Abstract

Tuberculosis (TB), an infectious disease affects millions of humans worldwide and is caused by the Mycobacterium tuberculosis. A label free electrochemical DNA biosensor for the detection of M. tuberculosis using gold electrode modified by self-assembled monolayer of thiol is presented in this paper. Single-stranded DNA probe was immobilized on the surface of self-assembled monolayer gold electrode with the assistance of cysteamine and glutaraldehyde, which was further used to hybridize with the target sequence and non-complementary target sequence. Differential Pulse Voltammetry (DPV) was used to characterize the self-assembled monolayer on the gold electrode and also to study the immobilization of ssDNA probe and hybridization with the complementary sequence (target ssDNA). The hybridization reaction on the gold electrode surface was detected by monitoring a guanine oxidation signal at potential +0.21 V. Electrochemical DNA biosensor using gold electrode modified of thiol (Au-SAM) can be used to determine hybridization between ssDNA probe and ssDNA target sequence of M. tuberculosis with sensitivity value is 0.5152; detection limit is 3.47 µg.mL⁻¹ and quantification limit is 11.56 µg.mL⁻¹

Keywords: DNA Biosensor, gold electrode, Self-Assembled Monolayer, Mycobacterium tuberculosis, guanine oxidation

1. Introduction

Mycobacterium tuberculosis is an airborne contagious disease that is transmitted by coughing, sneezing, or even talking. Once a person becomes infected, any condition that weakens the immune system can trigger the development of active M. tuberculosis. According to the WHO, TB can be defined as a disease of poverty which affecting young adults in their productive years. Furthermore, majority of the TB death occurred in the developing world. This infectious disease is among the three greatest causes of death among women mostly between 15 to 44 years old. Annually, TB infection led to the death of approximately 3 million people worldwide. At least 8 million new TB infected patients were reported worldwide. Theoretically, one TB patient can transfer infection to at least 10-20 people from his surroundings. Therefore, early and quick diagnosis will be of great help to isolate the patients

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and control the disease. Many diagnosis for the detection of TB infection has been carried out by conventional methods such as acid fast staining (Ziehl-Neelsen), culturing on Lowenstein-Jensen media, and also the molecular assays such as Real Time PCR (RT-PCR) and Enzyme Linked Immunospot (ELISpot). The conventional methods are time consuming and insensitive, (mycobacterium culture usually require 4 to 8 weeks to obtain good growth). Some other molecular methods, such as latex agglutination, Enzyme Linked Immunosorbent assay (ELISA), radiometric detection and gen-probe amplified M. tuberculosis direct test (AMTDT), are more sensitive and rapid than conventional methods, however most of these methods are centralized in large stationary laboratories because complex instrumentation and highly qualified technical staff are required.

Electrochemical DNA biosensor has aroused great interest in recent years for its simplicity, higher sensitivity and cheaper equipment. Electrochemical transducers are powerful tools for the detection of DNA hybridization reaction. DNA biosensors are made by immobilizing single stranded (ss) DNA probes on different transducers for measuring the hybridization between the DNA probes and their complementary DNA strands. The immobilization of DNA probe onto the transducer plays an important role in the performance of the DNA Biosensor.

Different methods have been used to immobilize the DNA onto electrode surface. Silva et al. (2010) had been using self assembled monolayer (SAM) by the assistance of cysteamine (cys) and glutaraldehyde (glu) to immobilized DNA probe on surface of gold electrode and used cyclic voltammetry to characterize SAM on the gold electrode. Silva et al. (2010) reported SAM is a good strategy for linking biomolecules, because this technique allows an easy formations of monolayers, reasonable stability for extended period, allowing several reliable measurements. However, SAM also presents limitations, such as, immobilization of enzymes are very much sensitive towards changes in analytical parameters (pH, temperature), a minor change in one of these parameters can sometimes be responsible for losing the biological activity. The hybridization can be exhibited via the redox signal of an electrochemical indicator, which can be an organic molecules, metal complexes, enzymes, redox labels or nanoparticles or using technique direct oxidation.

Issa et al., (2010) had been used electrochemical DNA biosensor using methylene blue on Screen Printed Carbon Electrode (SPCE) for the detection of M. Tuberculosis. Hamdan et al. (2012) was developed electrochemical biosensor for the alternative detection of M. tuberculosis using Pencil Graphite Electrode (PGE) and methylene blue as electroactive intercalator.

