Laccase immobilized on screen printed carbon electrode as quercetin biosensor

Abstract
Rapid electrochemical quercetin detection by oxidation of oxygen from laccase-quercetin enzymatic reaction from *Trametes versicolor* is proposed. Three different methods for laccase enzyme immobilization on screen printed carbon electrode (SPCE) were compared for the development of quercetin biosensors. In this work, glutraldehyde and nafion were deposited onto enzyme electrode surface and were evaluated by cyclic voltammetric and chronoamperometric. The measurements were performed in 0.05M acetate buffer (pH 5.0). The modified biosensor for quercetin detection based on laccase/glutaraldehyde/SPCE gives highest sensitivity (341.2µAmM⁻¹cm⁻²) at concentration of laccase (100mgnL⁻¹) and at applied potential +0.2V. The range of detection is from 1.0 to 20.0 µM (R²=0.99) with a detection limit of 3.74µM and response time of 2.4s.

Keywords: biosensor, quercetin, SPCE, laccase, cyclic voltammetric, chronoamperometric

Introduction
Biosensors can be defined as an analytical device or system consisting of an immobilized biological material in intimate contact with a suitable transducer. Biosensors are smart alternative methods due to their special features such as selectivity, low expenditure, miniaturization, ease to use, time saving and simplicity of operation. The biological part namely called the sensing element that responds to the compound being measured is biological in nature. A specific biological reaction is converted by a transducer to produce a current response which is relative to the concentration of a substrate can be processed into a useable signal. In recent years, there are growing interests attaining the biologically active compounds from natural sources that lead to the development of biosensor. The protective effects of diets rich in tropical against diseases and certain cancers have been attributed partly to the antioxidants such as flavonoid because its shows variety of essential bioactivities. Studies on organic compounds that undergo bioactivity properties such as antioxidant, antiviral, and so on for medical benefits have become trend nowadays. This is due to the abundant species of plants which contains various polyphenols, terpenes and organic acids. Quercetin is a chemical compound from flavonol, one of the seven groups in the flavonoids family, a powerful antioxidant, and is regularly consumed by humans in edible fruits and vegetables at levels of up to 16mg per day as discussed elsewhere. Refer to Van Acker et al apart from flavonoids’ therapeutic effects, quercetin is also an excellent scavenger of free radicals which benefits humans immensely, therefore much attention has been paid to both biological and physicochemical properties of quercetin over the past few decades. The discovery on screen printed electrode (SPE) opens new paths in the field of sensors. SPE is an entrenched and simple technology for mass production of single electrode for biosensor application which applies electrochemical method providing convenience, simple and inexpensive approach. Many established research laboratories fabricate SPE for their own use and some companies produce mass SPE production that offer numerous designs with better quality. In this case, normally SPE manufacture provides the market with autentic characteristics such as different inks, substrates and design of SPE that have specific electrochemical behavior stated by Morrin. Studies that had been carried out by Portaccio et al for the detection of Bisphenol A, reported that thionine was used as electrochemical mediator coupled with a nanostructured carbon black. By means of cyclic voltammetry, the interaction of thionine adsorbed on modified screen printed electrode with laccase/BPA reaction products has been studied. Besides that, in the review by Renedo et al discussed on SPE that had been widely used in many fields such as foods, pharmaceutical and environmental. Enzyme modified SPE and immunosensor are also extensively studied for analysis of metals, pesticides, phenolics, hormones, genetics, drugs, cholesterol and glucose. Therefore, the increasing need for specific sensors to enable fast routine measurements in many fields of analysis elevates SPE usage in biosensors application among researchers in various fields, such as food, medicines, agriculture and pollution monitoring.

Enzyme electrode is one of the most intensively investigated biosensors because enzymes are highly selective and respond quickly to a specific substrate. Most widely used biosensor in SPE analysis of phenolic substances is based on immobilization enzymes such as peroxidase and tyroxidase. However, in this work using laccase as bio-recognition for detection of quercetin is being studied. The phenomenon of direct electron transfer (DET) in enzymes was first described for laccase by Fiera et al. Electron transfer between analyte, an enzyme and electrode surface is fundamental process of electroanalytical methods. Laccase are copper-containing enzymes that has been received much attention from researchers in last decades due to its ability to catalyze the oxidation of a wide variety of organic and inorganic substrates including mono-, di-, and polyphenols, amino phenols, methoxy phenols, aromatic amines and ascorbate with the concomitant four electron reduction of oxygen to water, which make it very useful for application to several biotechnological processes. Studies by Odaci et al for detection of phenolic acid using laccase from *Trametes versicolor* was carried out on commercial oxygen electrode and ferroene-modified screen-printed graphite electrodes. Proposed biosensors were analytically evaluated with real samples of human plasma and it is specific for a class of compounds, not for a single compound and this characteristic has been exploited to obtain an analytical device for a wide spectrum of phenolic compounds.

