

# High Sensitivity of HSA Capacitive Immunosensor using Poly-*para*-phenylenediamine Modified Gold Electrode คาพาซิทีฟอิมมูโนเซนเซอร์ที่มีความไววิเคราะห์สูงสำหรับตรวจวัดฮิวแมนซีรั่มอัลบูมิน โดยการโมดิฟายด์อิเล็กโทรดทองด้วยโพลีพาราฟีนิลีนไดอะมีน

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#### Abstract

The highly sensitive capacitive immunosensor detects the capacitance change at the electrode-electrolyte interface when the immobilized antibody on the electrode surface binds to the antigen in the sample. One of the important steps for the construction of the capacitive sensor is the immobilization of antibody via an insulating layer on the electrode surface. This layer is generally based on a self-assembled monolayer (SAM) of alkanethiol compounds, however, the self-assembling process requires a relatively long period of time. In this work electropolymerization of *para*-phenylenediamine (*p*-PD) on the gold electrode surface was investigated as an alternative for the immobilization of antibodies. Anti-human serum albumin (anti-HSA) was immobilized for the capacitive detection of HSA with the flow system. Both concentration of the monomer solution and the number of electropolymerization scan were optimized. Under the optimized conditions, a wide linear concentration range,  $1.0 \times 10^{-15}$  to  $1.0 \times 10^{-10}$  M, was obtained with a very low detection limit of  $1.0 \times 10^{-15}$  M. This sensor could be reused up to 31 times with relative standard deviation (RSD) lower than 3.0%. Moreover, an easy and very fast of the poly-*para*-phenylenediamine (*Pp*PD) modified a gold electrode was obtained.

**Keywords:** Affinity biosensor, Capacitive immunosensor, Poly-*para*-phenylenediamine, Human serum albumin, Electropolymerization

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## บทคัดย่อ

คาพาซิทีฟอิมมูโนเซนเซอร์ที่มีความไววิเคราะห์สูงโดยการวัดค่าการเปลี่ยนแปลงค่าความจุไฟฟ้า ที่รอยต่อของอิเล็กโทรดและสารละลายเมื่อแอนติบอดีที่ตรึงบนผิวอิเล็กโทรดจับกับแอนติเจนในตัวอย่าง ขั้น ตอนสำคัญของการสร้างคาพาซิทีฟเซนเซอร์คือการตรึงแอนติบอดีผ่านชั้นที่มีสภาพเป็นฉนวนบนผิวอิเล็กโท รด โดยทั่วไปจะตรึงแอนติบอดีผ่านการโมดิฟายด์อิเล็กโทรดด้วยชั้นเซลฟ-แอสเซมเบิลโมโนเลเยอร์ของสาร ประกอบอัลเคนไธออลซึ่งใช้เวลาค่อนข้างนาน งานวิจัยนี้ได้ศึกษาการตรึงแอนติบอดีบนขั้วอิเล็กโทรดทอง ที่โมดิฟายด์ด้วยชั้นโพลิพาราฟีนิลีนไดแอมีน และตรวจวัดฮิวแมนซีรั่มอัลบูมินด้วยระบบไหลผ่านโดยใช้ อุปกรณ์ตรวจวัดชนิดคาพาซิทีฟ และได้ศึกษาหาความเข้มข้นของโมโนเมอร์และจำนวนการสแกนที่เหมาะสม ในขั้นตอนการเกิดโพลิเมอร์ ภายใต้สภาวะที่เหมาะสมพบว่าเซนเซอร์ที่พัฒนาขึ้นนี้มีช่วงความเป็นเส้นตรง ที่กว้าง คือ 1.0×10<sup>-15</sup> ถึง 1.0×10<sup>-10</sup> โมลาร์ มีขีดจำกัดการตรวจวัดค่อนข้างต่ำที่ 1.0×10<sup>-15</sup> โมลาร์ โดยขั้ว อิเล็กโทรดที่โมดิฟายด์ ด้วยชั้นโพลิพาราฟีนิลีนไดแอมีนสามารถใช้งานซ้ำได้ 31 ครั้ง และมีค่าเบี่ยงเบน มาตรฐานสัมพัทธ์ น้อยกว่า 3.0 เปอร์เซ็นต์ นอกจากนี้การโมดิฟายด์ผิวอิเล็กโทรดด้วยชั้นโพลิพาราฟีนิลีน ไดแอมีนสามารถทำได้ง่ายและรวดเร็ว

