IMMUNOSENSOR BASED ON INK PRINTED ELECTRODE FOR STAPHYLOCOCCAL ENTEROTOXIN DETECTION

Maria Gardenny Ribeiro PIMENTA-MARTINS
Roselayne Ferro FURTADO;
Rosa Fireman DUTRA;
Luiz Guilherme Dias HENEINE;
Ricardo Souza DIAS;
Maria de Fátima BORGES;
Carlucio Roberto ALVES

1Researcher Professor - State University of Ceara, Fortaleza, Brazil - gardennyrp@yahoo.com.br
2Researcher - Embrapa Tropical Agroindustry, Fortaleza, Brazil - roselayne.furtado@embrapa.br
3Researcher Professor - Federal University of Pernambuco, Recife, Brazil - rfireman@yahoo.com.br
4Researcher - Ezequiel Dias Foundation, Belo Horizonte, Brazil - ricardo.dias@funed.mg.gov.br
5Researcher Professor - State University of Ceara, Fortaleza, Brazil - alvescr@pq.cnpq.br

ABSTRACT: Staphylococcal enterotoxin is one of the more aggressive enterotoxins produced by Staphylococcus aureus strains and it is a common cause of food poisoning. Analytical methods that are sensitive, low cost and easy to use are needed to evaluate the food quality. This work describes the development of a label free immunosensor based on screen-printed AuNPs/carbon and the characterization of its analytical response for staphylococcal enterotoxin B (SEB) detection. The biosensor was constructed from self-assembled monolayer of thiols and protein A for the oriented immobilization of the polyclonal antibodies against SEB. As electrons mediator, potassium ferrocyanide was used. The electrochemical detection was direct with the parameters following: -0.2 to 0.6 V with the pulse amplitude of 0.075 V and the pulse width of 75 ms. The immunosensor showed detection and quantification limits of 0.4 µg mL$^{-1}$ and 1.6 µg mL$^{-1}$, respectively. The immunosensor showed quite satisfactory performance in contaminated and non-contaminated cheese samples.

Keywords: Screen-printed carbon (SPCE). Staphylococcal enterotoxin B. Biosensor. Food safety. Analysis.

RESUMO: Enterotoxina estafilocócica é uma das mais graves enterotoxinas produzidas por Staphylococcus aureus e é uma causa comum de intoxicação alimentar. Métodos analíticos sensíveis, de baixo custo e de fácil uso são necessários para avaliar a qualidade dos alimentos. Este trabalho descreve o desenvolvimento de um imunossensor nao marcado com base eletrodo impresso de carbone/AuNPs e a caracterização de sua resposta analítica para detecção de enterotoxina estafilocócica B (SEB). O biosensor foi construído a partir de monocamada de tiol e proteína A, para a imobilização orientada dos anticorpos policlonais contra a SEB. Como mediador de elétrons foi usado ferrocianeto de potássio. A detecção eletroquímica foi direto e utilizou-se os seguintes parâmetros: -0,2 a 0,6 V com a amplitude de pulso de 0,075 V e a largura de pulso de 75 ms. O imunossensor mostrou limites de detecção e quantificação de 0,4 µg mL$^{-1}$ e 1,6 µg mL$^{-1}$, respectivamente. O imunossensor apresentou desempenho bastante satisfatório em amostras de queijo contaminadas e não contaminadas.


*corresponding author:
E-mail address: alverscr@pq.cnpq.br
Telephone number: +55 85 3101 9766

Recebido em: 25/05/2014 - Aprovado em: 30/06/2014 - Disponibilizado em: 30/07/2014
1. Introduction

Staphylococcal enterotoxins (SEs) are a family of structurally related proteins, produced by Staphylococcus aureus strains (Khreich et al., 2008). SEs are low-molecular weight proteins (26-30 kDa), heat stable, resistant to gut proteases and stable in a wide range of pH (4-10) (LE LOIR; BARON; GAUTIER, 2003; Omoe et al., 2005). These characteristics make these proteins quite resistant to food processing and storage and the proper conditions of the digestive system. Staphylococcal enterotoxin B (SEB) is a cause of serious food poisoning and represents a potential agent of biological terrorism.

Traditional techniques, such as radioimmunoassay, fluorescence-labeled antibody assay, and enzyme-linked immunosorbent assay (ELISA) are widely used for detecting SEB. However, these traditional immunological techniques involve time-consuming procedures, harmful biological markers, and expensive instruments (LANCETTE; BENNETT, 2001; TSAI; LI, 2009; Velusamy et al., 2010). An unquestionable trend for monitoring biological agents and food contaminants is through novel methods that present rapid analytical response, and that are sensitive, portable and low cost (PETRENKO; SOROKULOVA, 2004).

