A bisulfite treatment and PCR-free global DNA methylation detection method using electrochemical enzymatic signal engagement

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\textbf{ABSTRACT}

In this paper we report on a bisulfite treatment and PCR amplification-free method for sensitive and selective quantifying of global DNA methylation. Our method utilizes a three-step strategy that involves (i) initial isolation and denaturation of global DNA using the standard isolation protocol and direct adsorption onto a bare gold electrode via gold-DNA affinity interaction, (ii) selective interrogation of methylation sites in adsorbed DNA via methylation-specific \textit{5}m\textit{C} antibody, and (iii) subsequent signal enhancement using an electrochemical-enzymatic redox cycling reaction. In the redox cycling reaction, glucose oxidase (GO\textsubscript{X}) is used as an enzyme label, glucose as a substrate and ruthenium complex as a redox mediator. We initially investigated the enzymatic properties of GO\textsubscript{X} by varying glucose and ruthenium concentration to delineate the redox cyclic mechanism of our assay. Because of the fast electron transfer by ruthenium (Ru) complex and intrinsic signal amplification from GO\textsubscript{X} label, this method could detect as low as 5% methylation level in 50 ng of total DNA input. Moreover, the use of methylation-specific \textit{5}m\textit{C} antibody conjugated GO\textsubscript{X} makes this assay relatively highly selective for DNA methylation analysis. The data obtained from the electrochemical response for different levels of methylation showed excellent interassay reproducibility of RSD (relative standard deviation) < 5% for \( n = 3 \). We believe that this inexpensive, rapid, and sensitive assay will find high relevance as an alternative method for DNA methylation analysis both in research and clinical platforms.

1. Introduction

Development of a reliable methodology for epigenetic biomarker detection is a key demanding factor in accurate diagnosis and prognosis of the cancer progression (Portela and Esteller, 2010). DNA methylation is a well-known epigenetic change that could be occurred via hypermethylation or hypomethylation (Ehrlich, 2002). Increased oncogene expression, activating transcription and altering the genomic stability often refer to hypomethylation, and it is observed in metastatic tissues and primary tumors. To profile DNA methylation, generally bisulfite treatment is used to convert all the DNA without changing the epigenetic mark, which could allow subsequent analysis of DNA bases by well established molecular biology techniques, such as direct sequencing (Paul and Clark, 1996), pyrosequencing (Tost and Gut, 2007), methylation-sensitive melting curve analysis (MS-MCA) (Worm et al., 2001) and methylation-sensitive high-resolution melting (MS-HRM) (Wojdacz and Dobrovic, 2007) etc. Despite high reliability of these methods, their inability to detect \textit{5}-methylcytosine \textit{(5-mc)} oxidation products (Wu and Zhang, 2017), incomplete conversion of cytosine to uracil (Genevreux et al., 2008; Grunau et al., 2001; Rein et al., 1997), primer design (Li, 2007; Thomassin et al., 1999), and biased PCR amplification (Luo and Preuss, 2003; Warnecke et al., 1997; Wojdacz et al., 2014) make the overall detection method more challenging and time-consuming.

