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ELECTROACTIVE SURFACES BASED ON CONDUCTING POLYMERS FOR CONTROLLING CELL ADHESION, SIGNALING, AND PROLIFERATION

Edwin W. H. Jager1,3, *, Maria H. Bolin1,3, Karl Svennersten2,3, Xiangjun Wang1,3, Agneta Richter-Dahlfors2,3, Magnus Berggren1,3

1Department of Science and Technology, Linköping University, SE-601 74 Norrköping, SWEDEN
2Department of Neuroscience, Karolinska Institutet, SE-171 77 Stockholm, SWEDEN
3Strategic Research Center for Organic Bioelectronics, SWEDEN

ABSTRACT

We report on a variety of electroactive surfaces for the control of in vitro cell adhesion, proliferation, and stimulation. Planar cell culture substrates have been coated with the conducting polymer PEDOT and by switching the redox state, adhesion and proliferation of MDCK epithelial cells was controlled as well as stem cell seeding density. Electronically active 3D-scaffolds based on electrospun PET nano-fibers coated with PEDOT have been used as a substrate to culture SH-SY5Y neuroblastoma cells and to induce Ca\(^{2+}\) signaling. Finally, we report on micromechanical stimulation of cells using an electroactive topography surface based on micropatterned polypyrrole.

KEYWORDS

Organic electronics, conducting polymers, cell adhesion, surface energy, PEDOT, PPy, cell stimulation.

INTRODUCTION

Traditionally, in vitro cell studies are performed using planar and rigid Petri dishes comprising the cell culture and culture medium. Cellular responses upon exposure to various biomolecules are studied by adding these in solution to the medium. This is different from the in vivo situation where cells grow in a complex 3D environment. In this environment, they receive chemical and electrical signals from neighboring cells and from the extracellular matrix (ECM). In order to better mimic nature, a number of artificial surfaces have been developed such as micro- and nanotexturing [1, 2], fibers [3, 4], and coatings [5].

Electronics operate in a fundamentally different manner from biological systems: electronics is the domain of electrons and bio-systems are the domain of ions and molecules. Organic bio-electronics aims to combine and interface these worlds by utilizing the simultaneous electronic and ionic conduction present in conducting polymers.

Conjugated or conducting polymers (CP) such as poly(3,4-ethylenedioxythiophene) (PEDOT) and polypyrrole (PPy) can be electrochemically oxidized and reduced. This will change the material properties of the polymer such as conductivity, surface energy (wettability), color, and volume. The volume change can be utilized in CP-actuators in various schemes such linear actuation, bulk volume change (piston), and bending mode by assembling the CP in a bilayer configuration. Furthermore, the CP-actuators can be miniaturized using standard photolithography and microfabrication methods [6]. In addition, the change in surface energy can be used to make wettability or surface energy switches [7]. It has been shown that this switching of the redox state of CPs influences the adhesion and proliferation of cells [8]. Finally, during synthesis of the CP a charged counter ion can be included in the polymer matrix. This counter ion can then be released upon electrochemical activation. By exchanging the counter ion for a biomolecule, electronically controlled release of substances can thus be achieved [9].

PLANAR SURFACE SWITCH

Device fabrication

When switching the redox state of PEDOT electrodes, i.e. oxidizing one side and reducing the other, the surface energy will change. Using this switching it was investigated whether the adhesion and proliferation of cells adhered on the surface could be influenced.

Electronically functionalized cell culture dishes were produced using chemical polymerization of PEDOT in petri dishes. A solution of EDOT monomers, the oxidant iron(III) p-toluenesulfonate (Tosylate), and the basic inhibitor pyridine in butanol was spin coated inside the petri dishes [10]. The petri dishes were then heated to facilitate chemical polymerization and to evaporate the pyridine. Residual iron was removed by sequential washes in butanol, isopropanol, and deionized (DI) water; thereafter the films were blow dried. The surface was then mechanically divided into two electrodes (Figure 1).

Figure 1. Functionalized petri dish consisting of two PEDOT coated electrodes that can be electrochemically switched, i.e. one electrode is oxidized and the other electrode reduced.

Upon applying a low potential of 1.0-1.5 V in the
presence of an electrolyte, such as cell culture medium, the device functions as an electrochemical circuit. At the anode the PEDOT is oxidized and at the cathode the PEDOT is reduced. Since the PEDOT is electrochromic, the redox switch can be monitored visually as well. The color of the reduced electrode changes from blue to deep blue while the oxidized electrode turns more transparent.

Using this surface switch, two cell systems were studied. We investigated electronic control of the seeding density of c17.2 neural stem cells and adhesion and proliferation of MDCK epithelial cells.

**Neural stem cells**

Adhesion is an essential parameter for stem cells. It regulates the overall cell density along the carrying surface which further dictates the differentiation scheme of stem cells towards a more matured and specified population. C17.2 neural stem cells were seeded on the biased (1.5 V) PEDOT-coated dishes. After 2-4 h hours, the cells were fixed, stained with 4’, 6-diamidino-2-phenylindole (DAPI), and counted. We found that the c17.2 cells preferred to attach to the oxidized surface (Figure 2a). The intrinsic adhesive properties of the cells were not changed by the surfaces they were presented. As the oxidation state of the conjugated polymer electrodes was controlled the seeding density could be varied by a factor of 2 (Figure 2b). Along the oxidized PEDOT:Tosylate-electrodes, a relatively lower density of, and less tightly bonded, Human Serum Albumin was observed as compared to reduced electrodes. We found that this favours adhesion of the specific stem cells studied [11].