## **คำสำคัญ** : แอฟฟินิตีไบโอเซนเซอร์ คาพาซิทีฟอิมมูโนเซนเซอร์ โพลิพาราฟีนิลีนไดแอมีน ฮิวแมนซีรั่ม อัลบูมิน อิเล็กโตรโพลิเมอร์ไรเซชัน

## Introduction

Human serum albumin (HSA) is a negatively charged, non-glycosylated globular protein with a molecular weight of 67 kDa. Reported to be the most abundant protein in the body, HSA accounts for 60% of the total proteins in plasma, and is synthesized exclusively in the liver, primarity in the polysomes of hepatocytes (Lu et al., 2014). Clinically, albumin was used in past decades to maintain vascular volume in patients with cirrhosis due to its regulation of oncotic pressure. Today, combined with other therapeutic approaches, the volume-expanding properties of albumin are still believed to be beneficial for patients with cirrhosis. Beside its applications in therapy, HSA is regarded as a standard biomarker, with its levels in serum, urine and saliva serving as diagnostic and prognostic criteria. The normal concentration of HSA in blood serum is 35-50 g/L. In diseased conditions, low levels of albumin in serum (pypoalbuminemia < 30 g/L) may reveal malnutrition, liver disease, nephrosis, gastrointestinal protein loss, shock, edema and cardiovascular disease. On the other hand, high serum levels of albumin (hyperalbuminemia > 55 g/L) (Cieplak et al., 2015).

The detection of HSA using a flow injection capacitive immunosensor relied on the immobilization of anti-HSA via a self-assembled monolayer (SAM) of HSA on the gold electrode. The advantages of SAM is that it can provide a reproducible, ultrathin and well-ordered layer, suitable for the immobilization of biological elements (Ferretti et al., 2000; Riepl et al., 1999). However, SAM also has some drawbacks. For example, a relatively long time is needed for



the fabrication, 12 hours of thioctic acid and mercaptosuccinic acid, and 24 hours for thiourea (Shervedani and Hatefi-Mehrjardi, 2007). The cleanliness of the electrode surface also affects the quality of the assembled monolayer (Altintas et al., 2012). Alternatively, a polymer film modified electrode is an interesting approach because it is a simple and fast method that can provide a homogeneous and chemically stable layer with strong adherence to the electrode surface (Rezaei et al., 2015).

In affinity biosensor application, both conducting and non-conducting polymers have been used. However, an electropolymerized non-conducting polymer is more appropriate, because its film thickness is self-controlled during electropolymerization, and a very thin and uniform film can be obtained (Zhang et al., 1996). Non-conducting polymer also have high resistivity (Yuqing et al., 2004) and this is suitable for the capacitive transducer where an insulation electrode surface is required. One interesting polymer is poly-*para*-phenylenediamine (*PpPD*) because there are two amino groups in a monomer unit that can be used to immobilize the biological element.

PpPD, a non-conducting polymer, is used in this work. PpPD presents two amine groups in a monomer unit, a very high surface concentration of reactive groups for the immobilization of biomolecules (Tran et al., 2014). In addition, PpPD film exhibit good stability over a wide pH range (Wu et al., 2005). The deposition of PpPD onto a gold electrode surface can be carried out using electropolymerization process where the appropriate potential with the certain number of scan is applied to the gold electrode immersed in the *para*- phenylenediamine (*p*PD) monomer solution. The monomers are oxidized to form radical cations, followed by coupling reaction to form oliogomers that eventually lead to the deposition of the polymer on the electrode surface (Peng et al., 2009). From electopolymerization, it allows the reproducible and precise formation of a polymer deposited on the surface of the electrode whatever their size and geometry (Cosnier, 2003). Degree of coverage can be controlled by either the number of the scan of the electropolymerization process or the concentration of the monomer (Pournaras et al., 2008).

