REAL-TIME MONITORING OF CELL CULTURES WITH NICKEL COMB CAPACITORS

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Abstract. The aim of the study was to present a method for assessing the condition of cell culture by measuring the impedance of cells cultured in the presence of nickel. For this purpose, an impedance measurement technique using nickel comb capacitors was used. The capacitor electrodes were made using a thin film magnetron sputtering. In the experimental part, the culture of cells of mouse fibroblasts on the prepared substrate was performed. The cell culture lasted 43 hours and showed that the presented technique allows it to be used to analyze the effect of nickel on cells.

Keywords: BioMEMS, ECIS, nickel, thin films

MONITOROWANIE HODOWLI KOMÓRKOWYCH W CZASIE RZECZYWISTYM PRZY ZASTOSOWANIU NIKLOWYCH KONDENSATORÓW GRZEBIENIOWYCH

Streszczenie. Celem pracy było przedstawienie metody oceny stanu hodowli komórkowej poprzez pomiar impedancji komórek hodowanych w obecności niklu. W tym celu zastosowano technikę pomiaru impedancji z wykorzystaniem niklowych kondensatorów grzebieniowych. Ciekawostkowymi elektrodami kondensatora wykonano metodą rozpylania magnetronowego. W części eksperymentalnej przeprowadzono hodowli komórek mysich fibroblastów na przygotowanym podłożu. Hodowla komórkowa trwała 43 godziny i wykazała, że przedstawiona technika mogła być zastosowana do analizy wpływu niklu na komórki.

Słowa kluczowe: BioMEMS, ECIS, nikiel, cienie warstwy

Introduction

Impedance measurement methods can be used to assess cell status. These techniques are non-invasive and label-free electrochemical, enabling to obtain sensitive and quantitative results. The most typical are three measurement techniques [10]:

a) impedance flow cytometry,

b) electric impedance spectroscopy,

c) electric cell-substrate impedance sensing (ECIS).

These methods are based on impedance measurement, which is defined as the complex ratio of the voltage to the current in an alternating/direct current circuit. Usually a small sinusoidal AC signal is used as excitation and the electrical current response is measured. During the measurement, biological cells are suspended in medium or adhered to a substrate. To properly analyze the cell-medium-electrode system, models of electrical circuits were created. Research work presented in this paper relates to the ECIS technique, which is based on a single-shell model, in which the cell membrane blocks the current when cells adhere to the electrodes of the substrate. Due to cell proliferation and spreading, the measured impedance changes [4]. The possibility of long-term monitoring of cell culture means that it is a method which can be widely used in biological research.

1. ECIS to monitor cell status

The technique of monitoring cell culture using impedance measurement in the ECIS system is based on the use of thin-film capacitors produced on a biocompatible substrate. This technique has already been used for various types of researches: cell apoptosis [2], cell proliferation [11], drug screening [6], toxicity testing [9], cancer research [5] and stem cell differentiation [3].

Figure 1 shows typical resistance and capacitance characteristics (impedance components) for a cell culture cycle on the example of animal fibroblasts. When cells are added to the medium, the number of cells is increased by cell growth and division. Impedance increases as a result of the increase of electrodes area covered with a non-conductive cell membrane (phase 1). The number of cells is closely related to the corresponding normalized impedance value [1]. Cell proliferation and cell apoptosis or necrosis change the morphology of the cells that cover the electrodes, and thus can be detected quantitatively. During phase 2, cell culture stabilizes. And in phase 3, cells die, their adhesion decreases, therefore in the measurement circuit electrical resistance decreases, and capacity increases.

The fabrication of electrodes requires the use of microelectronic technologies, because their size should be about the size of biological cells (~10 µm in diameter). The biocompatible test substrate with capacitor electrodes is the main sensor element of the ECIS system. Polyethylene terephthalate (PET) and polycarbonate (PC) are used as the starting substrate. Thin film electrodes are usually fabricated of gold or platinum due to their biocompatible properties and good electrochemical parameters.

The measuring capabilities of the ECIS system also allow testing the effect of the presence of various metals on cell cultures. However, it is necessary to produce dedicated test substrates with thin-film capacitors made of various materials. Some other metals offer different chemical, physical and electrical properties than gold and platinum, which are also the expensive materials. In biomedical microdevices, metal elements are also made of nickel, aluminum, tungsten, silver alloys, aluminum alloys and Indium-Tin Oxide (ITO) [7].

Fig. 1. Resistance (top) and capacitance (bottom) characteristics of mouse fibroblast cells measured by a standard ECIS sensor array at 4 kHz and 64 kHz, respectively.
The main problem in the selection of material in biomedical applications is the assessment of the impact of its presence on the tested cells or substances [8]. Biocompatibility is a very important issue, especially for implants intended for long-term use. The materials used in BioMEMS devices usually have lower requirements, but also need to be tested for toxicity, degradation, corrosion and dissolution at ambient temperature. There are many applications in which the contact of substances of biological origin with metallization is short-lived and often one-off, e.g. physiological and biochemical sensors, devices for drug delivery. In such cases, it seems advisable to use a cheaper material, even if it is less biocompatible than gold or platinum typically used. The aim of this work is to perform a test culture on the made substrate with nickel electrodes, expecting that the obtained results of electrical parameters should be similar to a typical cell culture on a commercial substrate with gold electrodes. The prepared substrates can be used to test the effect of metal on biological substances using the ECIS measuring platform to monitor the condition of cells cultured in the presence of nickel electrodes. In addition, they will replace highly biocompatible materials with a less biocompatible but cheaper replacement.

