Electrochemical DNA biosensor for detection of pork 
(Sus scrofa) using screen printed carbon-reduced 
graphene oxide electrode

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ABSTRACT: The identification of pork in foodstuff is critical regarding the counterfeiting of meat and kosherness, which is a particular concern for certain religions. In this study, we developed an electrochemical detection method of pork DNA without the use of DNA amplification by using screen printed carbon-reduced graphene oxide (SPC-RGO) electrode. The probe DNA of CytB gene of S. scrofa mtDNA was immobilized on the SPC-RGO surface by passive adsorption. Differential pulse voltammetry (DPV) was used to characterise the probe-target DNA hybridisation based on the target’s guanine oxidation signal. The Plackett-Burman and Box Behnken designs were used to select the factors that influence the hybridisation of probe-target DNA and to optimise each parameter. The following findings regarding the several factors that influence the hybridisation process and optimum condition were obtained: 5.0 µg/ml of probe DNA, 6.0 min of immobilisation time of probe DNA, 20.0 min of probe-target hybridisation time, a scan rate at 0.5 V/s, the pulse amplitude at 50.0 mV, and the washing time of the electrode being as long as 40 s. The limit of detection was obtained at 1.76 µg/ml for the linear range of 0–10.0 µg/ml target DNA while the relative standard deviation (RSD) was 2.25%. The DNA biosensor was tested on the isolated DNA samples from pork, chicken and beef while the voltammetry response reveals that it can distinguish the samples. These results indicate that the proposed electrochemical DNA biosensor has the potential to develop the detection method of pork content in the food samples.

KEYWORDS: DNA biosensor, pork, voltammetry, SPC-RGO electrode

INTRODUCTION

The adulteration or preparation of meat products by mixing meats with cheaper meats of different species sources has been commonly practiced in many countries. Today, consumers demand high-quality food products with the appropriate labeling of ingredients for various reasons, including medical motives, personal preferences (e.g., vegetarians) or religious prohibitions such as for Jews and Muslims. Rising consumer demand underscores the need for the development of more swift and reliable methods to identify species in food commodities such as detecting pork in food. On this note, instead of protein, a DNA analysis would be preferable to identify species due to the nature of protein being easily denaturised while processing [1–4].

DNA-based methods have become a consideration for researchers, managers and regulators. This method involves the detection, identification, quantification and monitoring of the falsification of species in raw and processed meat [5]. There are several detection and quantification methods for the identification of pork in food products that rely on DNA-based analyses. The polymerase chain reaction (PCR), real-time PCR, PCR-restricted fragment length polymorphism (PCR-RFLP), real-time-multiplex PCR, and species-specific PCR were used extensively [3, 4, 6]. Most recently, duplex droplet digital PCR has become more frequently used in identifying fraudulent meat products [7, 8].

There are numerous advantages to DNA-based analysis, including its rapidity, sensibility, simplicity and capacity for widespread speculation on the future availability of inexpensive and accurate means for identifying and quantifying each declared or
undeclared component in finished commercial products [5, 9]. Recent developed DNA-based methods include DNA sensors, DNA biochips and DNA microarray technology. These methods constitute a modern approach that enables the examination of complex mixtures of PCR products and may potentially identify a wide array of species simultaneously [5, 10, 11].

A modified gold nanoparticle’s DNA biosensor with citric acid-tannic was utilised for porcine detection in mixed meat spectroscopically. The visual change was rapid and the species detection was performed within ten minutes without any instrument. However, the method was solely qualitative, and the detection limit of 4–6 µg/ml was considerably higher than conventional and real-time PCR [12, 13]. A chemiluminescent optical fibre genosensor was also developed for the detection of pork meat, which can detect a 1% quantity in mixture samples [14]. The new electrochemical DNA biosensor based on the bioconjugate of gold nanoparticles-DNA biosensor has also been reported, which was selective towards 10% of the pork DNA in the mixture [15].

The DNA-based electrochemical biosensor has gained attraction due to its simplicity, sensitivity, selectivity, and economical equipment. The use of graphene as a transducer in several electrochemical DNA biosensor studies has been successfully developed due to its unique feature. Graphene (or graphene oxide) is an excellent material as an anchor for biomolecular detection because of its large surface area (theoretically 2630 m²/g) and unique sp² (sp²/sp³) bond [16]. Based on the differences of binding affinity of single-stranded DNA (ssDNA) and double-stranded DNA (dsDNA) to the graphene layer, graphene has been successfully adopted as a means of distinguishing DNA strands [17]. Graphene has a larger surface area with better electrical conductivity than a glassy carbon electrode and is suitable for use as a sensing medium [18].