To avoid these complications, numerous bisulfite free methods for DNA methylation analysis have been proposed. Efforts have also been directed towards the development of platforms that are rapid, cost effective and involve simple operation to extend their application to routine clinical testing (Kurinomaru and Kurita, 2017; Zhang et al., 2015). Among these techniques, enzyme-linked immunosorbent assay (ELISA) is used to quickly assess global DNA methylation using 100 ng to 2.0 µg of total DNA. However, high variability of this assay is only suitable for the rough estimation of DNA methylation (Kurdyukov and Bullock, 2016). Another common approach of DNA methylation detection is the use of affinity capture agents such as methyl binding domain (MBD) proteins or antibodies raised against \textit{5}-methylcytosine, which have also gained much interests (Haque et al., 2017a; Krejcova et al., 2017). While MBD proteins have high affinity towards...
catalytic response is directly proportional to the amount of the surface-enhanced chronocoulometric (CC) responses via EN redox cycling. The presence of the glucose and ruthenium complex, the electrode generates \([\{\text{Ru(NH}_3\}_6\}^{3+}\) undergo a fast outer-sphere electron-transfer reaction electrochemical enzymatic (EN) redox cycling reaction (Singh et al., 2012; Lai et al., 2019; Singh et al., 2013; Zhang et al., 2017). Another approach to overcome these limitations, mediated electron transfer can be achieved on various materials to enhance the enzymatic catalytic signals (Akanda and Ju, 2017; delle Noci et al., 2008; Haque et al., 2017a; Kaatz et al., 2013; Yu et al., 2016a, 2016b, 2016c). Most of these assays were developed based on either nanomaterials (NMs) or enzyme to improve the sensitivity and selectivity. Due to their large surface area and biocompatibility, numerous studies have been reported that utilizes a wide variety of nanomaterials of different morphologies, such as, nanoshells, nanospheres, nanowires etc that increase the performance for detection of DNA methylation. However, the unstable nature and loss of enzyme activity during immobilization process of the NMs affects the thermal and chemical stability, sensitivity, and reproducibility of the biosensor. (Daneshpour et al., 2016; Yin et al., 2014; Zhang et al., 2007). Beside NMs, various enzymes e.g. alkaline phosphatase (AP), horseradish peroxidase (HRP), glucose oxidase (GOx) etc are also considered most popular groups of tracer labels. These enzymes have been immobilized on various materials to enhance the enzymatic catalytic signals (Akanda and Ju, 2017; delle Noci et al., 2008; Haque et al., 2017a; Kaatz et al., 2012; Lai et al., 2009; Singh et al., 2013; Zhang et al., 2017). Another advantage of using the enzymes is direct electron transfer between the electrode and enzyme label, which could improve the signal in a biosensor. However, a large electron-hopping distance between the electrode and the redox center of the enzyme label could create few challenges for efficient electron transfer in biosensor (Chen et al., 2007). To overcome these limitations, mediated electron transfer can be achieved using a redox mediator that conveys electrons between the electrode and the redox center present in solution. This process is known as electrochemical enzymatic (EN) redox cycling reaction (Singh et al., 2013). In this reaction, electroactive molecules such as Ru complexes ([Ru(NH)₆]³⁺) undergo a fast outer-sphere electron-transfer reaction at the electrodes as well as electron-transfer reactions with redox enzymes (Chen and McCreery, 1996).

Herein, we introduced a simple and sensitive assay for the detection of global DNA methylation using EN redox cycling reaction. In our assay, extracted genomic DNA was denatured at 95 °C to obtain ssDNA. Then, 5 µL of denatured ssDNA samples were dropped onto a screen-printed gold electrode (SPGE) followed the protocol of REPLI-g whole genome amplification mini kit (Qiagen, Hilden, Germany) and purified following the protocol DNeasy Blood & Tissue Kits (Qiagen).

2.1. Reagents and chemicals

Antibody-GOx conjugation kit (ab102887) was purchased from Abcam, Inc. (Cambridge, MA, USA). GOx-conjugated 5mC antibody were prepared in phosphate buffer (50 mM, pH 7.4) according to the manufacturer’s procedure. Hexaammineruthenium (III) chloride (RuHe₆)₂, glucose, bovine-serum albumin (BSA), and all reagents for buffer solutions were obtained from Sigma-Aldrich, Co. and New England Biolab (NEB). Tris buffer comprised 50 mM tris(hydroxymethyl)aminomethane (pH 8.0). Phosphate- buffered saline (PBS, pH 7.4) comprised 10 mM phosphate, 0.138 M NaCl, and 2.7 mM KCl.

2.2. Isolation and purification of genomic DNA

Two ovarian cancer cell lines (SKOV 3, OVCAR 3) and one normal cell line (MeT-5A) were purchased from American Type Culture Collection (ATCC), USA. A blood and cell culture DNA mini kit (Qiagen, Hilden, Germany) was used to extract and purify the DNA from these cell lines. Jurkat genomic (100% methylated) DNA was purchased from New England BioLab. Whole genomic amplified DNA (WGA) was prepared by following the protocol of REPLI-g whole genome amplification mini kit (Qiagen, Hilden, Germany) and purified following the protocol DNeasy Blood & Tissue Kits (Qiagen).

