USA).

#### Solutions, buffers and media

Electropolymerization was done using 0.05 M pyrrole (used as received) solution in water. Dopant counter-ions were added in a 10-mL aliquot of the pyrrole solution to obtain concentration of 0.1 M (Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, PO\textsubscript{4}\textsuperscript{3−}, or DS) or 10 mg/mL (PSS). Dopamine solutions for sensor characterization were prepared by diluting a 1 mM stock solution prepared in nitrogen purged PBS (pumped for at least 20 minutes before use). Purging of the stock solution was continued throughout the sensor characterization. Each dilution (e.g. 100 µM, 10 nM) was prepared immediately before use. Buffers for exocytosis measurements were prepared using cell
culture tested water. The low K\textsuperscript+ buffer contained 10 mM HEPES, 5 mM glucose, 1.2 mM Mg\textsuperscript2+, 2 mM Ca\textsuperscript2+, 150 mM Na\textsuperscript+ and 5 mM K\textsuperscript+. In the high K\textsuperscript+ buffer, the concentration of Na\textsuperscript+ and K\textsuperscript+ were 5 mM and 450 mM, respectively. Both buffers were sterile filtered before use. All cell media were prepared using DMEM/F12. The growth medium was supplemented with 15% HS, 2.5% FBS, 1% P/S, and 0.5% HEPES. The differentiation medium had a lowered serum contents (0.5% HS and 0.5% FBS). The L-DOPA (100 µM) medium was prepared by adding L-DOPA from a 10-nM stock solution in PBS into the differentiation medium. The stock solution was prepared by first dissolving L-DOPA (40 mg/ml) in 0.5 M HCl followed by dilution with PBS and sterile filtration.

Electrode microchip fabrication

The microchips having 12 electrode sets, each comprising an interdigitated electrode (IDE), a counter electrode (CE) and a reference electrode (RE), all in gold (see Fig. 1), were fabricated using standard cleanroom-based micromachining techniques. A 670 nm layer of thermal oxide (SiO\textsubscript{2}) was grown on a silicon wafer (one side polished) in a drive-in furnace at 1050 °C in the presence of water vapor (wet growth). The gold structures, i.e., electrodes, leads and contact pads, were photolithographically defined in positive photoresist (AZ\textregistered 5214E from MicroChemicals GmbH, Ulm, Germany) using an image reversal process, followed by metal deposition (10 nm thick Ti adhesion layer and 150 nm thick gold layer) through electron beam evaporation and lift-off in acetone. Prior to metal deposition, the SiO\textsubscript{2} was etched for 100 s in the areas having opened resist using buffered HF solution to form ca. 150 nm deep isotropic undercuts\textsuperscript{46} that eliminate the formation of lift-off ears at the edges of the metal structures. The non-active gold areas (the leads connecting the electrodes to the contact pads) were passivated with 500 nm thick silicon nitride deposited using plasma-enhanced chemical vapor deposition. A second photolithography step coupled with reactive ion etching of the silicon nitride was used to expose the active electrode areas and contact pads. Removal of the final photoresist was achieved by ultrasonication in acetone followed by intermediate rinsing with ethanol and final rinsing with deionized water.