Thus, this work reports the capacitive immunosensor detection of human serum albumin. Anti-human serum albumin was immobilized on the gold electrode surface via an electropolymerization PpPD film using flow injection system with a highly sensitive capacitive transducer.

#### Objective

The aim of this work is to detect HSA with study the effect of the PpPD formation, i.e., number of scan and concentration of polymer were optimized. The effect of the immobilization, i.e., the time to activate the amine group of the polymer surface was also evaluated. The performance of the PpPD modified electrode was tested with HSA.



## Materials and methods

## Materials

Human serum albumin (HSA) and glutaraldehyde were purchased from Sigma-Aldrich (Steinheim, Germany). *para*-phenylenediamine (*p*-PD) was from laboratory UNILAB reagent (Sydney-Melbourne, Australia). Polyclonal rabbit anti-human serum albumin (anti-HSA) was from Dako (Denmark). 1-dodecanethiol was obtained from Aldrich (Milwaukee, USA). All buffers were prepared with deionized water treated with a reverse osmosis-deionizing system (Pentair, Inc., USA). Before using, buffers were filtered through a nylon membrane filter (pore size 0.2 mm, 47 mm diameter) (Vertical<sup>®</sup>, Albet, Spain) and degassed, respectively. Other chemicals were analytical reagent grade and were used as received.

## Methods

## Gold electrode preparation

Gold rod electrodes (99.99% purity) with a diameter of 3.0 mm were cleaned by dipping in the piranha solution (conc.  $H_2SO_4$  : 30%  $H_2O_2$  equal to 3:1 %v/v) for 20 min followed by rinsing with distilled water. Then, they were polished using alumina slurry (5, 1, 0.3 mm), on a smooth polishing cloth until a mirror-like surface was obtained and subsequently washed with distilled water. The gold electrodes were placed in a plasma cleaner (Model PDC-32G, Harrick, New York, USA) to remove organic and inorganic molecules adsorbed on the gold electrodes surface.

## Antibody Immobilization

In the PpPD film modified gold electrode surface, the monomer of *p*-PD was prepared in 10 mM sodium acetate buffer pH 5.18. The monomer concentrations and the number of scans for electropolymerization were optimized simultaneously. Since a 5.0 mM *p*PD in acetate buffer has earlier been used, the monomer concentration closed to 5.0 mM, i.e., 1.0, 5.0, 10 and 15 mM were studied. Electropolymerized was carried out using cyclic voltammetry at 8, 10, 15 and 20 scans using the potential range from 0.0 to 0.8 V vs. Ag/AgCl with a scan rate of 50 mVs<sup>-1</sup>. The P*p*PD coated electrode was cleaned by rinsing with distilled water and treated with 5.0 % (v/v) glutaraldehyde in 10 mM phosphate buffer pH 7.00 at room temperature for 20 min to activate the aldehyde group. Twenty microliter of 0.5 mg ml<sup>-1</sup> anti-HSA was placed on the modified electrode for 24 hour at 4°C and washed with 10 mM Tris-HCl buffer pH 7.00. The electrode was finally immersed in 1.0 M ethanolamine pH 8.50 for 20 min followed by 10 mM 1-dodecanethiol ethanolic solution for 20 min to block any pinholes on the gold electrode surface. The electrochemical behavior of each immobilization step was studied by cyclic voltammetry (CV).



#### Capacitance measurement

Experimental setup for the flow-injection capacitive system is shown in Figure 1. The measurement was performed in a three electrodes system connected to a potentiostat (Model EA161, EDAQ, New South Wales, Australia). The modified gold electrode was used as working electrode together with a Ag/AgCl reference electrode and a Pt wire counter electrode.



Figure 1 Schematic diagram showing an experimental setup of flow-injection immunosensor system.