The application of experimental design for the detection of pork by electrochemical DNA biosensors has never been previously reported. Plackett-Burman (PB) design and Box-Behnken (BB) response surface methodology has, on the other hand, been successfully applied in various experimental designs with complex design parameters involving two or more parameters by producing robust design models. Herein, we report a voltammetric DNA biosensor for pork detection based on the guanine oxidation signal of target DNA using SPC-RGO electrodes, and the application of PB and BB design experiments to obtain optimised parameters. The scheme of SPC-RGO DNA biosensor is indicated in Fig. 1.

**MATERIALS AND METHODS**

**Materials**

The DNA probe used in this study was based on Ref. [13]. Twenty nucleotide swine specific probe of CytB S. scrofa mtDNA nucleotide between 567 and 586: 5’-TACCICCTCIAACCCITAC-3’ (guanine base was substituted with inosine). The target DNA complementary sequences: 5’-GTACGGCTGCGAGGGCGGTA-3’. The oligonucleotide sequence was synthesised by IDT (Integrated DNA Technologies Pte. Ltd. Singapore). Commercial graphene oxide (GO) (Graphenea SA ES A7502260) was re-dispersed with redistilled water, NaCl, K₃[Fe(CN)₆] and acetic buffer saline (ABS), while phosphate buffer saline (PBS) was purchased from Merck (Germany). DNeasy Mericon Food (Qiagen, Cat. 3695140) and restriction enzyme Sul1 (R0138S) came from New England Biolabs (USA) while the SPCEs (Cat. DRP 110) were from Dropens (Germany).

**Apparatus**

Cyclic voltammetry (CV) and differential pulse voltammetry (DPV) measurements were conducted using transducer Metrohm® Autolab Type III Potentiostat/Galvanostat with NOVA 1.10 software (Metrohm, Switzerland). A pH meter (Mettler Toledo InLab pH combination polymer electrodes), microcentrifuge (Thermo Scientific MicroCL 17R, USA), BDA digital compact gel documentation system, a multi-mode reader (Tecan Infinite M200 PRO, Switzerland) and an UV Biophotometer Eppendorf (Germany) were also used. Finally, DESIGN-EXPERT software version 9.1 (Stat ease Inc., USA) was used for processing data of PB and BB design.

**Modification of SPCE with GO and electrochemical characterisation**

The SPCE was modified with three different concentrations of GO: SPC modified with 1000 µg/ml of GO (SPC-RGO 1000), SPC-RGO 500 and SPC-RGO 500 with 0.25 M NaCl. Briefly, 40.0 µl of GO (that was already sonicated for 15 min) was dropped onto the SPCE, respectively. The GO was electro-deposited on SPCE and characterised by cyclic voltammetry (CV) by observing the redox activity of the electroactive species.
Fig. 1  The scheme of the SPC-RGO based pig DNA biosensor. The presence of target DNA (black) and the absence of target DNA can be distinguished by a differential pulse voltammetry signal.

\[
\text{[Fe(CN)]}^3^-/[\text{Fe(CN)]}^4^- \text{ using } 10 \text{ mM K}_3[\text{Fe(CN)}]_6 \text{ containing } 100 \text{ mM of KCl. The CV was done for seven cycles at a potential range at } -1.6 \text{ to } +0.4 \text{ V for } 120 \text{ s, at a frequency } 50 \text{ Hz, amplitude } 0.04 \text{ V and voltage step at } 0.004 \text{ V [20].}
\]

**Immobilisation of the probe DNA and hybridisation of probe DNA-target DNA**

The probe DNA (30 µl of 5.0 µg/ml, diluted in ABS pH 5.0) was dropped onto SPC-RGO and incubated for 6 min at room temperature. It was washed with ABS pH 5.0 for 40 s. Afterwards, x µl of y µg/ml of the target DNA (x and y were based on experimental design) (diluted in PBS pH 7.2) was dripped onto SPC-RGO-probe DNA, followed by incubation for 20 min and then washed with PBS pH 7.2 for 40 s. After this process, the target DNA was hybridised to the probe DNA.

