Toxicity studies of agarwood essential oil in vero cells using electrical impedance sensor

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INTRODUCTION

This article is an extension of the work presented during the IMEDITEC17 Conference in September 2017. This article provides more comprehensive details on the previous work as the study developed further by comparing the response of the natural product against Vero cells with our previous testing of natural product against breast cancer cell lines, MCF7 (Abbas et al., 2017).

Natural products have traditionally been used for medicinal purposes in Asian communities. Toxicity studies typically use animal testing to predict the harmfulness of a particular substance to human health. For this study, in lieu of animal testing, we utilized cell-based biosensors to evaluate the toxicity of natural products. The cell-based biosensors were fabricated on a printed circuit board with copper electrodes and equipped with PDMS cell culture chambers. Two different electrodes (interdigitated and circular) were designed. Vero cells were used to represent normal healthy cells. The cells were first cultured on the biosensors and then inoculated with natural products. Taxol (chemo drug – positive control) and DMSO (negative control). Impedances of these biosensors were afterwards recorded at six-hour intervals for 80 hours to determine the growth of the cells. It was found that as compared to Taxol, natural products have substantially low toxicant values.

Keywords: Impedance biosensor, natural product, agarwood, vero cells, AD5933

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study found that when wounded, agarwood trees initiate a defence system that produces secondary metabolites that were proven to have health benefits, which includes terpenes, phenolics and alkaloids (Mohamed et al., 2012). Researchers have also found that agarwood possess potent antioxidant, anti-inflammatory and anticancer properties against colorectal carcinoma cells (HCT 116) and pancreatic cancer cells (MIA PaCa-2), which were mediated via apoptotic mechanism (Dahham et al., 2015a; Dahham et al., 2015b).

This method was shown to be simpler but as reliable as the rabbit skin test (Miyamura et al., 1974). Vero cells were cultured and maintained inside T-flask until it reached confluency before being seeded into the biosensor. All drugs were introduced 24 hours following the cell seeding procedures.

In this study, we present a printed circuit board impedance biosensor with copper electrodes based on our previous work (Mansor et al., 2015b) for toxicity studies of natural products on normal healthy cells (Vero cells). For comparison, the Vero cells were also exposed to Taxol (chemo drug – positive control) and DMSO (negative control). The impedance biosensor operates on the basis that healthy cells adhere to the sensor’s electrodes impeding current flow, resulting in high impedance. A high impedance also correlates to a high cell index, indicating that more cells adhere on the electrode surface. Conversely, low impedances reflect that less cells are attached to the electrodes, and this occurs when the cells are dead. Dead and unhealthy cells are usually non-adherent and slowly detach themselves from the electrodes leading to a decrement in cell index.

**MATERIALS AND METHOD**

**Cell cultures**

Vero cells (ATCC® CCL-81™ (ATCC, Manassas, VA, USA), a type of normal cell line, was grown and maintained to confluence in Dulbecco’s Modified Eagle Medium, DMEM (Gibco, Paisley, UK) supplemented with 10% heat inactivated fetal bovine serum (FBS; Gibco). The Vero cells were incubated under standard cell culture environment in CO₂ incubator at 37°C with atmosphere containing 5% carbon dioxide, CO₂. Once confluent, the cells were detached from the culture flask using accutase and were then resuspended in fresh media. The suspended cells were counted and the media of density 1x10⁵ cells/ml in 100 µL was seeded on each biosensor.

**Gelatin preparation and cell adhesion coating**

Gelatin coating is necessary for long term (7-14 days) maintenance of cells. Gelatin coating was performed before seeding feeder cells into the chamber. For preparation of a 1% gelatin, 0.1 gram of gelatin was taken out of the container and put into 100 ml of double distilled water. Next, it was sterilized in an autoclave sterilizer at 121°C for about 15 minutes. In a biosafety cabinet, 0.1% gelatin was added to each chamber to coat it. After gelatin was poured into each chamber, it was incubated for a minimum of 30 minutes at a temperature of 37°C and the excess was discarded. Each of the chambers have to be dried for at least two hours before cell seeding. Ready chambers were sealed with parafilm for storing purposes.

**Agarwood and toxicity test**

In this study, a total of three treatments were sampled at 10 µl. Natural product, i.e. the agarwood branch ethanolic crude extract, was seeded at a final working concentration of IC₅₀ 6 µg/ml; Taxol, a commercial cancer drug, was seeded at a final working concentration of IC₅₀ 2.3 µg/ml; and DMSO, a negative control was seeded at a concentration of 10% (v/v). All IC₅₀ values were based on the response of MCF-7 breast cancer cells from different studies. All treatments were introduced 24 hours subsequent to cell seeding. The response of the cells to the drugs was plotted as cell index number in the next section.