Electrochemical biosensors are an efficient method for detection of analytes in different types of samples including those from food (PIMENTA-MARTINS et al., 2012; LI et al., 2012). In this method, it is possible to associate the specificity of bioreceptors to analytical sensitivity of electrochemical techniques. Electrochemical biosensors based on screen-printed technology have been successfully used in the analysis of contaminated food (DOMINGUEZ-RENEDEO; ALONSO-LOMILLO; ARCOS-MARTINEZ, 2007; YANG et al., 2010).

The technology for preparation of screen-printed electrodes (SPEs) is simple, inexpensive, versatile, and also suitable for mass production of disposable electrodes (DAI et al., 2007; GARCÍA-GONZÁLEZ et al., 2008). SPEs are made by printing electrically conductive inks, especially carbon on an inert support. In this case, they are called screen-printed carbon electrodes (SPCEs). The use of carbon inks is particularly attractive for manufacturing printed electrodes (due to low cost, lead to low background currents and a wide potential window for electrochemical devices) (WANG et al., 1996; MORRIN; KILLARD; SMYTH, 2003). Moreover, theses disposable electrodes can be modified with nanoparticles. The large surface area-to-volume ratio of nanoparticles such as gold, silver, and carbon nanotubes are important to improve the electron transfer of
the biochemical reaction of the biosensor resulting in higher sensibility of the method and to facilitate the assembly of molecular structures on them (PINGARRON; YÁÑEZ-SEDEÑO; GONZÁLEZ-CORTÉ, 2008).

The first step for the construction of biosensors is the modification of the surface to obtain an optimum condition of biomolecule immobilization (RASOOLY; RASOOLY, 1999). Self-assembly has often been applied to the formation of the thin and organized film of biomolecules on a gold surface. In this paper, we used a thiol of short chain that has two functional groups (SH and NH₂), one bound to gold nanoparticles deposited on a SPCE surface and another group bound to Protein A from Staphylococcus aureus. Protein A is a highly stable receptor, capable of binding to the crystallizable fragment (Fc) of immunoglobulins from a large number of species. This step is important for an oriented immobilization of antibodies and it makes a more specific detection of the antigen by immobilized antibodies per portion possible (LEONARD et al., 2003; MUZZUCHELLEI et al., 2010; VELUSAMY et al., 2010).

The possibility to miniaturize the staphylococcal enterotoxin B (SEB) immunosensor using a transducer element that is small in size and disposable such as the Screen Printed Electrodes (SCE) represents an important goal in the development of an amperometric staphylococcal enterotoxin B (SEB) device. We previously showed an immunosensor for staphylococcal enterotoxin detection based on a gold surface (PIMENTA-MARTINS et al., 2012). Recently, we have adapted the methodology for disposable electrodes. In this work, the results inherent to the development of the biosensor with antibodies against staphylococcal enterotoxin B (SEB) on screen printed carbon electrodes are reported. In specific, electrochemical characterization of the construction of biosensors and the analytical response for staphylococcal Enterotoxin B (SEB) were studied.

2. Experimental

2.1. Reagents and apparatus

Electrodag PF-407 C carbon ink was acquired from the Acheson Henkel Corporation (USA). The reagents N-hydroxysuccinimide (NHS), 1-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride (EDC), Cysteamine, Hydrogen tetrachloroaurate (HAuClO₄), and the proteins: Bovine Serum Albumin (BSA) Staphylococcal enterotoxin B (SEB), anti-Staphylococcal B, and Protein A from soluble Staphylococcus aureus were purchased from Sigma-Aldrich, St. Louis, Mo, USA.

Unless indicated, all the antibodies and antigen solutions was prepared in 0.01 mmol L⁻¹ phosphate buffer saline (PBS) at pH
Ultra-pure water (18 MΩ cm) used to prepare all solutions was obtained from a Milli-Q water purification system (Millipore Inc., USA).

The electrochemical experiments were carried out using Autolab PGSTAT 12 potentiostat/galvanostat (Eco Chemie, The Netherlands) with General Purpose Electrochemical System software (GPES 4.9). In the experiments, the SPCEs with AUNPs were used as working electrodes, helical platinum wire was used as a counter electrode, and an Ag/AgCl electrode was used as reference. The electrodes were set up in a glassy electrochemical cell with 10 mL volume.