Here we report the used of label-free electrochemical DNA biosensor using gold electrode modified by SAM to study oxidation signal of guanine in various concentration of DNA target and to determined the specificity of DNA probe. Differential Pulse Voltammetry (DPV) was used to characterize SAM on the gold electrode and also to study the immobilization of ssDNA probe and hybridization with the complementary sequence (target ssDNA)

2. Materials and Methods

2.1. Materials

2.1.1 Apparatus

Differential pulse voltammetry was carried out using potensiostat Metrohm µAutolab type III connected to PC with NOVA software. Three electrode system were used consisted of an Ag/AgCl as reference electrode, platinum electrode as an auxiliary electrode and gold electrode (0.5 mm) as working electrode.

2.1.2. Chemicals

All DNA oligonucleotides (probe, target and non-complementary sequences) were synthesized by First Base Asia. Probe sequences: 5'-IAC III CAA TCC All IC-3'; Target DNA sequences: 5'- GCC CTG GAT TGC CCG TC -3'; Non-complementary DNA sequences: 5'-GAG CTG TGA AAT TTG GTG CC-3'. Glutaraldehyde (Glu) (25%), 2-aminoethanethiol or cysteamine (Cys) (95%) were purchased from Sigma. Alumina, ethanol (99.5%), hydrogen peroxide (30%), propanol, sulfuric acid were from Merck. The phosphate buffer (0.1 mol L⁻¹; pH 7.0) was used as electrolyte.
2.2. Methods

2.2.1. Pretreatment of the gold electrode

The gold electrode was mechanically polished with alumina slurry followed by rinsing with distilled water and sonication in pure ethanol and water (1:1), for 2 minutes. After mechanical cleaning, the gold electrode suffered a chemical treatment by immersion in a ‘piranha solution’ (H₂SO₄/H₂O₂, 1 : 3 v/v) for 10 minutes at room temperature. Afterwards, the gold electrode was immersed in phosphate buffer (PB) and 10 cycles were carried out between + 0.2 and + 1.5 V at 50 mV s⁻¹. Finally, the electrode was rinsed thoroughly with distilled water for 10 minutes and after exposed to UV radiation for 15 minutes.

2.2.2. Self-assembled monolayer and ssDNA immobilization

The pretreated electrodes were immersed into 25 mmol L⁻¹ ethanolic solution of Cys for 2 hours at room temperature (25°C). After that, the electrode was washed with distilled water and incubated in a Glu solution (2.5% of Glu in 0.1 mol L⁻¹ PB pH 7.0 at 4°C for 50 minutes). The Au-SAM electrode was then exhaustively washed with ultrapure water and incubated with ssDNA probe. The Au-SAM-ssDNA probe was incubated for 1 hour at room temperature (25°C). Then, the electrode was washed for 2 minutes twice with PB (pH 7.0) to remove the non-binding ssDNA probe. The immobilization of ssDNA on Au-SAM was scanned by differential pulse voltammetry in PB buffer solution.

2.2.3. DNA hybridization

The probe-modified electrode was incubated with a solution containing complementary or non-complementary DNA sequence to the ssDNA probe for 30 minutes to form a hybrid double stranded DNA (dsDNA). After that, the modified gold electrode was washed with PB (pH 7.0) to remove the non-hybridized DNA probes, and then scanned by differential pulse voltammetry in PB buffer solution. A scheme of immobilization and hybridization is shown in Figure 1.

3. Results and Discussion

3.1. Pretreatment of the gold electrode

The gold electrode was pre-treated to remove various contaminants on gold electrode surface, as well as activate the gold electrode before modified by self assembly monolayer (SAM), as reported by Silva et al. (2010)
uncontaminated gold surface is important for the good chemisorption of thiol group on gold. The use of self-assembled monolayer allows a connection between thiol group on gold and DNA probe which bind to an aldehyde ligand. Target DNA (cDNA) than will hybridized to probe DNA resulted in the possibility to determined DNA hybridization by guanine oxidation signals (Figure 1).

Modification of electrode can be done in two ways: (i) adsorption of thiols on gold surface followed by the adsorption of specific receptor on thiols monolayer (embedding procedure), or (ii) adsorption of functionalized thiols on gold surface, where thiols play as receptor. In both cases, the performances of modifications are dependent on the gold surface quality. If the gold surface is the smooth-faced one, its cover degree is higher, the obtained monolayer is well ordered, and the electrodes have great analytical performances.

### 3.2. Guanines oxidation signal

In direct label-free electrochemical detection, the hybridization events trigger a changes of an electrical signal. This detection technique was eliminate the need of additional indicator and detection steps. In direct label free, in situ detection can be accomplished by monitoring changes in the intrinsic redox activity of the nucleic acid target (Figure 2). Among the four nucleic acids bases, the Guanine base is most easily oxidized and is most suitable for such indicator-free hybridization detection. To omitted the Guanine signals in probe sequences, Guanines in the probe sequences were substituted by Inosine base which is complement to Cytosine and has low oxidation signal, therefore the oxidation signals only come from Guanine in target DNA sequences which hybridized to probe DNA.

