Non-invasive early diagnosis of oral squamous cell carcinoma using piezoelectric biosensor

Irene E. Rieuwpassa, Sumintarti, Yu-Ri Kim, Megatriani Matandung, Gabriel SA. Matongan

Abstract

Objective: To develop a device capable of distinguishing Oral squamous cell from other types of lesion so early diagnosis can be made non-invasively.

Methods: The piezoelectric buzzer was coated with IgA EBV enzyme conjugate and antigen substance from Bionan. The sensor was designed label free so further substance won’t be needed. The body of the device was made from white acrylic with 4mm thickness. The body was divided into 3 compartments each with different purpose. The main component consists of reagent coated piezoelectric buzzer and sensor. Both sensors are attached to Arduino UNO, breadboard, LIPO UBEC battery and LCD display network. The arduino UNO will read and process the difference between initial frequency and vibration frequency after the antigen were bind by the reagent, and shows the level of IgA in the LCD display.

Results: The device worked and manage to show the level of IgA and presence of EBV in saliva but still need further improvement to make it more specific and valid

Conclusion: Piezoelectric biosensor have shown promise to make early non-invasive diagnosis for OSCC.

Keywords: Oral squamous cell carcinoma, Piezoelectric biosensor


Introduction

Oral Squamous Cell Carcinoma (OSCC) is the manifestation of ulcerative lesion, which could be found in oral mucosa basal tissue. OSCC is the 6th most common found malignancy in the world; approximately 500,000 new cases of OSCC are found every year. OSCC has one of the highest mortality rate in comparison to other type of carcinoma due to difficulty in diagnosing it at early stage. Nearly all of OSCC advances from precancerous lesions such as leukoplakia and erythroplakia. Clinicians find it difficult to differentiate OSCC from precancerous lesion. Due to this difficulty, clinicians often are only able to diagnose OSCC when it has already reached advance lesion. Therefore, mortality rate for this cancer is high because treatment starts after the lesion has already advanced hence, there is an urgent need of marker to differentiate OSCC from precancerous lesion. These markers are vital to help increase the disease prognosis and life expectancy because it enables early diagnosis and treatment.

Most clinical diagnosis requires additional examination such as biopsy for patients who could be diagnosed with OSCC, but this procedure is very invasive. Therefore, these days people seek for less invasive alternative to diagnose a disease, which is by using saliva. Like serum, saliva also contains hormones, antibodies, growth factors, enzymes, microbes and their products. Many of these constituents enter saliva through blood via passive diffusion, active transport or extracellular ultra filtration. Therefore, saliva can be seen in many cases as a reflection of the physiological function of the body. Some advantages of salivary testing for diagnosis are non-invasive, easy to use, inexpensive, safer to administer serum sampling, real-time diagnostic values, no need for trained medical staff, multiple samples can be obtained easily, collection and screening can be done at home, minimal risk of cross-contamination, more economical sampling and storage compared to serum, requires less manipulation during diagnostic procedures compared to serum, and commercial availability of screening assays. In addition saliva has been widely used in clinical research to detect diseases in oral cavity.

Saliva contains many immunoglobulin (Ig) one of them is Immunoglobulin-A. IgA is the second most common Ig in serum and found extensively in secretions such as saliva, mucous, colostrum, and tears. It is, therefore, implicated in humoral immunity. Table 1 since IgA is associated with local immune response and saliva is in direct constant contact with oral lesions, salivary IgA is proposed to accurately reflect the changes caused by precancerous lesion and OSCC in the oral cavity. The increase in salivary IgA in OSCC is proposed to be due to increased local infection, increased antigen inflammatory stimulus, increased local
synthesis, and local host reaction to the disease. People with no lesion, precancerous lesion, and OSCC has different level of salivary. One of the way to measure IgA level is through Enzyme-Link Immunosorbent Assay (ELISA), but ELISA has drawbacks in its requirement of sample manipulation, time, costly equipment, and operation expertise. Therefore, a device that can measure Ig level with ease, handy, and cheap are necessary. Piezoelectric biosensor can be a solution to this problem. A biosensor is an analytical device, which converts a biological response into an electrical signal. The term ‘biosensor’ is often used to cover sensor devices used in order to determine the concentration of substances and other parameters of biological interest even where they do not utilize a biological system directly. Biosensors function by coupling a biological sensing element with a detector system using a transducer. A biosensor can be made as ‘label-free’ meaning it doesn’t need any more reagents for the next usage once the piezoelectric sensor has been coated with reagent. Using biosensor as a mean of diagnosing OSCC by measuring level of salivary IgA can be very handy and easy.

Material and Methods
ELISA reagent (Bionevan China), Phosphate buffered saline (Biogear), distilled water, deionized water, pure ethanol, glutaraldehyde, bovine serum albumin (Erybank), piezoelectric passive AC buzzer sensor, soldering iron, lead, arduino UNO, acrylic board, rainbow cable male-female, rainbow cable male-male, oscillator 10MHz, battery, wind pump, LCD figure 1.