The transient current obtained from a potential step of 50 mV (1 pulse/min, pulse width 6.4 ms) was sampled and the first 10 points of the current response with sampling rate 40 kHz was used. From the linear least-square fitting of ln i(t) versus t, value of the capacitance,  $C_{total}$ , could be obtained from the slope (-1/R<sub>s</sub>C<sub>total</sub>) and R<sub>s</sub> can be calculated form intercept (ln (u/R<sub>s</sub>)) corresponding to t = 0 (Berggren *et al.*, 2001). The measuring flow cell had a dead volume of approximately 10 mL. The carrier buffer, 10 mM Tris-HCl buffer pH 7.00, was flowed through the measuring cell using peristaltic pump at a flow rate 50 mL min<sup>-1</sup> to provide a baseline capacitance. Then, HSA with a volume of 300 mL was injected into the flow system and the affinity binding at the electrode surface was performed at room temperature. The decrease in the total capacitance upon binding due to increase of the dielectric layer thickness (Figure 2). After that the regeneration solution (300 mL of HCl pH 2.50) was injected into the flow system to break the binding between anti-HSA and HSA. Then, the capacitance returned to its baseline, ready for a new analytical cycle.





Figure 2 Flow injection capacitive immunosensor system.

#### **Results and Discussion**

#### **Electrode** preparation

The degree of insulation of the modified gold electrode after each immobilization step was examined by cyclic voltammetry (Eco Chemie m-autolab B.V., Utrecht, The Netherlands) with 10 mM  $K_3$ Fe(CN)<sub>6</sub> for PpPD modified electrode in 0.10 M KCl between -0.3 to 0.7 V at a scan rate of 0.1 V s<sup>-1</sup> vs a Ag/AgCl reference electrode. Example of the cyclic voltammograms recorded from the PpPD modified electrode with the immobilized anti-HSA is shown in Figure 3. The CVs of the clean gold surface showed large oxidation and reduction peaks (Figures 3(a)). Redox peak decreased when PpPD (Figure 3(b)) was coated onto the gold electrode surface. The insulating property of the electrode surface was further increased when activated with glutaraldehyde and the anti-HSA was immobilized onto the modified electrode. Then reacted with ethanolamine pH 8.50. This step was to occupy all the remaining aldehyde groups that had not been coupled to the immobilized anti-HSA. After the treatment with the blocking thiol, the electrode surface was completely insulated as shown by the disappearance of the redox peaks (Figure 3(e)). This insulation property is a necessary condition for the non-faradaic detection of the employed capacitive system.





Figure 3 Cyclic voltammograms obtained from (a) bare gold, (b) PpPD, (c) glutaraldehyde, (d) anti-HSA and (e) blocking thiol, 1-dodecanethiol.

#### Activation time of glutaraldehyde

The amine groups of P*p*PD film were activated by glutaraldehyde to form the aldehyde groups which further formed the amide bond with the amine groups of antibodies. This activation time was evaluated to ensure that all amine groups of P*p*PD were activated. The activation time between 5 and 20 minutes of 5% (v/v) glutaraldehyde was investigated. The electropolymerization was performed using 5 mM of *p*PD and 10 scans. HSA  $(1 \times 10^{-15} \text{ to } 1 \times 10^{-10} \text{ M})$  was analyzed and the sensitivity (nF cm<sup>-2</sup> log M<sup>-1</sup>) was calculated and plotted against the activation time (min). From the result, the activation time of 5 min showed the lowest sensitivity because only a small amount of amine groups were activated resulting in a small amount of immobilized antibodies. Ten minutes of activation time provided the highest sensitivity while at 15 and 20 min the sensitivity was slightly decreased. This is because high glutaraldehyde loading may result in the immobilization of antibodies through multiple binding sites, thus reducing the flexibility (rigidity and binding) and the binding capacity of the antibodies as well as the accessibility of the target analyte to them. Therefore, the activation time of 10 min was chosen as an optimum value and used for the ongoing experiments.

#### Number of scan and concentration of para-phenylenediamine

The number of scan used for the electropolymerization of *p*PD was evaluated using cyclic voltammetry at 8, 10, 15 and 20 scans. The scanning was carried out between 0 and +1.5 V at a scan rate of 50 mVs<sup>-1</sup>. Number of scan and concentration of *p*PD were then optimized together using one concentration of HSA ( $1 \times 10^{-12}$  M). The concentrations of *p*PD were 1, 5 10 and 20 mM. Each concentration was studied with number of scan of 8, 10, 15 and 20.