**Voltammetric analysis of biosensor DNA**

The probe DNA on SPC-RGO was hybridised with various concentrations of synthetic target DNA (0–10 ppm). The measurement was done at the optimum condition obtained by differential pulse voltammetry analysis at the potential range from +0.5 V to +1.5 V in 0.1 M phosphate buffer solution pH 7.0. The DPV peak current was measured based on the guanine oxidation signal of the target DNA, which was hybridised to the cytosine in the probe DNA sequence. The guanine in the probe DNA sequence was substituted with the inosine, which does not show peak current in the range −1 V to +1.5 V. The limit of detection was calculated by measuring the average of blank responses, plus three times the standard deviation of the blank response.

**Determination of optimum experimental condition**

Determination of optimum experimental conditions was carried out using the factorial RSM Box-Behnken design level −1, 0, and +1 using MINITAB 17 statistical software. Eleven factors (X) were screened by applying PB, including GO concentration (A), probe DNA concentration (B), time to immobilise probe DNA (C), time to hybridise probe-target DNA (D), the scan rate (E), pulse amplitude (F), the number of CV cycles (G), the pH buffer of probe DNA (H), pH buffer of target DNA (I), temperature (J), and salt concentration (K).
DNA (J), the washing time (K), and pretreatment of electrode (L). The selected factors from the PB design were optimised by the Box-Behnken (BB) experiment design. The analytical parameters were then determined using the optimum condition of the BB results. The linearity range was determined by examining various concentrations of target DNA (0–10 µg/ml). Furthermore, the biosensor response was measured using the DPV at the potential range of +0.5 V to +1.5 V in a 0.1 M of phosphate buffer pH 7.

DNA extraction and application of voltammetric DNA biosensor for the detection of meat sample

Approximately 20 mg of mashed pork, beef, and chicken meat samples were weighed and placed into a 1.5-ml microtube. The total DNA was isolated following the procedures in the DNeasy Mericon food kit (Qiagen). The isolated DNA was then analysed by electrophoresis on 1% agarose gel (the data were not shown) and quantified using a UV spectrophotometer. The isolated DNA was cut with the SalI restriction enzyme to linearise the mtDNA following the procedure. The DNA concentration was measured by Biophotometer UV at 260 nm.

The purity of the DNA was then determined to calculate the ratio of absorbance at 260/280. The DNA samples were diluted five times to a total volume of 50 µl. DNA samples were denatured by heating at 95 °C for 5 min and 20 µl of DNA samples were dropped onto the SPC-RGO-DNA probe to be incubated for 1 h, followed by rinsing with 0.05 M phosphate buffer pH 7.0. The biosensor response was measured using DPV at the potential range −0.45 V to +0.1 V.

RESULTS AND DISCUSSION

SPCE modification and cyclic voltammetry characterisation

The SPCE was modified with three different concentrations of GO: SPC modified with 1000 µg/ml of GO (SPC-RGO 1000), SPC-RGO 500, and SPC-RGO 500 with 0.25 M of NaCl. Fig. 2 depicts the characterisation of the SPC-RGO using the ferric cyanide redox system by CV. The SPC-RGO 1000 showed a higher current response compared to other modifications. The success of electrodeposition and reduction of the graphene oxide in the solution were dependent on the average conductivity. The optimum conductivity of GO was about 4–25 mS/cm available from 500 µg/ml of GO:0.25 M NaCl (1:1) [21].

Fig. 2 Cyclic voltammogram of K$_3$[Fe(CN)$_6$] containing 100 mM of KCl on SPCE with and without RGO modification; (1) SPCE without modification, (2) SPCE with 1000 ppm of GO-Na$^+$, (3) SPCE with 500 ppm of GO-Na$^+$, and (4) SPCE with 1000 ppm of GO-Na$^+$. Fig. 2 also shows that the modification of SPCE with GO affects the current response because the GO increases the surface area of the electrode. The peak current generated by SPC-RGO 1000 was 2.3-times higher compared to that without GO modification. The electron transfer from the ferric cyanide redox system became easier on the SPC-RGO surface than SPC without GO electrodes.

Screening of significant factors and optimisation of experimental condition

The probe DNA used in this study was 20 nucleotides within the CytB gene of S. scrofa mtDNA. The CytB gene was used because it has low homology to the sequence of other species while mtDNA is present in high evolutionary values in abundant amounts of copy. The mtDNA genes were also protected from degradation attacks due to their protective mitochondrion forms and sizes [5].