2.2. Preparation of the screen-printed electrodes (SPCEs)

For the manufacturing of the SPCEs, adhesive plastic mold was fixed on the acetate strip substrate and it was over-coated with one layer of carbon ink (Electrodag® PF 407-C) using the appropriate stencils. The electrode surface (area = 9.62 mm²) was cured for 60 min. at 60 °C. Finally, the adhesive plastic mold was removed and the AuNPs were introduced onto the surface. Thereafter, the carbon ink thickness deposited on acetate was measured using a digital micrometer (Mitutoyo Corporation, Japan) and overall thickness was expressed as an average of ten readings taken randomly on each SPCEs sample.

2.3. Construction of the biosensor

Firstly, the SPCEs were pretreated in 100 mM KCl solution for the activation of the surface according to Alonso-Lomillo et al. (2009). Screen-printed carbon electrodes with AuNPs were immersed in 10 mM cysteamine solution for 2 h at room temperature. Afterwards, the SPCEs were rinsed with ethanol and water, and dried at room temperature (MENDES; CARVALHAL; KUBOTA, 2008). In the next step, 2 mM EDC and 5 mM NHS solution freshly prepared in acetate buffer (pH 5.0), reacted for 30 minutes. In sequence, 5 mg mL⁻¹ of protein A was added the EDC/NHS solution and left to react for 30 minutes. Finally, the modified electrode with cysteamine was incubated in protein A and EDC/NHS solution for one hour. Subsequently, the SPCEs were incubated in anti-SEB solution (100 mg mL⁻¹) for one hour. Then, 10 µL 1% BSA was dropped onto SPCEs for 1 h at room temperature. After each incubation step, the SPCEs were washed with 10 mM PBS and deionized water.

2.5. Immunosensor response

After the fabrication of immunosensor, the modified SPCEs were dropped in 10 µL
of SEB (1 mg L⁻¹, pH 7.4) for one hour. All differential pulse voltammetry (DPV) measurements were carried out in 50 mM PBS (pH 7.4) containing 4 mM K₃Fe(CN)₆ redox probe system at room temperature. These measurements were obtained from -0.2 to 0.6 V with the pulse amplitude of 0.075 V and the pulse width of 75 ms. The cyclic voltammetry (CV) measurements were carried out in 50 mM KCl and 4 mM K₃Fe(CN)₆ solution and scan speed of 0.050 mV s⁻¹.

3. Results and discussion

3.1. The principle of the electrochemical immunosensor for enterotoxin

Electrochemical immunosensors are a type of biosensor that is quite popular. Immunosensors are actually a new version of enzyme-linked immuno-sorbent assay (ELISA), with reduced cost, improved response speed, operation convenience, and comparable or even higher sensitivity than other conventional methods of analysis. In the development of electrochemical immunosensors, the amount of antibodies or antigens are immobilized at the electrode. The biological molecules form the surface sensor. The interaction of the analyte-substrate is measured through electrochemical techniques. These are based on the detection of a change in the electrical properties of the surface transducer as a result of the immunocomplex formation which causes a change in the transducer signal.

A redox probe is used in many cases, in order to detect this change (Figure 1). In the label-free immunosensors, the use of enzymes or another marker is dispensable. This fact confers the advantages of this single stage analysis, which leads to a lower cost and is easier to use than labeled-immunosensors. Direct measurements have been a success achieved in immunosensors adopting the cyclic voltammetry technique in the presence of [Fe(CN)₆]³⁻/⁴⁻ as a redox probe for the detection of microorganisms (CHO et al., 2008). In this work, the immunosensor was developed using the differential pulse voltammetry (DPV) technique and [Fe(CN)₆]³⁻/⁴⁻ as a redox probe. Generally, the detection limit obtained from the differential pulse voltammetry is two or three orders of magnitude lower than values obtained by cyclic voltammetry, reaching the range of 10⁻⁷ to 10⁻⁸ mol L⁻¹ (WELCH; COMPTON, 2006).

The K₃Fe(CN)₆ probe is a valuable tool for testing the kinetic barrier of the charge transfer between solution and electrode interface. When antibodies against Staphylococcal enterotoxin are deposited on SPCEs, the electrode active area decreases and hinders probe ions through their pathways into the electrode. Hence, the redox reaction of K₃Fe(CN)₆ decreases and consequently the peak current too. The same electric behavior
is observed when a blocking agent and the target molecule bind to a sensing surface. The approach to detection of the target analyte is illustrated in Figure 1.