The amine groups of PpPD were then activated with 5% (v/v) glutaraldehyde for 10 min. The electrodes with different number of scan were used to analyze HSA in the flow injection capacitive immunosensor. The capacitance change (nF cm<sup>-2</sup>) against concentration of monomer is shown in Figure 4. The highest capacitance change was obtained at 10 scans with 5 mM. For the lower scan (8 scan), the lower capacitance change might be the result of the lower number of amine groups resulting in the lesser amount of immobilized antibodies. Since the capacitive measurement is based on the capacitance change at the electrode/solution interface, a thin and insulated layer is required. The thicker insulating layer may provide a lower capacitance change. When the number of scan was higher (15 and 20 scans) the reduction of the polymer thickness also increased. Moreover, the higher number of amine groups may result in the higher loading of glutaraldehyde. Hence, the immobilization of antibody may occur through multiple binding site, reducing the flexibility and the binding capacity of the antibodies. Therefore, 5 mM of *p*PD with the number of scan of 10 were then selected as the optimum conditions for PpPD electropolymerization.



Figure 4 The capacitance change obtained from different concentrations of *p*PD and different number of scan.

#### Linear range and limit of detection

Under the optimum conditions of PpPD preparation, linear range and limit of detection were determined for HSA detection. The flow injection conditions were; flow rate 50 mL min<sup>-1</sup>, sample volume 300 mL and carrier buffer 10 mM Tris-HCl pH 7.00. The dissociation of HSA can be done using HCl pH 2.50. The linear range of capacitive immunosensor for HSA



was tested between  $1 \times 10^{-16}$  and  $1 \times 10^{-8}$  M. Each concentration was analyzed for 3 replications. The calibration curve between concentration of HSA (M) and capacitance change (nF cm<sup>-2</sup>) showed the linearity between  $1 \times 10^{-15}$  and  $1 \times 10^{-10}$  M. The limit of detection was determined followed IUPAC recommendation 1994, which was found to be  $1 \times 10^{-15}$  M (Figure 5).



Figure 5 The calibration curve of HSA detection plotted between capacitance change and HSA concentration.

#### Reproducibility

The reproducibility of capacitive immunosensor was tested by analyzing the same concentration of HSA ( $1 \times 10^{-12}$  M). After each injection, the regeneration step was included in the analysis cycle. The residual activity (%) of immobilized antibodies to its antigen after regeneration was calculated.

The residual activity (%) was plotted against the number of injection. For the first 31 injections of HSA, the average residual activity was 96%  $\pm$  2% (RSD = 3%). After 32 regeneration cycles, the average residual activity reduced to 88%. The electrode surface was then tested by cyclic voltammetry. A flat voltammogram similar to one obtained after electrode preparation was observed. This confirmed that the film on the electrode surface was not destroyed by the frequent regeneration. The results indicated that the decrease of residual activity after being used several times.

#### Detection of human serum albumin in real samples

Six serum samples were tested by flow capacitive label-free affinity immunosensor. These sample were diluted 100 times before injecting them directly into the system. The obtained signals were used to calculate the concentration of human serum albumin from the calibration curve. The same samples were tested by the immunoturbidimetric assay (Krystal et al., 1985). The result is shown in Figure 6, that mean the concentration of human serum albumin from the form all samples were corresponding between two methods.





Figure 6 Comparison of the concentration of human serum albumin in serum samples obtained from capacitive immunosensor and immunoturbidimetric assay.

#### Conclusions

This work studied the modification of PpPD on the electrode surface to be used in the capacitive immunosensor. The electropolymerization of PpPD film on the gold electrode was optimized. Anti-HSA and HSA binding was used for the optimization process and the performance of the modified electrode was tested using anti-HSA and HSA interaction. The optimum values of PpPD preparation were 10 min of activation time by glutaraldehyde, 10 scans of the electropolymerization with 5 mM of *p*PD. Under these optimum conditions, the modified electrode can be reused up to 31 times. The electropolymerization of the polymer film onto gold surface took only 15 min. This is the advantage over the SAM which needs 12-24 hours. The proposed PpPD modified electrode would be useful for the electrode preparation when the thin film and high population of amine groups are needed. Therefore, PpPD film is an interesting alternatively material for electrode surface modification in an affinity capacitive immunosensor.

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