2.3. Electrochemical detection of global DNA methylation

20 µL of isolated genomic DNA were denatured at 95 °C followed by the dilution with 5X saline sodium citrate (SSC) buffer and 1% BSA to obtain 10 ng/µL of ssDNA. Then, 5 µL of denatured ssDNA samples were dropped onto a screen-printed gold electrode (SPGE) (WE: gold (diameter: 4 mm), CE (platinum) and RE (silver) purchased from Dropens, Spain) surface and allowed to adsorb for 10 min. 1% BSA was used to block the uncovered electrode surface area with target DNA to minimize the non-specific adsorption of biological molecules. 5 µL of GOx-5mC antibody (20 ng/µL) was added onto the gold electrode surface and allowed to incubate for 30 min at room temperature with gentle shaking (unless otherwise stated) to facilitate the binding of the GOx-5mC antibody with the designated methylocytosine site of ssDNA. After the incubation, the electrode was washed with 10 mM PBS (pH 7.4) to remove loosely bounded or unbounded GOx-5mC antibody. A CH1040C potentiostat (CH Instruments, TX, USA) was used to perform the electrochemical measurements. The CC responses were recorded in the presence of 60 µL of glucose substrate solution (2 mM) mixed with 10 µM RuHe₆ in Tris buffer (pH 8.0) solution onto the gold electrode surface with a potential + 50 mV for 100 s. At least three replicates were measured at room temperature for each standard/sample and all data were subtracted from the background data.

2.4. Determination of the surface area of the electrodes

The effective working area of the electrodes was calculated by measuring the peak current obtained as a function of scan rate under cyclic voltammetric conditions for the one-electron reduction of [Fe(CN)₆]⁴⁻ (2.0 mM in PBS (0.5 M KCl)) and by using the Randles-Sevčík equation (Eq. (1)) (Bard and Faulkner, 2001; Shiddky et al., 2010),

\[
\text{current} = \frac{2.69 \times 10^5 \times \text{concentration} \times \text{scan rate}^{1/2}}{\text{surface area}}
\]
\[ i_p = (2.69 \times 10^4) n^{3/2} D^{1/2} C^{3/2} \]  

where, \(i_p\) is the peak current, \(n\) is the number of electrons transferred (Fe\(^{3+}\) \rightarrow Fe\(^{2+}\), \(n = 1\)), \(A\) is the active area of the electrode (cm\(^2\)), \(D\) is the diffusion coefficient of [Fe(CN)\(_6\)]\(^{3-}\) taken to be 7.60 \times 10^{-5} \text{ cm}^2 \text{s}^{-1}\), \(C\) is the concentration (mol cm\(^{-3}\)), \(v\) is the scan rate (Vs\(^{-1}\)).

#### 2.5. Glucose oxidase kinetics

The Steady-state kinetic experiments of glucose and RuHex were analysed using the Michaelis–Menten and Lineweaver–Burk models. The steady-state kinetic experiments were performed using 100 ng of GOD for both glucose substrate and Ru(NH\(_3\))\(_6\)\(^{3+}\) mediator by varying the concentrations of (A) glucose (0.01–5.00 mM) and (B) RuHex (0.01–100.0 µM) with fixed amounts of (A) RuH (10 µM) and (B) glucose (2 mM), respectively. To obtain the corresponding electrochemical signals from these samples, chronoamperometric responses were taken at 60 s by applying a fixed potential of + 50 mV.

Kinetic parameters were estimated based on the simplified Michaelis–Menten and Lineweaver–Burk equations (Eq. (2)) (Lehninger et al., 2005; Masud et al., 2017; Tanaka et al., 2018).

\[ \frac{J}{V} = \frac{K_m}{V_{max}} \frac{1}{[S]} + \frac{K_m}{V_{max}} \]  

In this equation, \(J\) is the rate of conversion, \(V_{max}\) is the maximum rate of conversion, \([S]\) is the substrate concentration, and \(K_m\) is the Michaelis constant which is equivalent to the substrate concentration at which the rate of conversion is half of \(V_{max}\). \(K_m\) denotes the affinity of the enzyme for the substrate.

#### 3. Results & discussion

##### 3.1. Assay principle

In this work, methylation specific antibody (5mC) was used to specifically recognize methyl group in genomic DNA. To read the methylation level, 5mC was conjugated with glucose-oxidase (5mC-GOx), and electrochemical-enzymatic redox cycle was performed. Briefly, double-strand genomic DNA were denatured to generate ssDNA and allowed to adsorb on SPGE. After DNA adsorption, 5mC-GOx was added onto the electrode surface to complete the immunorecognition process. Here, 5mC is specific to methylated site of the ssDNA and 5mC-GOx only attaches to the methylated DNA on the electrodes. In the presence of GOx, Ru(NH\(_3\))\(_6\)\(^{3+}\) (Ru\(^{3+}\)) was reduced to Ru(NH\(_3\))\(_6\)\(^{2+}\) (Ru\(^{2+}\)) and glucose was oxidised to gluconic acid. During this process, Ru\(^{2+}\) oxidised back to Ru\(^{3+}\) and completed the redox cycle. Therefore, increased chronocoulometric response in electrochemical readout is directly proportional to the amount of GOx, which is successively proportional to the methylation level in the sample. The overall reaction process is known as ping-pong mechanism and shown in Eqs. (3)–(5) (Gorski, 2001; Singh et al., 2013; delle Noci et al., 2008). The working principle of this assay is depicted in Fig. 1.