The measurement of the current resulting from the oxidation or reduction of electroactive species are based on chronoamperometric biosensors was implemented in this work. An electrochemical biosensor may be more attractive as compared to separation methods due simpler on-site analysis and fast response technique. Cyclic voltammetric and amperometric enzymatic electrodes hold a leading position among presently available biosensor systems.
These techniques offer huge information about analyte of interest. The selectivity of the amperometric devices is governed by the redox potential of the electroactive species present. Hence, the immobilization of laccase enzyme would improve the selectivity towards quercetin. The goal of this study is to develop a sensitive and fast carbon screen printed biosensor for quercetin analysis. In this work, SPCE was implemented in the development of the biosensor based on laccase using cyclic voltammetry and chronoamperometry for the detection of quercetin.

**Methodology**

**Chemical**

Quercetin dehydrate ($C_{15}H_{10}O_5$), calcium carbonate (CaCO₃), sodium sulphate (Na₂SO₄), acetic acid (C₂H₄O₂), isopropanol (C₃H₈O), gluteraldehyde 25% (CH₂(CH₂CHO)₂), Nafion (C₂HF₄O₃S.C₂F₄) and laccase (E.C. 1.10.3.2, 23.3Umg⁻¹) were purchased from Sigma-Aldrich. All chemicals used are analytical grade and were used as received without any further purification. All solutions were prepared with ultra-pure water of resistivity not less than 18.2 MΩcm⁻¹.

**Apparatus**

The electrochemical measurements were performed using an Autolab potentiostat (PGSTAT) (Netherlands). All measurements were conducted using three electrode configurations consisting of carbon working electrode area of 4 mm in diameter, carbon-graphite counter electrode and Ag/AgCl as reference. This commercially available SPE was utilized for comparison and sourced from Dropsens (Spain). The pH of acetate buffer and phosphate buffer were measured using pH meter (Mettler Toledo).

**Cleaning of SPCE surface**

It has been observed that cleaning of the SPCE is required before conducting any electrochemical performance. 1.0 M Sodium carbonate, Na₂CO₃ and 0.05 M phosphate buffer saline solution, PBS were used as cleaning agent and was crucial in producing reproducible peak current and good surface active electrode. The cleaning process is performed by oxidising the electrode surface at 1.4V for 90 seconds twice using sodium carbonate.

**Development of biosensors**

Three immobilization methods for the development of biosensors have been approached. Physical adsorption and cross-linked methods using gluteraldehyde (GA) and nafion were implemented respectively. Fabrication of biosensor by physical adsorption was carried out by air drying 20mg mL⁻¹ enzyme solution on surface electrode for 1 hour. Gluteraldehyde solution was prepared by dissolving 40µL GA (25%) in ultra-pure water up to 10mL. The nafion solution was prepared by dissolving 20µL nafion in 65µL isopropanol and 15µL water solution. The same concentration of enzyme was applied for both cross-linked methods at 20mgmL⁻¹ of laccase. The electrochemical measurements were done to study their sensitivity by mean of chronoamperometric detection of quercetin dehydrate in 0.05 M acetic buffer pH 5 as background electrolyte at working potential +0.2V. Approximately, 2µL of enzyme solution was dropped onto working electrode to cover the area of working electrode and the buffer were purged with nitrogen gas to remove any dissolved oxygen in buffer solution.

**UV-Vis Spectrophotometry**

UV-Vis spectra of quercetin were recorded in the wavelength range from 190 to 600nm using a UV-Vis spectrophotometer (Perkin-Elmer UV-Vis Lamda-35).

**Results and discussion**

The electrochemical behavior of quercetin by cyclic voltammetry

Some polyphenols have both oxidation and reduction peaks on their voltammograms showing reversibility on redox process. When the cyclic voltamogram for quercetin was investigated, it was realized that the top scan of the cyclic voltammogram represents the oxidation of quercetin and this is achieved by generating a cathodic current. Anodic current occurs by producing a peak at a particular electrode potential. On the reverse scan, reduced form of quercetin is oxidized back to its original form. In this situation, a positive peak potential is produced indicating to the anodic current value. A compound can be classified as powerful oxidation compound or reducing agent if oxidation takes place at lower potential.