**Sensor fabrication**

Copper electrodes were designed and fabricated on FR4 board using standard PCB fabrication processes i.e. UV exposure, development, etching and finishing, as shown in Fig 2. In this study, two different types of electrodes were modelled—interdigitated electrode and circular electrode—as shown in Fig. 3 with dimensions as given in Table 1. Each board contains eight sets of identical and parallel sensor designs that were used for the experiments. Sensors were kept in a dry place to avoid oxidation, and were cleaned with ethanol and PBS. All sensors were exposed to UV light for a night for sterilization process prior to ECM coating and cell seeding.
Fig. 2 Process flow for fabrication of the impedance biosensor; (a) Positive PCB board was exposed to UV light for 180 seconds to polymerize the photoresist; (b) Unpolymerized photoresist was developed in sodium hydroxide for a few minutes; (c) Unwanted copper was etched in ferric chloride solution, leaving only the traces of the design; (d) Photoresist layer was removed via stripping process; (e) Culture well made of PDMS was attached on the sensor for cell culture purposes; (f) Gelatin was coated on the electrodes to improve cellular adhesion on the electrodes.

Table 1: Size and dimension parameters of the electrode.

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<thead>
<tr>
<th>Electrode Design</th>
<th>Parameter</th>
<th>Dimension (µm)</th>
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<tr>
<td>Interdigitated</td>
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PDMS Fabrication

Sylgard 184 Silicone Elastome Kit was used when making Polydimethylsiloxane (PDMS). The elastomer and curing agent were weighed and mixed at a standard ratio of 10:1 and left overnight to solidify in a closed container. Once hardened, the PDMS was cut to fit the electrode size, punctured to make wells on top of the electrode, glued onto the sensor using another form of liquid PDMS and left overnight to dry and harden.

AD5933 Impedance Converter Analyzer

The impedance network converter integrated circuit AD5933 is the core of the measurement system. Based on its basic configuration, AD5933 can measure impedance from 1kΩ to 10MΩ in the frequency range of 1 to 100kHz (Yusoff, 2016). The Analogue Front End (AFE) can measure impedance at low frequencies through the instrumentation amplifiers, general application op-amp circuits and voltage references using the provided software. There are four steps in measuring the impedance with the evaluation board, which include system setup, system calibration, impedance measurement and data acquisition. The data transmission of the impedance can be done through the F/C port connected to a computer through USB connection.

T-Flask Protocol

As a point of reference for the biosensor testing, the conventional method of cell culture using T-flasks were also performed and microscopic cell counting using hemocytometer was done on the cells inoculated with the natural products. Initially, a total of 13 flasks were prepared and cell-counting was completed every six hours corresponding to the impedance measurements. The media of density 1x10⁵ cells/ml in 5ml was seeded inside each flask. The graphical scheme of the experimental protocol is shown in Fig. 4.
Experimental Setup for Biosensor Test

In the experiment, sensors were retained inside the incubator at all times except during impedance measurement. Every six hours, the sensors were taken out for data acquisition using AD5933 evaluation board, which was connected to a laptop. Cell impedance and phase at specific times were measured using an AC frequency of 180Hz. The impedance and phase were converted into resistance and cell index (CI). Fig. 5 shows the setup of the experiment.

RESULTS AND DISCUSSION

Cell Index (CI) representation of cell

Measurement using AD5933 evaluation board gives the magnitude of impedance and phase of the biosensors. Since CI is a much more accurate representation of cellular behaviour (Boyd et al., 2008) on the surface of electrodes, the impedance and phase were converted into CI number. CI is directly related to cell viability. Higher CI number indicates that more cells are adhering on the electrode surface. The CI for cellular growth can be expressed based on Eq. (1).

\[ CI = \max_{i=1,N} \left( \frac{R_{cell}(f_i)}{R_b(f_i)} - 1 \right) \]  \hspace{1cm} (1)

where \( R_b \) represents the frequency-dependent resistance of control measurement (without cells) and \( R_{cell} \) indicates the frequency-dependent resistance of the cells and electrodes. Measured resistances, \( R \) were extracted from \( Z \) and \( \theta \) using Eq. (2) and Eq. (3):

\[ |Z| = \sqrt{R^2 + X^2} \]  \hspace{1cm} (2)

\[ \theta = \tan^{-1}\left(\frac{X}{R}\right) \]  \hspace{1cm} (3)

Biosensor and T-Flask comparison

Fig. 6 shows the comparison of monitoring NP exposure on Vero cells using two techniques, i.e. conventional method of trypan blue exclusion and impedance biosensor. A similar graph trend can be seen from both methods where the highest number of cell count was shown at time of 60 hours, correlating to the highest measured CI. Fluctuation of CI in several measurements were due to limitation of growth caused by limited surface area. For this case, the T-flask offers larger surface area for adhesion compared to biosensors, where dead cells need to be detached before making space for the next adhesion process. These results indicated that the biosensor can be used to predict the response of cells towards drugs exposure. The IDT electrode design showed higher CI values ranging from 0.08 to 0.5 CI compared to that of circular electrode ranging only from 0.01 to 0.3 CI, suggesting a higher accuracy representation of cellular activities thus concluded as having better sensitivity between both designs.