![Figure 2](image.png)

Figure 2. Schematic characterization of label-free electrochemical detection.

Figure 3 shows the DPV voltammogram of blank (Au-SAM), Au-SAM-target DNA, Au-SAM-probe DNA and Au-SAM-ssDNA probe-ssDNA target hybrid. As shown in Figure 3, no peak current of guanine oxidation observed on 0.1 M PB solution (blank), and so do the ssDNA probe due to substitution of Guanine with Inosine. Voltammogram of the target ssDNA and target DNA-probe DNA hybrid showed the signal at +0.21 V and the peak current were 11.7 and 3.65 µA, respectively. Hybrid dsDNA gives the oxidation peak lower than the ssDNA due to the nitrogen bases in dsDNA seems to be hidden into a helical structure of dsDNA and the rigidity of the structure make the nitrogen bases far from the surface of the electrode, therefore the oxidation process was blocked. The signal of ssDNA target was higher due to the flexibility of Guanine base on the surface of electrode.
Fig. 3. The differential pulse voltammograms of blank solution, ssDNA probe, ssDNA target, hybridization of ssDNA probe-ssDNA target on Au-SAM in 0.1 M PB solution pH 7.0. Scanning potential –1.0 to +1.0 V. Scan rate 50 mV.s⁻¹

3.3. Specificity of DNA probe

The specificity of DNA probe for *M. tuberculosis* detection by using Au-SAM biosensor was tested by analyze the oxidation signal of DNA probe- DNA target hybrid and DNA probe- DNA non complementary mixture on Au-SAM. The results shown in Figure 4.

As shown of Figure 4, no peak current observed at +0.21 V for ssDNA-non complementary DNA, because hybridization between ssDNA probe and ssDNA non complementary doesn’t occured. However, DNA probe-DNA target was hybridize through the formation of hydrogen bond between complementary bases (G with C and A with T) and oxidation signal of Guanine in hybrid dsDNA was detected at +0.21 V. Therefore, this electrochemical DNA biosensor has a good specificity for specific oligonucleotide sequences.

3.4. Calibration curve

Calibration curve were obtained by hybridized 10 μg mL⁻¹ ssDNA probe with 0, 5, 10, 15 and 20 μg mL⁻¹ ssDNA target on Au-SAM. Each concentration was measured three times to obtain standar deviation of measurement. A calibration curve is shown in Figure 5. Voltammograms of each concentration of ssDNA target are shown in Figure
6. The peak currents produced by each concentration is varied, and there is a linear relationship between increased of concentration against peak currents height. The higher concentration of ssDNA target will be produced increased in the peak currents.

![Figure 5](image_url)

**Fig. 5.** Calibration curve of peak current vs ssDNA target concentration. Calibration curve was obtain by using ssDNA target 0, 5, 10, 15, 20 µg.mL⁻¹ using DPV on Au-SAM in PB buffer 0.1 M pH 7.0. Scanning potential -1.0 to +1.0 V and scan rate 50 mV.s⁻¹ (n = 3).

As seen in the Figure 5., the signal was linear between 0 and 20 µg mL⁻¹ with a correlation coecient of 0.986 for complementary ssDNA target. The regression equation was \( y = 0.489 x + 0.383 \). We have to calculate confidence interval for intercept (a) to test systematic errors in a measurement and for this measurement the 95% confidence interval for intercept is from -0.90359 to 1.67016, so that is indicate the value of intercept through the point of zero, so the regression equation had a adjustment to \( y = 0.5152 x \). The detection limit and quantification limit values were calculated using following equations: \( y_{LOD} = y_b + 3S_b \) and \( y_{LOQ} = y_b + 10S_b \). The detection limit value is 3.47 µg.mL⁻¹ and quantification limit value is 11.56 µg.mL⁻¹.

![Figure 6](image_url)

**Fig. 6.** The DPV voltammograms of Au-SAM-ssDNA-different concentration of ssDNA target in 0.1 M PB solution pH 7.0. Potential scanning – 1.0 to +1.0. Scan rate 50 mV.s⁻¹.

**Conclusions**

Label-free electrochemical DNA biosensor using gold electrodes modified of thiol (Au/Cys/Glu) can be used to determine hybridization between ssDNA probe and ssDNA target sequence of *M. tuberculosis* with sensitivity (slope value) is 0.5152; detection limit is 3.47 µg.mL⁻¹ and quantification limit is 11.56 µg.mL⁻¹.
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