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Human Stress Monitoring Through an Organic Cotton-Fiber Biosensor

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Selective detection of bioanalytes in physiological fluids, such as blood, sweat or saliva, by means of low-cost and non-invasive devices, is of crucial importance to improve diagnosis and prevention in healthcare. To be really useful in everyday life a sensing system needs to be handy, non-invasive, easy to read and possibly, wearable. Only a sensor that satisfies these requirements could be eligible for applications in healthcare and physiological condition monitoring. Herein an organic electrochemical transistor has been investigated as a simple, low-cost and e-textile biosensor, fully integrated on a single cotton yarn. The biosensor has been used for real-time detection of adrenaline, selectively compared to the saline content in human physiological fluids. The sensing mechanism is based on the oxidation of adrenaline at the Pt-gate electrode surface, with the formation of adrenalonequinone and adrenochrome. Two independent organic electrochemical transistors, characterized by different gate-electrode materials, detect saline and adrenaline concentration, respectively, in real human sweat. Measurements performed in real-time mode show the complete independence of adrenaline detection from NaCl and, hence, guarantee the simultaneous monitoring of both concentrations. The oxidation of adrenaline has been studied by means of absorption spectroscopy in air, with either silver and platinum working electrodes. Our results confirm that the oxidation reaction driven by the Pt-electrode leads to the formation of adrenochrome, while, with the Ag-electrode the oxidation is similar to the spontaneous one occurring in air. The cotton-based biosensor shows the possibility to monitor human performances (hydration and stress) in-situ and with a non-invasive approach, opening new unexplored opportunities in healthcare, fitness and working safety.

Introduction

Performing a continuous and selective detection of bioanalytes in physiological fluids, without invasive and complicated sampling-laboratory analysis, is an open challenge in prevention and diagnostics for healthcare applications. Hence, the idea to develop a wearable bio-sensing system, integrated in clothes, light, handy and non-invasive, could be an appealing solution. A wearable monitoring system offers several advantages, for example the prompt detection of molecules addressable to specific diseases or critical physiological state, as well as the ability to perform an early preventive diagnosis of a large number of pathological conditions. Moreover, the continuous monitoring of physiological parameters, like glucose, lactate and saline concentration through a low cost, non-invasive device, allows an active control on patients’ health and/or sport performances. Organic electrochemical transistors (OECT) are emerging as a feasible solution in biosensing applications thanks to their low cost, non-invasive and easy to read features. So far OECTs have been applied as sensors for simple analytes such as hydrogen peroxide, ions and with more complex bio-structures such as micelle, liposomes, DNA, and pigments. Extensive reviews about OECT bio-integration have been reported in literature recently. In addition, OECT can be easily integrated on cloths, realizing efficient e-textile organic devices. In recent times different approaches for the functionalization of cotton yarns (natural cellulose) have been used to realize conductive fibers, which have been applied as innovative textile-integrated sensing devices. Among the different strategies for the realization of an active layer, conductive polymeric materials, metal nanoparticles, or carbon nanotubes (CNTs), have been successfully employed. The output characteristics of textile-OECT have been demonstrated for the first time using a gel electrolyte for ionic exchange. Successively cotton-OECT demonstrated to work in liquid solutions and they were applied for monitoring saline concentration in human sweat. Despite of the high sensitivity and the very simple and handy structure, OECT actually lacks the required selectivity for the recognition of characteristic molecular species. Different approaches have been attempted to overcome this limitations, for example, a molecular membrane surrounding the active layer has been used to selectively detect potassium ions or lipid membrane allows to discriminate between monovalent and divalent ions.
In this paper we present the selective detection of adrenaline with respect to the saline content in human physiological fluids. Adrenaline is a functional neurotransmitter, playing a central role in many instinctive responses, especially under stress situations and strongly physical strengthens. A timely sensing of abnormal adrenaline concentration could be a fingerprint of a pathological situation, like panic or heart attack, or could identify a typical flight, fight and fright response. Moreover it could be used to monitor athletes, where the control of physiological performances during competition and training is required. The adrenaline sensing in a complex fluid (human sweat) is herein reported for the first time using OECT, enabling real-time monitoring of human physiological parameters. To detect adrenaline in an independent way with respect to the saline content, an innovative system of two OECTs, completely integrated in cotton fibers have been proposed. The sensing mechanism of adrenaline relies on the oxidation process at the platinum metal gate, leading to the formation of different intermediates and eventually adrenochrome. To make a step further towards real on-field, application, we applied the device to the study of real human sweat: measurements have been recorded in real-time using human sweat as electrolyte and monitored the OECT sensing upon injection of adrenaline during acquisition, showing the complete independence of adrenaline sensing respect to salt sensing. This innovative device is an useful tool for an in-situ and non-invasive analysis of human performances (hydration and stress), finding application in sports, health care and working safety.