2. Technology and experimental cultures

2.1. Technology of nickel electrode array

As part of an experiment analyzing the effect on a cell culture of a material other than gold or platinum, a test substrate with nickel capacitors was produced. The preparation of test substrates required a number of works in the field of thin film structure manufacturing technology. First, technological masks have been designed based on an electrode array with eight wells. The individual electrodes were made as comb capacitors in which the width of a single finger was 200 \( \mu \text{m} \).

The biocompatible substrate was cut out of 2 mm thick polycarbonate. The key step was the deposition of a nickel layer (100 nm) in the magnetron sputtering process using a Kurt J. Lesker NANO 36™ device. The magnetron sputtering technique is based on the phenomenon of individual particles deposition of material evaporated from the source under the influence of ionized energy by a strong electric field of gas. A vacuum is required to perform the deposition process. During the process, high temperatures are not used that would damage the base material (melting or polymerization). In this case, the process temperature does not exceed 60–70°C. Standard semiconductor technology uses a silicon monocrystalline substrate and the process temperatures reach above 1000°C.

The stabilizing conditions lasted 24 hours after placing the medium in the wells. Inoculation of arrays was carried out by 0.3 cm\(^2\) per well of cell suspension at \( \sim 1.2 \times 10^5 \text{cell/cm}^2 \). When the cells will be introduced to the well, they drift to the bottom of the dish where they attach and then spread on both the nickel electrode and the polycarbonate surfaces.

2.2. Preparation of the cell culture test

An experimental culture of animal cells was performed on the prepared testing substrate. For this purpose, cells of mouse fibroblast cell line were used. It was NCTC clone 929 [L cell, L-929, derivative of Strain L] (ATCC® CCL-1™) derived from ATCC organization which were cultured according to the instruction manual in complete Eagle MEM medium (Sigma Aldrich) supplemented with 10% Fetal Bovine Serum (FBS) Good HI, in an Galaxy 170R incubator, under controlled growth conditions, constant humidity and air saturation of 5% \( \text{CO}_2 \) (Fig. 4).

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3. Results and discussion

3.1. The results of electrical parameters measurements

During the experiment, the results of measuring resistance, capacitance and impedance were collected for a signal frequency in the range 62.5 Hz – 64 kHz. The experiment lasted for 43 hours. However, only the results obtained for the first 18 hours are interpretable. The obtained values were normalized and presented as their changes during cell culture. Each parameter reading is plotted as a point, in ohms for resistance and impedance or nanofarads for capacitance, in dependence of time.
Fig. 5. Resistance response measured by a nickel sensor array at 62.5 Hz for 18 hours.

Fig. 6. Impedance response measured by a nickel sensor array at 4 kHz for 18 hours.

Fig. 7. Capacitance response measured by a nickel sensor array at 64 kHz for 18 hours.

Fig. 8. Impedance response measured by a nickel sensor array at 4 kHz for 45 hours.

Fig. 9. Damaged nickel capacitor electrode without cells in the reference well after 67 hours (24 h stabilization of medium conditions and 43 h cell culture).

Fig. 10. Damaged nickel capacitor electrode with cells after 67 h (24 h stabilization of medium conditions and 43 h cell culture).

3.2. Metallization layer damage

After more than 20 hours of the cells growing, there was a sharp increase in resistance (Fig. 8) and impedance. Cell culture was continued for up to 43 hours despite receiving unusual results. Of electrode coverage. The decrease in capacity (Fig. 7) achieved during the initial 18 hours with increasing resistance should be interpreted as cell proliferation and in this respect both values complement each other and should be analyzed in parallel as a standard.
After the experiment microscopic images of the capacitor electrodes were taken (Fig. 10). They showed that during cell culture the metallization layer lost its adhesion to the substrate, which resulted in a rapid increase in the measured resistance.

However, during this time the second phase is visible, i.e. the stabilization of cell culture. After about 42 hours, the third phase (4 and 8 wells) begins, which is characterized by a decrease in the value of the measured resistance.

4. Conclusions and future work

In this paper, the developed measuring substrate technology with thin-film nickel capacitors is demonstrated. The substrates were checked for compatibility with the ECIS system. To verify that the presence of nickel would allow monitoring of vital functions of cells, the L929 cell line was performed. The obtained results of the electrical parameters show that after adding the cells to the wells, a typical cell culture cycle can be observed. This indicates the possibility of using a substrate with nickel capacitors to analyze the effect of this metal on cell culture. And in the future it will allow the use of cheaper material in many applications.

References

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