Polypyrrole electropolymerization, overoxidation and characterization

For the electropolymerization and electrode characterization, the electrode chips were placed in a micromilled poly(methyl methacrylate) (PMMA) holder (for details see ESI\textsuperscript{5}). The holder formed a 500-µL vial on top of the electrode chip to facilitate liquid handling during experiments. Interconnections between the electrode chip and the potentiostat (1010A eight-channel potentiostat from CH Instruments, Austin, TX, USA) were obtained using a tailor-made PCB having gold plated spring loaded pins (Mill-Max Mfg. Corp., Oyster Bay, NY, USA). Before use, the electrode chips were cleaned for 10 minutes in a solution containing 50 mM KOH and 25% H\textsubscript{2}O\text subscript{2}, followed by a potential sweep on the IDE working electrodes (WE) from -200 mV to -1200 mV in 50 mM KOH at 50 mV/s to remove the gold oxides formed during the chemical cleaning.\textsuperscript{47} The polypyrrole (PPy) electrode modification was achieved by potentiostatic electropolymerization (10 seconds at 700 mV) with or without dopant counterions. The electropolymerization potential was chosen to obtain a neutral state PPy film before proceeding with the overoxidation, which occurs at around 900 mV (data not shown). The IDEs of the microchip were used as WEs together with the RE and CE from each respective electrode set. Overoxidation of the PPy layer was carried out by cyclic voltammetry in a 0.1 M NaOH solution by applying 50 cycles in a potential window from 0 to 1000 mV at the sweep rate of 100 mV/s. The electrochemical behavior of the doped PPy-modified electrodes was characterized by cyclic voltammetry in a potential window from 0 to 700 mV at a potential sweep rate of 50 mV/s in solutions having various dopamine concentrations. A morphological comparison of the polymer layer was done by low-vacuum scanning electron microscopy (SEM) using a FEI Nova 600 NanoSEM system (FEI Company, Oregon, USA) at 5 kV (spot size 4.0). Atomic Force Microscopy (AFM) characterization was carried out with a Veeco di CP-II microscope (Veeco Instruments Inc., Plainview, NY, USA). The results showing the average thickness of the obtained PPy films are presented as ± standard error of mean (s.e.m.), with n being the number of characterized IDEs.

Cell culturing and differentiation

The cell-based experiments were conducted using rat pheochromocytoma (PC12) cells from Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (Braunschweig, Germany). Initially, PC12 cells were grown for 48 hours in PEI-coated T25 culture flasks from Nunc A/S (Roskilde, Denmark) using growth medium. PEI coating was done at room temperature for 2 hours using sterile filtered 50 µg/mL PEI diluted in PBS followed by rinsing twice with PBS. 24 hours prior to seeding the cells onto an electrode chip, the growth medium was changed for the differentiation medium to initiate the cellular response to NGF. Sterilization of all microfabricated cell culture substrates and materials was done by immersing them in 0.5 M NaOH for 30 minutes followed by rinsing with PBS trice. Electrode chips were coated with laminin (20 µg/mL laminin diluted in PBS) in a Petri dish for 2 hours to ensure cellular adhesion. Adsorption of laminin, and hence adhesion of cells, only in the central part of an electrode chip where the electrode sets are located was achieved using a hydrophobic pen from Dako Denmark A/S (Glostrup, Denmark). The pre-differentiated PC12 cells were rinsed with PBS and trypsinized for 5 minutes followed by 5-minute centrifugation at 850 rpm at 20 °C. The cell pellet was resuspended in the differentiation medium and the cells were seeded onto the coated electrode chips with a surface density of 10\textsuperscript{6} cells/cm². All cell culturing and differentiation was done in an incubator at 37 °C in a humidified atmosphere of 5% CO\textsubscript{2}/95% air. SEM imaging of the differentiated cells was done using a FEI Nova 600 NanoSEM system after fixation for 1 hour with 2% glutaraldehyde solution diluted in PBS followed by rinsing with PBS (twice for 15 min) and cell culture tested water (twice for 5 min). Cell counting on microchips was done using a Zeiss Axios Imager M1m microscope equipped with an AxioCam MRcS computer controlled CCD camera (Carl Zeiss AG, Göttingen, Germany).

Exocytosis measurements

After 4 days of differentiation, the differentiation medium was
replaced by the L-DOPA medium and the cells were kept in the incubator for one hour to increase the dopamine load in the vesicles. Before conducting the exocytosis measurements, each electrode chip with the differentiated PC12 cells was placed in a micromilled PMMA holder (for details see ESI†) to facilitate contact between the individual electrode sets and a tailor-made 24-channel potentiostat with a dedicated acquisition software. The open vial in the holder facilitated the addition of the necessary buffer solutions during exocytosis measurements. The medium was immediately replaced by 160 μL of the low K+ buffer to record a baseline for the measurements. Each array of interdigitated WEs was poised at 400 mV vs. the gold RE adjacent to the array. After a stable baseline had been recorded, exocytosis was triggered by pipetting 80 μL of the high K+ buffer directly into the vial to elevate the K+ concentration to 150 mM. The current-time traces corresponding to the oxidation of the dopamine released by the cells were obtained shortly after triggering the exocytosis. Recording of the exocytotic events was done simultaneously on each IDE array. Exocytosis experiments were carried out on 3 different electrode chips, each having non-modified IDE arrays and IDE arrays modified with overoxidized PSS-doped PPY. All the amperometric recordings were done at room temperature (~22 ºC). All calculated results from cell-based measurements are presented ± (s.e.m.), with n being the number of IDEs having cell populations subjected to exocytosis measurements.