Different analytical methods for the detection of adrenaline in human physiological fluids, like blood, sweat\textsuperscript{17} or saliva\textsuperscript{18} have been approached, including liquid chromatography (LC),\textsuperscript{19} spectrophotometry,\textsuperscript{20} fluorimetry,\textsuperscript{21, 22} capillary electrophoresis (CE),\textsuperscript{23, 24} amperometric,\textsuperscript{25} chemiluminescence (CL)\textsuperscript{26, 27} and electrochemiluminescence\textsuperscript{28} detection. Nevertheless, despite all these methods are regarded as highly sensitive, are also expensive, not portable and can perform off-line analysis only. The viability of methods and devices for a fast and sensitive detection of adrenaline in biological fluids is, on the other hand, desirable for daily screening and preventive diagnosis. So a wearable and highly sensitive device sensor could be also more appealing and readily applicable to patients or sport men.

**Experimental**

**Device fabrication**

The OECT device on cotton (hereafter referred as the cotton-OECT), has been fabricated on a cotton fiber (the yarn) by a simple soaking process, as reported in \textsuperscript{12}: the yarn was immersed into an aqueous solution of PEDOT:PSS (CleviosPH500, Starck GmbH) for 5 min, followed by a baking on hot plate at 150 °C for 3 hours. The PEDOT:PSS solution was previously modified with addition of ethylene glycol (20%) and dodecyl benzene sulfonic acid (DBSA) surfactant (12%) in order to increase electrical conductivity and decrease solubility in water.\textsuperscript{15, 20, 30} The process did not change significantly the flexibility of the cotton yarn (see Supporting Informations). The measured electrical resistance of the as-prepared yarn was 430 Ohm over a linear length of 1 cm: that value allows the use of voltages in the mV range. Figure 1a shows a FE-SEM image of a thin layer of PEDOT:PSS surrounding the cotton yarn. The layer appears uniformly distributed at the nanometer scale with an estimate thickness of about 50 nm; few layer borders could be seen on the side of the yarn. A drop of about 200 μL of the electrolyte solution is placed on the yarn between the source and drain contacts, allowing the liquid to reach the gate electrode. The channel of the cotton-OECT is defined by the overlapping of the liquid electrolyte with the polymer, as evidenced by the darker area on the cotton fiber reported in Figure 1b. After processing with PEDOT:PSS, the yarn still maintains its flexibility and can be easily integrated on cloth. In this respect, it is important to note that the output characteristics are not altered under bending. This property is very important for real textile applications, for example in the case of integration of the device in shirts used in fitness or daily routines.

To realize a cotton-OECT sensor with a selective sensing capability, two OECTs were integrated on the same fabric patch, very close each other but working independently. The devices are shown side by side in Figure 1b, one with a silver (Ag) wire as the gate electrode, the other with a platinum (Pt) wire. As previously described the black yarn at the bottom side of each device is the cotton fiber coated with the PEDOT:PSS polymer. The two cotton-OECTs can detect simultaneously and independently different kind of analytes; in particular the Ag cotton-OECT is sensitive to ions,\textsuperscript{3, 4} while the Pt-OECT is capable to react with adrenaline molecule. Electric measurements were performed contemporary on both devices, taking the electrolyte from the same common bath. In order to monitor the oxidation process of adrenaline at the Pt-gate electrode surface, spectroscopic investigations were also performed: absorption spectra (Jasco V-530 spectrophotometer in the 250-
650 nm range with 1 nm step) were recorded as a function of time to better understand the role of OECT detection in the oxidation rate of adrenaline. A 1 mM adrenaline solution in 0.1 M NaCl was used for monitoring the adrenochrome evolution, considering both the spontaneous adrenaline oxidation, as well as during OECT operation.

Device characterization

Figure 1c shows a schematic view of the cotton-OECT electrical circuit and the sensing mechanism involved in the detection of adrenaline: adrenaline undergoes an electro-oxidation to adrenaline-quinone and to adrenochrome at the surface of Pt-gate electrode (Figure 1d); as a consequence a faradic current flowing in the source-gate circuit is generated and hydrogen protons are released in the solution. The OECT response depends on the potential drops occurring at the gate/electrolyte and electrolyte/polymer interfaces. Under the OECT faradaic regime of operation the potential drop at the electrolyte/gate interface decreases and the effective gate voltage ($V_{gs,eff} = V_g + V_{g,drop}$) increases, the latter forcing H⁺ cations to move toward the polymer surface and de-dope PEDOT:PSS. The sensing mechanism is similar to the OECT-based sensor for hydrogen peroxide, dopamine or melanin pigment reported recently, meaning that catecholamine molecules, like adrenaline, are suitable to be studied by OECT technology.