**Glucose + GOx (Ox) \rightarrow GOx(Red) + Gluconic acid**  

\[ 2\text{Ru}^{3+} + \text{GOx}(\text{Red}) \rightarrow \text{GOx}(\text{Ox}) + 2\text{Ru}^{2+} + 2\text{H}^+ \]  

\[ 2\text{Ru}^{2+} \rightarrow 2\text{Ru}^{3+} + 2e^- \]  

#### 3.2. Assay optimization

To get the optimum performance of the assay, several experimental conditions (i.e. the amount of target DNA, pH, incubation time etc.) need to be optimized. Previously, we have optimized the major adsorption parameters such as adsorption time, pH of the solution and amount of input DNA to obtain the maximum electrochemical responses between methylated and unmethylated DNA targets (Haque et al., 2017b; Koo et al., 2015; Sina et al., 2014). In the current assay, we have adopted all these optimized parameters and unless otherwise stated, we directly adsorbed 50 ng of target DNA (diluted in 5X SSC buffer, pH 7.4) onto the unmodified SPGE for 10 min. Conventional BSA blocking was also used to prevent the non-specific adsorption of target DNA onto the gold electrode surface. As can be seen in Fig. S1, the difference between the responses at the BSA-blocked and −unblocked electrodes is about 22 µC cm\(^{-2}\), which is almost 1.15-times higher than that of the electrodes with 1% BSA blocking. Notable, the total charge from 50 ng of WGA at the electrode without BSA blocking is significantly higher which indicates the non-specific adsorption of the biomolecules onto the electrode surface.

In the case of enzymatic redox cyclic reaction, Ru\(^{3+}\) undergo fast outer-sphere electron transfer reaction at electrodes which create lot of background noise (Akanda et al., 2012; Singh et al., 2013; Tanimoto and Ichimura, 2013). Thus, it is important to optimize the concentration of glucose substrate solution and Ru\(^{3+}\) solution. To obtain the maximum signal to noise ratio, electrooxidation of Ru\(^{3+}\), the enzymatic reduction of Ru\(^{3+}\) and enzymatic oxidation of glucose by GOx should be faster. However, without GOx in the system, the direct electro-oxidation of glucose and direct electron-transfer between glucose to Ru\(^{3+}\) should be slower to maintain a low background noise. These requirements were investigated prior to use our assay for detecting methylation level. We used Tris buffer as the electrolyte for the electrochemical readout step (pH 8.0) was selected from a previously published work (Singh et al., 2013)) and compared the electrochemical responses in the presence or absence of glucose oxidase in the system. Fig. S1 shows the comparison of chronocoulometric responses obtained from Tris buffer (pH 8.0), 2.0 mM glucose (in tris), 2.0 mM glucose and 10 µM RuHex (in tris) in the absence and presence of GOx. In this assay, we applied + 50 mV of potential to measure the electrochemical readout which is much higher than the formal potential of Ru\(^{3+}/\text{Ru}^{2+}\) (ca.-150 mV) to faster the electro-oxidation process in the system. This is due to the CC charge that generates during Ru\(^{3+}/\text{Ru}^{2+}\) redox cycling is controlled only by mass transfer of Ru\(^{2+}\) to the electrode not by the kinetics of Ru\(^{3+}\) electrooxidation (Gorski, 2001; Singh et al., 2013). However, we found that the surface response on Tris buffer and glucose substrate solution without RuH and GOx shown almost similar response (20 µC/cm\(^2\) vs 21 µC/cm\(^2\), intercept at Y-axis) but obvious improved (charge density of about 35 µC/cm\(^2\)) response was observed when RuHex was added in the solution (blue line). Once GOx present in the redox system, the overall enzymatic reaction improved and the response in the CC readout was also improved (charge density of about 250 µC/cm\(^2\)). These results signify the slower electrochemical oxidation and electron transfer in glucose and between glucose and Ru\(^{3+}\) when GOx is not present. However, oxidation, reduction improved the enzymatic redox reaction thus shown a high CC response in the Fig. S2.