The immobilisation of probe DNA onto SPC-RGO electrodes occurs due to the strong adsorption of the ssDNA strand on GO shown by high fluorescence quenching efficiency of GO [22]. These passive adsorptions would immobilise the biomolecules onto the electrodes by utilising hydrophobic, hydrophilic and other physical interactions.

The screening of factors that influence the experiment using the PB design was obtained via the Randles-Sevic equation for voltammetry analysis. The GO concentrations were chosen between 1000 and 4000 ppm based on previous research [23]. DNA probe concentrations were between 5 and 20 ppm based on the effectiveness of DNA concentration on the surface of the graphite electrode [24].
The immobilisation and hybridisation time was chosen between 5 and 20 min based on previous research for the effectiveness of the analysis period\textsuperscript{[24]}. The lowest and highest values of voltammetry parameters as scan rate, pulse amplitude and cycle number were chosen based on the effectiveness of the deposition of GO onto SPCE. The pH and washing time for experiment optimisation and pre-treatment was done to make SPC-RGO more positively so that it can absorb negative phosphate groups from DNA\textsuperscript{[24]}.

The calculation of regression coefficients is initiated upon a collection of 12 PB design runs and calculated responses. The results were interpreted using the first-degree polynomial model, which can be presented in the following equation:

\[
Y = 0.8317 - 0.3533 A + 0.2367 B + 0.2217 C + 0.4911 D - 0.0500 E - 0.2167 G + 0.0172 H - 0.2033 J + 0.4050 K + 0.5384 L.
\]

This equation based on 
\[ Y = \beta_0 + \cdots + \beta_i X_i, \]
where
\[ Y \] is predicted response (the peak currents),
\[ \beta \] is the intercept of mean,
\[ X_i \] is the setting (A–L factors), and
\[ \beta_i \] are the respective coefficients. An analysis of variance (ANOVA) was performed in order to determine which factors significantly affected the peak current. The ANOVA (\( F \)-test) showed that the second model is well adjusted to the experimental data (the data were not shown).

The coefficient of variation indicates the degree of precision to which the treatments were compared. However, because the number of degrees of freedom for the error term is small in saturated designs, the power of classical ANOVA was too low\textsuperscript{[25]}. For this reason, a graphical tool, the effect probability plot of the estimates, was used to identify possible significant effects and to estimate the standard deviation of the effects. Significant effects in normal plots are detected through visual inspection. A graphical representation of the significant effect probability is shown in Fig. 3 as generated by the software program Design Expert 9.1\textsuperscript{[26]}.

By using the effect probability plot in Fig. 3, we were able to identify 7 important factors of the experiment: the probe DNA concentration (B), immobilisation time (C), hybridisation time (D), scan rate (F), pulse amplitude (H), washing time of electrode (K), and the pretreatment of the electrode (L). These important factors are marked with red squares in the plot (Fig. 3).

Based on the PB design result shown in Fig. 3, 7 variables were chosen for further optimisation by using the BB design, excepting pretreatment of the electrode because it was one of the most important factors. Therefore, all experiments were conducted with the pretreatment of the electrode. The experiment consisted of 48 experimental runs (data were not shown) to optimise the peak current as the responses. Table 1 presents the experimental BB design with independent variable values.

![Graphical representation of the significant effect probability in a normal plot of the estimates of pork DNA biosensor generated by the software program Design Expert 9.1.](image-url)

**Table 1** The optimisation of experimental conditions using Box-Behnken design with the independent variable values.

| Factor              | Unit       | Level |
|---------------------|------------|-------|
| B-probe DNA concentration | µg/ml     | −1    | 0   |+1 |
| C-immobilization time | min       | 5.00  | 12.50 | 20.00 |
| D-hybridisation time | min       | 5.00  | 12.50 | 20.00 |
| F-scan rate         | V/s       | 0.50  | 0.85  | 1.20 |
| H-pulse amplitude   | mV        | 20.00 | 35.00 | 50.00 |
| K-washing time      | s         | 4.00  | 22.00 | 40.00 |
parameter.
By using the ANOVA, the statistical significance of each coefficient of regression equation was checked by Fischer’s value (F-value) and probability value (p-value) which, in turn, indicate the interactions of the variables. The F-value and the p-value obtained were 2.66 and 0.0286, respectively. The large F-value 2.66 indicates the significance of the term. This model was also significant with a p-value of 0.0286, which meant that only 2.86% of the data occurs in noise. Optimisation was then performed to search for the values of different independent variables that were considered optimal, effective and efficient to achieve the desired result [25,27]. The optimisation process often involves a single response; in this research, the expected response was obtained through the maximum current response.