The output currents were acquired by sweeping $V_{ds}$ between 0 and -0.5 V with steps of 0.1 V, at fixed $V_{gs}$ values between 0 and 0.5 V ($V_{gs}$ steps of 0.1 V). Transfer characteristics of OECT were acquired by measuring $I_{ds}$ vs. $V_{gs}$ using $V_{ds} = -0.2$ V and pulsing $V_{gs}$ between 0 V and 0.5 V with 0.1 V steps. They have been expressed as current modulation $\Delta I_{ds}/I_{ds,0}$ vs. $V_{gs}$, $I_{ds} = -0.2$ V.

The current values were determined from current transient measurements considering the quasi-steady state current level. The application of $V_{ds}$ induces a drift of the holes along the PEDOT:PSS channel, generating a drain-source current ($I_{ds}$). Upon application of a positive gate voltage ($V_{gs}$), cations ($Y^+$) from the electrolyte enter the PEDOT:PSS channel causing its de-doping according to the Equation 1:

$$\text{PEDOT}^+:\text{PSS}^- + Y^+ + e^- \leftrightarrow \text{PEDOT}^0 + Y^+: \text{PSS}^- \quad (\text{Eq. 1})$$

In the “de-doping process” a decrease of the module of drain current $I_{ds}$ (less holes available for conduction) is induced as a consequence of cations incorporation into the PEDOT:PSS backbone and a reduction of the oxidized PEDOT to PEDOT$^0$. This process is reversible and when $V_{gs}$ is switched off ($V_{gs}=0$ V), ions diffuse from PEDOT:PSS to the electrolyte increasing the number of conducting holes and, consequently, $I_{ds}$. Such process is referred to as “doping”.

Results and discussions

Figure 2 shows the basic output characteristics ($I_{ds}$ vs. $V_{ds}$ at different gate voltages, $V_{gs}$) of the cotton-OECT device with Pt-gate electrode measured in NaCl 0.1 M.

Transfer characteristics of OECT are shown in Figure 3a for adrenaline diluted in NaCl 0.1 M at 1x10⁻³ M and 1x10⁻⁹ M concentrations. The transfer curves of the cotton-OECT with a Pt-gate electrode and characterized with two concentrations of adrenaline (1x10⁻³ M and 1x10⁻⁹ M), shift to a lower gate voltage (overall shift > 300 mV) for the higher concentration of adrenaline (1x10⁻³ M). This indicates that the effective gate voltage ($V_{gs,eff}$) is increased due to the electro-oxidation of adrenaline at the Pt gate electrode surface. The source-gate current ($I_{gs}$) was acquired simultaneously during transfer measurements because the gate current analysis provides information about electron transfer reaction occurring at the gate electrode, as well as operational mode of the OECT. The catechol-type structures belonging to the reduced state of adrenaline undergoes an oxidation at the gate electrode providing electrons to the $I_{gs}$ current of the gate circuit.

In Figure 4 a real-time acquisition of the drain-current $I_{ds}$ as a function of time is reported. The plots show the different sensing capabilities of the device to adrenaline with the Ag- and Pt-gate electrodes. Measurements have been realized using real human sweat as the electrolyte bath and injecting a 10 μl droplet of adrenaline in it at a fixed time.
In Figure 4a $I_{ds}$ has been measured with Ag-gate electrode ($V_{ds}=-0.2\,V$, $V_{gs}=+0.4\,V$): two successive injections of a 10µL droplets of adrenaline solution, at 550 s and 580 s, were done without any evident variation of the $I_{ds}$ current. In Figure 4b $I_{ds}$ has been measured by the same procedure, but with the use of a Pt-gate electrode, resulting in an evident decrease of $|I_{ds}|$ (y-axis is negative in the figure).

The sensor reacts almost instantaneously (less than 1 s) upon the liquid injection, with a relative change in the current signal ($|I_{ds}|$) of about 200% respect to the base signal. Real-time measurements are intuitive and provide clear evidence that the reaction at the electrode is completely different for the Ag or Pt gate electrodes. In a similar way, it has been demonstrated in previous publications that it is possible to detect the injection of a droplet containing NaCl salt in real-sweat with the Ag-gate electrode, while no signal change occurs with a Pt-gate. The combination of these two complementary results allows us to selectively detect a rush of adrenaline in a saline solution, like human sweat, without confusing it with a change in saline concentration, because they are detected independently by the two different electrodes.

In Figure 5 the current modulation of $I_{ds}$ (here expressed as $(I_{ds}-I_{ds,0})/I_{ds,0}$) is reported as a function of adrenaline concentration. Adrenaline has been diluted in NaCl 0.1 M and 0.01 M to show the independence of the detection of adrenaline from NaCl concentration. These values fall in the physiological range of saline concentration in human sweat, i.e. 30-60 mmol.