The concentration of glucose and RuHex solutions are also need to be optimised to achieve the optimal catalytic performances. We constructed Michaelis–Menten model and Lineweaver–Burk model experiments by changing the glucose substrate concentration between 0.01 nM and 5.00 mM, and RuHex concentration between 0.01 µM and 100.0 µM, respectively. In Fig. 2(A), maximum current density (\(I_{\text{max}}\)) before saturation for glucose was obtained at 2.0 mM concentration (RuHex 10.0 µM, GOx 100.0 ng/µL), on the other hand, RuHex showed the \(I_{\text{max}}\) for the EN reaction at 10 µM (glucose 2.0 mM, GOx 100.0 ng/µL) in Fig. 2(B). We further calculated the \(V_{\text{max}}\) and \(K_m\) for both glucose and ruthenium complex from Lineweaver–Burk (shown inset in Fig. 2). The measured \(V_{\text{max}}\) and \(K_m\) values for glucose were 2.04 µM\(^{-1}\) and 0.27 mM, respectively which were much higher than RuH (\(V_{\text{max}} = 1.72 \mu\text{M}\) and \(K_m = 0.22 \mu\text{M}\)). These findings suggested that, the oxidation of glucose by GOx enzyme was faster than the Ru\(^{3+/2+}\) transformation. From these experiments, we selected glucose concentration as 2.0 mM and RuHex concentration 10.0 µM for our entire experiment. Then we optimised the GOx concentration and...
incubation period for the assay. Fig. S3(A) shows that the charge density increased with the GOx concentration from 10.0 ng to 100.0 ng. After 100 ng to 1.0 µg of GOx concentration, the charge response did not change much, thus we have selected 100.0 ng of GOx for our assay. The incubation period for antibody was selected 30 min as the charge density started to be stable after 30 min Fig. S3(B).

3.3. Selectivity of the assay

The use of target specific antibody in biosensor as recognition element increases the selectivity of the assay. To investigate the assay specificity, we incubated equal quantity (50 ng) of methylated (Jurkat) and unmethylated (WGA) DNA on SPGE electrodes surface and measured the CC response for both samples in the presence of 5mC-GOx antibody. Fig. 3(a) shows a significant difference (more than 6 times) in charge density between two samples (Jurkat is 246 µC/cm² and WGA 37 µC/cm² Fig. 3(b)). For further verification, we also carried out two additional sets of experiment where only PBS was used instead of target DNA (referred as NoT) and a positive sample without 5mC-GOx antibody (referred as control). Both the NoT and control showed negligible responses in electrochemical measurements (17 µC/cm² Vs 18 µC/cm²). The control studies demonstrate that the assay response is very much dependent on the presence of methylation present in the sample. Altogether the response from controls and synthetic samples clearly demonstrates excellent level of specificity of our assay toward the detection of DNA methylation with negligible background response.

3.4. Sensitivity of the assay

As samples collected from patients with cancer usually contain a mixture of methylated and unmethylated DNA, it is therefore important to detect the degree of methylation in a high background of unmethylated DNA samples. To evaluate our assay performance for detecting heterogeneous DNA methylation pattern, we measured CC responses for samples with various degree of methylation. The samples were prepared by mixing Jurkat (100% methylated) and WGA (0% methylated) DNA to get 0–100% methylated samples. Under the optimum experimental conditions, the CC responses for various methylation levels show a linear relationship between the charge density and the methylation percentage (Fig. 4, charge density for each percentage was calculated by subtracting the control value from that of the sample value). The calibration plots in Fig. 4(b)(inset) showed good linearity from 5% to 100% of methylation with a correlation coefficient, R² of 0.963 (n = 3). From the charge density values, the limit of detection (LOD) was estimated to be as low as 5% of methylation which was easily distinguishable from the control or 0% of methylation (i.e. control Vs 5% was 18 µC/cm² Vs 57 µC/cm²). A relative standard deviation (% RSD) for three independent measurements were calculated to be < 5.0%, suggesting the good reproducibility of assay toward the qualitication of heterogeneous DNA methylation pattern.