![Figure 2](image1.png)

**MDCK kidney cells**

In order to study the effect of the redox state of the conducting polymer on epithelial cell adhesion and proliferation, Madin Darby canine kidney (MDCK) epithelial cells were seeded on the biased (1.5 V) PEDOT electrodes and incubated at 37°C for 24 h. Hereafter, the MDCK cells on the electrodes were stained for immunofluorescence microscopy analysis of adherent cells.

![Figure 3](image2.png)

The extracellular matrix protein fibronectin is important for cell adhesion. The contribution of fibronectin to the cellular response of the PEDOT redox state was investigated [12].

**FIBER SURFACE SWITCH**

The previous devices demonstrated that cellular adhesion and proliferation can be electronically controlled by changing the redox state of the conducting polymer surface. However, these devices still present a planar geometry different from the 3D environment of cells in vivo, including the fibrous ECM.

![Figure 4](image3.png)
mats were fabricated using electrospinning. Hereafter, vapor phase polymerization was employed to coat the nano-fibers with a PEDOT:Tosylate layer [10]. The nano-fiber mats were spin coated with an oxidation solution comprising Fe(III) tosylate in butanol and pyridine. Hereafter, the substrates were exposed to EDOT monomer vapor resulting in a coating of PEDOT on the nano-fibers (Figure 4).

Two PEDOT coated nano-fiber mats were mounted into a custom-made petri dish resulting in a similar set-up as the planar surface switches. Electrical contacts were made with conducting copper tape.

Undifferentiated SH-SY5Y human neuroblastoma cells were cultured on the PEDOT coated nano-fiber mats for 24-48 h. The SH-SY5Y cells adhered well and showed healthy morphology (Figure 5). These electrically active scaffolds were used to induce Ca\(^{2+}\) signaling in SH-SY5Y neuroblastoma cells. It was shown that PEDOT coated nano-fibers can be utilized as 3D host environments that combine excellent adhesion and proliferation for neuronal cells with the possibility to externally regulate their signaling, such as electrically stimulated Ca\(^{2+}\) signaling [13].

![Figure 5. SEM image of SH-SY5Y cell growing on PEDOT coated fibers. Scale bar = 10 µm.](image)

TOPOGRAPHY SWITCH

Next, electroactive mechanical control of 3D surface was introduced by developing micromechanical topography switches. Microstructured polymer surfaces were prepared for electrochemically induced morphology switching. This was achieved by exploiting the large perpendicular volume change of PPy [14]. Micropatterned PPy was combined with inert SU-8 microstructures [15]. The surface consists of SU-8 square pillars (100 µm wide and 16.8 µm high) surrounded by a PPy mesh (100 µm wide and 12.7 µm thick). In order to see the effect of the raising PPy mesh in the SU-8 pillar area, contact angles were measured. The dynamics of the topography switch upon sequential reduction-oxidation switching, were recorded \textit{ex situ} by SEM and Dektak profilometry (Figure 6).

In the microstructured surfaces redox switching results in an alteration of the PPy thickness, \textit{i.e.} the depth between the SU-8 pillars alters. In the first switch cycle, the PPy thickness was varied from 12.7 µm to 13.8 µm upon reduction to the neutral state. We clearly observed how application of the cathodic potential leads to the raising of the PPy mesh. The expansion ratio of the initial reduction-oxidation cycle was 9%. This is slightly less then what has been observed by others, where ratios of 20-30% have been measured on micropatterned PPy structures [14]. As shown in Figure 6e the volume change is not fully reversible, which is generally the case for PPy-microactuators [6].

![Figure 6. SEM images of (a) SU-8 square pillars, (b,c) PPy electropolymerised (12.8 µm) forming a mesh surrounding the SU-8 pillars (16.8 µm) (b) as-prepared and (c) after reduction. Scale bar = 100 µm. (d) Dektak profile of the micro-patterned surfaces; before (black solid line) and after addition of PPy, and after the first reduction switch. Insets in the top right corner of the SEM are (a) 148º, (b) 129º, and (c) 44º contact angles of the droplets on the as-prepared surfaces. (e) The contact angle on the surfaces at different EC states.](image)
In the second redox cycle, the corresponding values were from ~74º down to ~26º, respectively.

A second generation of topography switches dedicated for micromechanical stimulation of cells has been designed and fabricated. MDCK cells were grown overnight on fibronectin coated topography switches. Cells were then loaded with Fluor-3 and monitored using confocal microscopy. Micromechanical stimulation of the topography switch induced Ca²⁺ signaling of the seeded cells.

REFERENCES

CONTACT
* E.W.H. Jager, tel: +46-11-363446; edwin.jager@itn.liu.se