The OECT response has been measured at increasing concentrations of adrenaline, ranging from $10^{-8}$ M to $10^{-3}$ M. In Figure 5a the current modulation, acquired at $V_{gs}=+0.3\,V$ by using the Ag-gate electrode, is reported for the two NaCl solutions. The curves do not show any dependence on adrenaline concentration, being constant within the experimental errors.

A radically different behavior is observed when Pt-gate electrode is employed (Figure 5b). The current modulation with the Pt-gate is constant down to $10^{-6}$ M adrenaline concentration and then increases monotonically for higher values. It is worth noting that the increase in modulation is proportional to the adrenaline concentration. The dynamic sensing range at $V_{gs}=+0.3\,V$ is between $1x10^{-6}$ M and $1x10^{-3}$ M, with a signal about 2 times larger. The different concentrations of salt do not affect the sensing properties of the OECT sensor, and probe the independence of the adrenaline sensing with respect to NaCl sensing. We have previously reported that a cotton-OECT may measure the saline content of real human sweat. In that case the sensing mechanism relied on the redox occurring at the Ag-gate electrode surface. In the case of adrenaline, instead, the sensing mechanism is due to the oxidation of adrenaline at the Pt-gate surface, which makes the cotton-OECT device to work under a faradaic operation regime.

To further investigate the oxidation of adrenaline induced in the OECT respect to a self-oxidation in air, absorption spectroscopy have been performed, using a dedicate experimental setup.
Adrenaline undergoes a self-oxidation resulting in the formation of adrenaline-quinone and then of adrenochrome, through an oxidation pathway that produces different intermediates (adrenaline semi-quinone/quinone, leucochrome, leucochrome quinone). The exact determination of these intermediates is not trivial and is beyond the scope of this work; nevertheless according to UV-VIS analysis reported in Figure 6, we have found meaningful differences between adrenaline self-oxidation in air (Figure 6a) and the corresponding oxidation during OECT sensing (Figure 6c).

Figure 6. Absorption spectra recorded at different times monitoring the oxidation of adrenaline. (a) Adrenaline auto-oxidation in air. (b) Adrenaline oxidation during OECT measurement with Ag gate. (c) Adrenaline oxidation during OECT measurement with Pt gate. (d) Evolution of the characteristic adrenochrome peak at 480nm as a function of time.

Figure 6c shows the simultaneous build up of the peak at 305nm and of the broad band centered at 480nm, which are the typical features of adrenochrome. The same features emerge by a far less extent during the self-oxidation in air (Figure 6a) and during the oxidation induced by a working Ag-gate electrode in OECT (Figure 6b). Hence the oxidation induced by the Pt-gate results faster and produces adrenochrome while the Ag-gate is similar to self-oxidation. No redox reactions take place at the Ag gate and the sensor response to adrenaline is almost negligible (Figure 6a). In these cases, the characteristic peak at 305nm is not visible, but a progressive shift of the absorption maximum at 290nm towards the visible range is observed and can be addressed to the convolution of the characteristic adrenaline peak at 280nm with a new one emerging at 295nm, reported by Sun et al. in their study of adrenaline auto-oxidation. Figure 6d shows the evolution of adrenochrome as a function of time by monitoring the absorption maximum of the characteristic peak centered at 480nm: interestingly no substantial difference is measured between self-oxidation and Ag-gate OECT, while platinum significantly increases the rate of adrenaline oxidation to adrenochrome.

Conclusions

In this work a simple, low cost, completely e-textile integrated biosensor for adrenaline sensing has been realized. The device is based on a cotton yarn functionalized with PEDOT:PSS and a Pt wire used as the gate electrode. The detection mechanism is based on the oxidation of adrenaline at the Pt-gate electrode surface, with the formation of adrenaline-quinone and then of adrenochrome. A setup with two independent cotton-OECT with Pt-gate and Ag-gate electrodes gives the opportunity to detect independently the saline concentration and adrenaline concentration in real human sweat. Real-time measurements show the complete independence of adrenaline sensing from NaCl sensing and the simultaneous monitoring of both the concentrations by means of the selective detection on the two different electrodes. The oxidation of adrenaline has also been studied with absorption spectroscopy in different conditions: in air, and using either silver and platinum electrode. Our results confirm that the adrenaline oxidation reaction is driven by the presence of Pt electrode, which leads to a faster and direct production of adrenochrome, while with Ag electrode the oxidation process simply follows the spontaneous and slower oxidation of this molecule in air.

Acknowledgements

GT acknowledges the projects BioNiMed (Multifunctional Hybrid Nanosystems for Biomedical Applications) from Fondazione Cassa di Risparmio di Parma (CARIPARMA) and the N-Chem project within the CNR–NANOMAX Flagship program. NC acknowledges Marco Pola for technical assistance.

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