The oxidation of quercetin dehydrate studied by cyclic voltammetry demonstrated 3 oxidation peaks as in Figure 1, which occurred at potentials 0.20, 0.35 and +0.65V (anodic peak potential, $E_p$). Quercetin shows a peak at +0.65 V due to the fact that, it contains hydroxyl group attached to the 3 position of B ring shown in Figure 3. These three oxidation peaks are associated with oxidation of the 5 functional OH groups of quercetin. A reduction peak 4 at about +0.18V could also be seen corresponding to reduction of the oxidation products formed in peak 1. This cyclic voltammogram clearly shows the reversible character of quercetin first electron transfer oxidation reaction. Therefore, a huge number of information on the mechanism of polyphenol oxidation can be obtained by investigating their cyclic voltammogram. Based on this information, the typical cyclic voltammogram of quercetin (Figure 1) is investigated to learn more about the electrochemical behaviour of quercetin (Figure 2). Quercetin has hydroxyl groups attached to the ring structures which may be electrochemically oxidised. Electrochemical studies reveal trends in the electron donating abilities of flavonoids. It showed that the catechol moiety is more easily oxidizable than the resorcinol group. Quercetin contains catechol moiety namely, the 3', 4'-dihydroxyl electron-donating group at ring C, the oxidation of the catechol moiety occurs first at a low positive potential and is associated with peak 1 in the cyclic voltammogram. In quercetin, the –OH groups present on the C ring are responsible for the first oxidation peak. The oxidation process involves a two electron – two proton reversible reactions and forms o-quinone as shown in Figure 3.

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Determination of biosensor sensitivity by chronoamperometry for detection of quercetin

Enzyme loading study showed the amount of sufficient enzyme concentrations that resulted in the significant sensitivities of the developed quercetin biosensor. Table 1 shows the sensitivity, regression coefficient (R²), maximum current at saturation (V_max), Michaelis-Menten constant (K_M) and limit of detection (LOD) for enzyme loading study. This study was performed by chronoamperometric at applied potential +0.2V and the enzyme was deposited onto electrode surface by physical adsorption at concentration ranging from 20 to 100 mg mL⁻¹. The result shows that 100 mg mL⁻¹ gives higher sensitivity and V_max at 341.2 µA mM⁻¹ cm⁻² and 2.0 µA respectively. The modifications on electrode surface were also investigated by using nafion and gluteraldehyde. Table 2 shows the enormous difference between both architectures in the same enzyme concentration at 20 mg mL⁻¹ and GA offers better sensitivity than nafion at 26.61 µA mM⁻¹ cm⁻². This is due to effect of nafion as the polymer that has been used as a proton conductor for proton exchange membrane. Nafion also have hydrophilic properties to be selective to cation charge. This resemblance to a semi-infinite diffusion process was consistent with the slow charge transport in nafion. Therefore, the employment of Nafion is actually to enhance the selectivity of phenol not the sensitivity. As a result, this membrane gives different sensitivity and kinetic constant values.

Calibration by UV-Vis Spectrophotometry

In comparison, the detection of quercetin was examined by UV-Vis spectrophotometry. Two characteristic absorption bands of quercetin were present at 243 and 369 nm and the calibration graph was carried out at maximum adsorption wavelength, λ_max shown in Figure 4. Based on analysis, it can be concluded that the detection of quercetin using UV-Vis spectrophotometer takes longer time analysis and consumed lots of solvent compared to biosensor technique.

Table 1 Quercetin based on concentration immobilized on SPCE

| Enzyme loading (mg mL⁻¹) | 20   | 40   | 60   | 80   | 100  |
|--------------------------|------|------|------|------|------|
| Sensitivity (µA mM⁻¹ cm⁻²)| 4.1  | 193.5| 182.9| 190.7| 341.2|
| R²                       | 0.987| 0.949| 0.991| 0.990| 0.990|
| V_max (µA)               | 0.62 | 1.75 | 0.55 | 0.80 | 2.00 |
| K_M (µM)                 | 48.30| 4.20 | 1.40 | 0.80 | 2.40 |
| Limit of detection (mM)  | 3.19 | 3.12 | 3.10 | 3.49 | 3.74 |

Table 2 Quercetin based on laccase (20 mg mL⁻¹) immobilized by gluteraldehyde and nafion on SPCE

| Immobilization          | Physical adsorption | Nafion (1.0%) | Gluteraldehyde (0.1%) |
|-------------------------|---------------------|---------------|-----------------------|
| Sensitivity (µA mM⁻¹ cm⁻²)| 4.13                | 4.94          | 26.61                 |
| R²                      | 0.987               | 0.997         | 0.971                 |
| V_max (µA)              | 0.415               | 0.135         | 0.462                 |
| K_M (µM)                | 0.048               | 0.028         | 0.007                 |
| LOD (mM)                | 3.174               | 3.895         | 3.420                 |

Conclusion

A biosensor based on laccase/gluteraldehyde immobilized on SPCE has good sensitivity and response time for quercetin detection. A range of detection from 1 to 20 µM (R² = 0.99) with a detection limit of 3.74 µM and response time of 2.45s were observed. Thus, the development of laccase immobilized on screen printed carbon electrode was proven to be sensitive and fast for the detection of quercetin.

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Conflicts of interest

The authors declare that there is no conflict of interest.
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