Sensor Coating
Sensor was washed before immobilization procedure starting. Sensor was consequently washed by deionized water and pure ethanol and let to dry in the next step, glutaraldehyde was solved in deionized water up to concentration 5% w/w and 50 µl of the solution was spread over electrode placed into dark and wet box for at least 5 hours and then washed and dried again. Enzyme conjugate, substrate A and B (Bionevan) was spread over the sensor and let incubate in a dark and wet chamber overnight (12 hours). The surface was then washed by a mild stream of PBS, dried and blocked by 5 mg/ml of bovine serum albumin (Erybank) for five hours in dark and wet chamber. The finished immunosensor was washed by a mild stream of by PBS with 0.1% (w/w) Tween 20 prior to use. The electrodes were stored in dry state in a paper box at 4°C until use in the experiments.

Device body
The device body was made using white colored 4mm thick acrylic. The body is comprised of 3 compartments; the first compartment was designed to store the main components, the second compartment to store the battery and arduino network, and the third to store the mini vacuum. The acrylic was cut into smaller pieces and glued together. Small holes was made for cables, switch, saliva holes and wind holes.

Main Components Assembling
The main component consists of IgA EBV reagent coated piezoelectric buzzer also piezoelectric sensor. Before saliva was dripped above the buzzer, sensor will read the buzzer’s initial vibration frequency, after the saliva has been dripped above the sensor, reagent will bind the specific antigen and increase the buzzer’s weight thus slowing it vibration’s frequency. The shift in buzzer’s vibration will be processed by arduino.

Components network assembling
The network consist of arduino UNO, breadboard, LIP UBEC 5V battery, and LCD display. The network were combined with the main components to process the data collected from the piezoelectric sensor. The results were processed by the arduino UNO system and then the results will be shown on the LCD Display figure 2.

Mini Vacuum and hole drilling
This component is used to suck the saliva after use. The holes were made using drills; holes made were used for cables, saliva, and wind holes.

<table>
<thead>
<tr>
<th>Subject</th>
<th>Salivary IgA level (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy/normal</td>
<td>75.51</td>
</tr>
<tr>
<td>Precancerous lesion</td>
<td>89.53</td>
</tr>
<tr>
<td>OSCC</td>
<td>94.47</td>
</tr>
</tbody>
</table>

Table 1  Salivary IgA level in different subjects
Results

Figure 3 the immunoglobulin biosensor device was successfully assembled. The prepared immunosensor were tested for IgA level on normal and aphous stomatitis patients. After 2 ml of saliva was collected from each category, the saliva was then poured on the hole which directly falls on the sensor. The buzzer then vibrates the sensor and the difference in frequency was counted and processed on the arduino UNO. The data collected will be shown on the monitor and to ease the use, the monitor will only present negative or positive according to the range of IgA level counted by the arduino UNO. After each sample IgA level is identified, the sensor is washed with buffered solution to prevent bias in the next IgA level count. The device worked well seeing that it didn't show positive results in patients with stomatitis, which is another type of white lesion.

Discussion

This device was made using piezoelectric biosensor principle. Piezoelectric is a physical phenomenon that leads to the ability of certain materials to produce electrical current when given mechanical pressure. Physicochemical transducer alters physical or chemical changes by antigen and antibody reaction into electrical current. Because this device uses specific antigen, hence this sensor is sensitive only to one certain antibody, in this case Immunoglobulin A in human saliva. This device work by the vibrating the sensor that has been coated before and measure the difference in frequencies between the vibration before and after antibody binds with antigen. Antigen and antibody (IgA) binds when the saliva was poured on top of the sensor. Before saliva was poured on top of the sensor, the sensor was first vibrated using oscillator and the frequency was measured, after the saliva was poured and IgA was bound the frequency changed due to the weighed added on top of the coat, the difference in frequency is then calculated by Arduino and the results will appear on the screen whether the subject are tested positive or negative.

In order to simply test the sensitivity of the device, to investigate whether the device will test positive on any subject, the device was tested on both normal and patient with oral lesion that could possibly have an increase in IgA level namely aphous stomatitis. Some studies have shown that patient with stomatitis has higher IgA level than normal patient without the lesion because IgA plays an important role in protection against infection. Henceforth the device was tested on stomatitis subject. The device shows negative results for both stomatitis subjects because the number of salivary IgA in stomatitis patient is still lower. Further studies need to be conducted to validate the data, calibration, and to make the sensor more specific.

Conclusion

Piezoelectric biosensor have shown promise to make early non-invasive diagnosis for OSCC.
Acknowledgment

The authors would like to say our deepest gratitude to Hasanuddin University Faculty of Dentistry and Electrical Engineering for all the support provided during this research.

Conflict of Interest

The authors report no conflict of interest.

References


This work is licensed under a Creative Commons Attribution