**Figure 1.** Schematic illustration for sensing of staphylococcal enterotoxin in 4 mM K$_3$Fe(CN)$_6$ and 25 mM PBS solution pH 7.

3.2. Electrochemical characterization

The average thickness of the deposited carbon ink on acetate film was 54.5 µm (CV = 1.6%, n = 3). Analysis of the coverage of the ink printed electrode can be conducted by cyclic voltammetric investigations (FURTADO *et al.*, 2012). The area of the redox peak of the redox probe [Fe(CN)$_6^{3-}/4-$] can be used in the characterization of layers with respect to their degree of coverage. Figure 2 shows that after the formation of each layer, a decreasing of the cathodic electric current and the anodic peak of the redox probe [Fe(CN)$_6^{3-}/4-$] occurs. Parallel to this, there was a noticeable separation of oxidation and the reduction peaks of the overlapping voltammograms. In this case, the separation of the redox peak is higher if there is an insulating of the electron flux. All these events are associated with the binding of the biomolecules to the surface electrode.

**Figure 2.** Cyclic voltammograms in 4 mM K$_3$[Fe(CN)$_6$] and PBS buffer solution pH 7.4: steps immunosensor manufacture: (____) bare surface - SPCE; (- - - -) cysteamine (10 mM); (____) protein A (0.05 mg mL$^{-1}$); (……) anti-SEB (100 mg mL$^{-1}$) immobilization and BSA blocking (1%). The measurements were carried out in 50 mM KCl and 4 mM K$_3$Fe(CN)$_6$ solution and scan speed of 0.050 mV s$^{-1}$.

3.2.1. Calibration curve

Calibration curve was obtained by using the differential pulse voltammetry (DPV) measurements. The analytical response of the AuNPS/SPCEs incubated in different concentrations of SEB prepared in PBS pH 7.4 was generated by the increasing the insulating of the electron flux of the redox probe until the surface. The calibration
equation obtained by DPV presented a good linearity with the correlation coefficient of 0.98 ($P < 0.001$, $n = 5$) according to Figure 3.

The limit of detection of 0.4 µg mL$^{-1}$ was estimated considering three times the standard deviation of the measurement of the blank divided by the slope of the calibration curve. The quantification limit of 1.6 µg mL$^{-1}$ was estimated considering ten times the standard deviation of the measurement of the blank divided by the slope of the calibration curve.

Figure 3. Calibration curve of immunosensor for SEB detection in 50 mM PBS (pH 7.4) and 4 mM K$_3$Fe(CN)$_6$. Measurements were obtained from -0.2 to 0.6 V with the pulse amplitude of 0.075 V and the pulse width of 75 ms. Insert: Linear curve of the calibration plot.

3.2.2. Detection of SEB in cheese samples

Attempting to make use of this SEB immunosensor in a complex matrix as Coalho cheese type, commercial samples were acquired for evaluating the performance of the immunosensor. Firstly, samples were prepared according to manufacturer’s instructions from the SET-RPLA test (Oxoid®), which uses the rapid reversed passive latex agglutination method. The positive and negative samples thus classified by the kit, were analyzed by the immunosensor. According to Figure 4, the immunosensor was capable to distinguish the contaminated and non-contaminated cheese samples. These results indicate that the immunosensor can be a low cost alternative method for analytical detection of SEB.

Electrochemical biosensors are often preferred over other transducers due to the ease of the miniaturization and automation system. In this sense, the methodology developed based on AuNPS/SPCE could be adapted to miniaturized electronic systems aiming at a future commercial application for monitoring the quality of food.

Figure 4. Assessment of the immunosensor in contaminated and non-contaminated cheese samples. Measurements were obtained from -0.2 to 0.6 V with the pulse amplitude of 0.075 V and the pulse width of 75 ms.
4. Conclusion

A label free immunosensor for SEB detection was satisfactorily developed based on SPCE modified with AUNPs. The device showed good performance in the contaminated cheese samples as an alternative method of analysis. The results obtained demonstrated that by using this developed system, it is possible to detect low concentration of SEB in few minutes. Moreover, this study leads to additional exciting investigations on rapid detection of a variety of microorganisms and its toxins.

Acknowledgment

The authors would like to thank the Brazilian agencies FUNCAP, CNPq and Embrapa for their financial support.

5. References


WANG, J.; PEDRERO, M.; SAKSLUND, H.; HAMMERICH, O.; PINGARRON, J. Electrochemical activation of screen-