This result is apparently in agreement with the theoretical concept discussed in a study conducted by Varshney in 2009 (Varshney and Li, 2009). Their article suggested that IDT electrodes show better sensitivities compared to other coplanar electrode designs. The main advantage of IDT electrodes is that they provide uniform and confined electric field distribution between the electrodes, allowing the IDT to detect smaller morphological changes of cells.

![Fig. 5 Right—Sensors and T-flask positioned inside the incubator. Left—Impedance and phase measurements were taken using AD5933 evaluation board and laptop. Four sets of double experiments were conducted on a single board: cell+natural product (blue box), cell+taxol (red box), cell+DMSO (green box) and control cell without drugs (yellow box).](image)

![Fig. 6 Comparison of cell counting using T-Flask method and cell indexing using biosensors](image)
Cytotoxicity evaluation of NP, Taxol and DMSO using impedance biosensor

Cytotoxic evaluation of NP, Taxol, and DMSO were performed on IDT electrode-design sensor as it has better sensitivities compared to the circular design. From the results in Fig. 7, both Taxol and DMSO showed similar trendlines. Taxol did not show major toxicity effects towards Vero cells at the concentration tested, which was our target for positive control. However, the results did indicate a slightly lower CI when exposed to Taxol compared to DMSO, which tells us that there were still less viable cells on the electrodes for Taxol over DMSO.

Meanwhile, NP showed slower growth between 0-60 hours whereby the maximum CI was achieved at 60 hours. This indicates that NP suppresses the growth rate of Vero cells (causing longer lag phase). There is a possibility that NP produces similar effects as Taxol, which targets the microtubule of the cells and prevents them from producing the protein needed for cell attachment. However, this effect is not necrotic, as the cells can still adhere and proliferate on the electrode surface, producing a higher CI number compared to the peak of Taxol and DMSO curves. Earlier findings imply that NP has cytostatic effects on Vero cells. Based on these studies, it can be concluded that the impedance-based biosensor shows great potential to be used as a tool to study the toxicity of natural products. Nevertheless, more work needs to be done to impart more data on cytotoxicity testing of different drugs and on different adherent cells to demonstrate the compatibility of the biosensor to various drug studies.

Cytotoxicity comparison of NP on Vero and MCF-7

A more recent study was conducted in order to investigate the effects of agarwood plant material against MCF-7 breast cancer cells (Abbas et al., 2017). The study concluded that the uninfected agarwood branch ethanolic extract exhibited potent cytotoxic effect against MCF-7 cells. The experiment managed to show cell density reduction as well as morphology change due to the exposure to agarwood branch ethanolic extract (ABEE) of which the IC<sub>50</sub> value was estimated to be at 8 µg/ml and 6 µg/ml for optimized extract. Meanwhile, Taxol, a commercial drug for cancer, was also tested in the experiment as positive control whereby the estimated IC<sub>50</sub> value was 2.3 µg/ml. Fig. 8 shows the population density change when MCF-7 was treated with ABEE. The cytokinetic study showed reduction of cell generation and specific growth rate in ABEE-treated cells as compared to control. Meanwhile, in a separate experiment by using assay techniques, ABEE was also observed to cause reduction in Vero cells (normal green monkey kidney), however the effects were seen only at higher concentrations of ABEE (64 µg/ml). Noticably, contradicting results may be seen from the biosensors and cell counting techniques in reference to different experimental works. The same reduction response of the ABEE on Vero cells were attained in both techniques but occurred at different levels of concentration, suggesting that the experimental design of the biosensor should be refined. This includes optimizing the ratio of medium volume and cell seeding density towards the growth surface area; this is a parameter to be concerned about, such that the outcome between these two techniques can be correlated.

Fig. 7 Cytotoxicity of NP, Taxol and DMSO on vero cells using IDT electrode sensor.

Fig. 8 Images of MCF-7 population density a) control group and b) ABEE group after 48 hours of incubation. Density of cells in ABEE treated group was less and morphology dissimilarity can also be detected when compared with the control group (treated with 10 % dimethylsulfoxide) (Abbas et al., 2017).
CONCLUSION

In this study, PCB board sensors had been fabricated based on two designs, the IDT and circular electrode. IDT was observed to have better sensitivity based on a higher CI number compared to the latter, hence a more accurate representation of cellular behaviour. The impedance-based biosensor has been shown to have great potential as a real-time and cost-effective tool to study the toxicity of natural products.

As with natural products, ABEE exhibits reduction on both MCF-7 and Vero cells indicating that the same set of response can indeed occur but at different concentrations of IC_{50}, ABEE can safely be regarded as less toxic than Taxol and DMSO. However, optimizations on the experimental framework of the biosensor should be done in order to have a correlated study between these two approaches.

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