Based on data processing, immense desirability value was also obtained, which was 0.558 and used as the optimum value of the process. The value of desirability lies between 0 and 1, which describes the proximity of the response to the ideal amount. If the response lies at an unacceptable interval, the value of desirability is 0. Moreover, if the response is at a range reaching the ideal value, desirability is 1.0. The response between the tolerance intervals is that of desirability 0 and 1 [29]. The optimisation goal is not to obtain a desirability value of 1.0, but to find the best conditions that bring together all the functions.

The optimum condition of experiments with the highest desirability value were as follows: B = probe DNA concentration (5.00 µg/ml), C = immobilisation time (6.0 min), D = hybridisation time (20.0 min), F = scan rate (0.5 V/s), H = pulse amplitude (50.0 mV), K = washing time (40.0 s). The peak current of 1.72 (µA) was then obtained as the optimum condition of experiments (the data were not shown).

**Voltammetric measurement of the target DNA based on Box-Behnken optimisation**

Fig. 4 shows the peak current linearity of the target DNA with various concentrations under optimum conditions. The linear relationship between target DNA concentration and the peak current of the Guanine oxidation was I (µA) = 0.2068 [target DNA] + 0.0622 while the R² value was 0.9836.

After determining the range of confidence in the intercept for finding out whether there is a systematic error in the measurements, the intercept confidence range was calculated with a 95% confidence level between −0.2044 to 0.3287. The intercept value passed 0 points, following which the regression equation was adjusted to y = 0.2148x. The slope of the equation was then used to calculate the limits of detection (LoD) and limit of quantitation (LoQ). By using the equation LoD = 3S_b/m, where S_b is the standard deviation of the blank, and m is the slope of the equation, the detection limit of the measurement was obtained at a value of 1.76 ng/µl. RSD for five times measurement of 10.0 µg/ml target DNA was 2.25%.

The previous study shows that a gold nanoparticle-probe DNA bioconjugate based on electrochemical biosensor for detection of Sus scrofa mtDNA using methylene blue indicators [15] had a lower detection limit than this study. Nevertheless, this proposed method has the advantage of being simpler. Its simplicity is found in the immobilisation system only by simple adsorption with the detection
of hybridisation based on the target’s internal base electroactive properties.

Guanine is the most electroactive part of the DNA molecule. The substitution of guanine in the probe’s DNA sequences with inosine enables the detection of the guanine oxidation in the target DNA. This label-free electrochemical detection has eliminated the external labels or indicators and significantly shortened the assay time, hence increasing interest [24, 30–32].

The comparison of the analytical performance of the proposed DNA biosensor with previous biosensor research is shown in Table 2. It can be concluded that, based on the detection limit, the proposed method can be used as an alternative to determine DNA in raw meat samples in a simple way.

Application of voltammetric DNA biosensor for the detection of meat sample

The isolated mtDNA from pork, chicken, and beef meat were characterised by electrophoresis and the spectrophotometer (data were not shown). The UV absorption measured the quantity and purity of DNA at a wavelength of 260 nm and 280 nm. The absorption ratio of A260:A280 was 1.85, indicating that the isolated DNA was pure or not contaminated with a protein. The DNA was then cut using a SalI restriction enzyme to linearise the mtDNA. Restricted DNA was used to determine the response and selectivity of the electrochemical DNA biosensor.

Voltammograms of guanine oxidation signal generated from the hybridisation of probe-sample DNA (pig, chicken, and cow DNA) were shown in Fig. 5. The result shows that the peak current signal of the hybridised probe-pork mtDNA sample is four-time higher compared to that of chicken and beef samples. The chicken and beef mtDNA will not hybridise with the probe DNA because it does not contain a complement base of the probe sequence. However, there might be several base pairs of sequence matches. Therefore, the current responses were observed, but lower. The difference in peak current height can then be used to ensure that the sample contains pig DNA.

CONCLUSION

Based on the selected factors and optimisation with Placket-Burman and Box-Behnken experiment design, the voltammetric DNA biosensor using SPC-RGO can be used to detect pig DNA in raw sample. The factors affecting the experiments were probe DNA concentration, the immobilisation time of probe DNA, the hybridisation time of probe-target DNA, the scan rate, pulse amplitude, washing time and pre-treatment of electrodes. The importance of this study will serve as a baseline for developing other alternative methods for monitoring food adulteration, especially for kosher or halal meat.

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