Conclusions
This work demonstrates the obtained enhanced sensitivity in dopamine detection by using polystyrene sulfonate (PSS) doped electropolymerized and overoxidized polypyrrole (PPy) film as an electrode modification. The presented investigation on how overoxidation and doping of the conducting PPY film with different counter-ions influences the resulting morphology and the sensitivity to dopamine led to the proposal of three possible factors contributing to the enhancement of dopamine detection: a three-dimensional polymer network with oxygen functionalities i) can guide the orientation of the catechol moiety, facilitating an improved electrochemical behavior previously shown for thiol modifications, ii) increase the fraction of detected dopamine, resulting in current amplification, and iii) decrease the possibility of electrode fouling caused by dopamine polymerization. The application of PPY/PSS electrode modification to detection of released dopamine from populations of differentiated rat pheochromocytoma (PC12) cells demonstrates how the optimized and characterized electrode modification provides a basis for future biological studies on, for instance, characterization of neural stem cell differentiation to dopaminergic neurons.

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Notes and references

References

Figures and Figure Captions

Fig. 1 A) An individual set of reference electrode (RE), working IDEs (WE) and counter electrode (CE); B) a microfabricated chip containing 12 sets of electrodes with individual contact pads on the outskirts of the chip for electrical connections (for a complete view of the analytical system setup, see ESI†).

Fig. 2 SEM (A, C, E, G) and AFM (B, D, F, H) images of electropolymerized PPy films on IDEs, showing typical morphologies for A, B) small-counter ion doped films (in this case Cl\textsuperscript{−}); C, D) overoxidized small-counter ion (Cl\textsuperscript{−}) doped films; E, F) large-counter ion (PSS\textsuperscript{−}) doped films; G, H) overoxidized large-counter ion (PSS\textsuperscript{−}) doped films.
Fig. 3 Typical cyclic voltammograms of a polypyrrole modified gold IDE set in 100 µM dopamine (in PBS) before and after overoxidation (no dopant counter-ions were added). The integrated RE and CE for the IDE set were used to obtain the three-electrode configuration. The potentials were adjusted vs. gold RE, the potential sweep rate was 50 mV/s (sweep direction from 0 to 700 to 0 mV).

Fig. 4 A) Comparison of typical cyclic voltammograms in 100 µM dopamine (in PBS) from sets of IDEs modified with overoxidized PPy doped with different counter ions (Large Counterion refers to PSS, and Small Counterion to PO₄³⁻). The integrated RE and CE for each IDE set were used to obtain the three-electrode configuration. The potentials were adjusted vs. gold RE and the potential sweep rate was 50 mV/s (sweep direction from 0 to 0.7 to 0 V). B) Voltammogram obtained from the overoxidized PPy/PSS modification as a response to 10 nM dopamine.
**Figure 5**

Fig. 5 SEM images of differentiated PC12 cells on PSS-doped overoxidized PPY-modified electrodes showing A) a single PC12 cell, B) a population of PC12 cells and C) a magnified view of the cell population in B.

**Figure 6**

Fig. 6 Typical amperometric current-time traces, corresponding to dopamine release from differentiating PC12 cells, obtained during an exocytosis measurement on a bare gold IDE (red line, Au) and on an IDE modified with PSS-doped overoxidized PPY (blue line, PPY/PSS). The black line (CONTROL) corresponds to the signal recorded on an IDE without PC12 cells. Inset shows the average charge accumulated during measurements on non-modified (Au) and modified IDEs (PPY/PSS). The error bars represent s.e.m. (n = 5).
A microelectrode modification was developed using electropolymerized polypyrrole, testing several counter ion dopants for sensitivity increase towards dopamine, and used for the detection of dopamine from a population of cells.