![Fig. 1. Schematic representation of the assay to detect global DNA methylation. Initially, cell lines were grown and extracted using commercial kit. Then, denatured ssDNA was adsorbed onto a SPGE surface followed by immunorecognition of methylated DNA using the GOx-5mC antibody. Subsequent detection of the genomic DNA methylation pattern was performed through enzymatic redox cyclic reaction.](image)

![Fig. 2. Steady-state kinetic analyses using the Michaelis–Menten model (main) and Lineweaver–Burk model (inset) for the GOx by varying the concentrations of (A) glucose (0.01–5 mM) and (B) Ruthenium complex (0.01–100.0 µM) with fixed amounts of (A) RuHex (10 µM) and (B) glucose (2 mM), respectively.](image)
The sensitivity (5% methylation), reproducibility (< 5.0% RSD), input DNA amount (50 ng), and time (1 h) required for our assay is highly comparable with the recently published DNA analysis methods (Bhattacharjee et al., 2018b; Haque et al., 2017a; Wee et al., 2015). For example, Wee et al. have demonstrated that 5% of methylation can be detected from 50 ng of DNA input within 2 h of assay time (Woe et al., 2015). Recently, Povedano and colleagues have reported a sandwich type assay for detecting global and regional DNA methylation patterns within 1 h (Povedano et al., 2018). In this work, for detecting global DNA methylation, a sandwich immunosensor involving the use of two different antibodies was fabricated. Anti-5mC was first immobilized on the surface of carboxylic acid-modified magnetic beads which capture ssDNA sequence bearing 5-mC marks. A second antibody conjugated with HRP was then used to recognize any ssDNA as well to produce electrochemical signal. In contrast, our method is based on the direct adsorption of target DNA onto a bare gold electrode. The adsorption of ssDNA target onto the bare gold electrode follows the gold-DNA affinity interaction (Koo et al., 2015). The use of the direct adsorption of target DNA samples on a bare gold electrode/surface rather than the conventional approach of using recognition and transduction layers in DNA biosensors is substantially important and novel, as it significantly simplifies the method by avoiding the complicated chemistries underlying each step of the sensor fabrication. Additionally, in our method, an electrochemical-enzymatic redox cycling reaction was used for the first time to achieve global DNA methylation analysis.

### 3.5. Analytical application in cell line samples

To verify the applicability of our assay in analyzing complex biological samples, we examined DNA samples derived from two ovarian cancer (SKOV 3 and OVCAR 3) and one normal cell lines (MeT-5A) (Fig. 5). First, DNA was extracted and purified from these cell lines. Fully methylated (100% methylation) Jurkat and fully unmethylated (0% methylation) WGA DNA were used as internal controls. As expected, both the cell lines (charge density for SKOV 3 and OVCAR 3 was about 177 and 146 µC/cm² respectively) showed relatively higher CC response compared to that of fully unmethylated WGA (18 µC/cm²) sample. As expected these responses were much lower compared to that of fully methylated Jurkat DNA (227 µC/cm²) (Fig. 5A). These variations suggest the present of the different level of methylation pattern in these cell lines. Furthermore, the charge density (Fig. 5B) derived from these cell lines are easily detectable compare to those of the control Jurkat and WGA samples. These findings suggest that the methylation level of normal cell line MeT-5A (67 µC/cm²) is much lower (about 18%) than that of the other two cancer cell line (SKOV 3 and OVCAR 3). The level of CC responses also indicate that the methylation patterns in both cancer cell lines (SKOV 3 and OVCAR 3) could exhibit more than 75% of methylation (by comparing the data shown in Fig. 4(B) and Fig. 5). A similar level of methylation patterns was also observed for these cell lines in other published work (Tomar et al., 2016).

### 4. Conclusions

In conclusion, we have developed a sensitive and selective biosensor for the detection of global DNA methylation. We took the advantages of 5mC antibody for selective binding to the 5-mC site and GOx enzyme to enhance the sensitivity via enzymatic redox cycling reaction. The assay also avoids the use of bisulfite treatment or PCR amplification steps. It uses the direct adsorption of target DNA samples on a bare gold electrode rather than the conventional approach of using recognition and transduction layers in hybridization based DNA biosensors, which makes it relatively simpler in compare with the conventional hybridization based methylation assays. Under the optimal conditions, this biosensor exhibited high sensitivity (5% of DNA methylation) with...
Fig. 5. Analysis of ovarian cancer cell lines and normal cell line. (A) chronocoulometrics response for different cell line and (B) represent charge density comparison between different cell lines.

the %RSD of < 5% (for n = 3) for the analysis of heterogeneous DNA samples derived from ovarian cancer cell lines. We believe that the method is not just limited to ovarian cancer DNA, but can easily be extended towards detection of global DNA methylation in samples collected from a diverse range of cancer types. In addition to this, we predict that our electrochemical method could have broad applications in the field of epigenetic research and with further improvements, can potentially find its relevance as a low cost, easy to operate and portable diagnostic tool during cancer diagnosis and treatment.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.bios.2018.10.020.

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