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The expression of microRNAs (miRNAs) is related to some cancer diseases. Recently, miRNAs have emerged as new candidate diagnostic and prognostic biomarkers for detecting a wide variety of cancers. Due to low level, short sequences and high sequence homology among family members, the quantitative miRNAs analysis is still a challenge. A novel electrochemical biosensor with triple signal amplification for the ultrasensitive detection of miRNA was developed based on phosphatase, redox-cycling amplification, bimetallic Pd-Pt supported graphene functionalized screen-printed gold electrode, and two stem-loop structured DNAs were used as target capturers. The proposed biosensor is highly sensitive due to enhanced electrochemical signal of Pd-Pt supported graphene and sufficiently selective to discriminate the target miRNA from homologous miRNAs in the present of loop-stem structure probes with T4 DNA ligase. Therefore, this strategy provided a new and ultrasensitive platform for amplified detection and subsequent analysis of miRNA in biomedical research and clinical diagnosis.

1 Introduction

MicroRNAs (miRNAs), small endogenous noncoding RNAs with approximately 22 nucleotides, have attracted numerous researchers’ attention due to their important roles in a variety of biological processes such as cell differentiation, apoptosis, proliferation, and immunological response. Recently, miRNAs have emerged as new candidate diagnostic and prognostic biomarkers for detecting a wide variety of cancers. Many methodologies, including Northern blotting, miRNA arrays, and real-time reverse transcription polymerase chain reaction, have been developed to detect miRNA expression. Unfortunately, these methods may be low sensitive, labor-intensive, or required a large amount of RNA samples, etc. Another great challenge for quantitative miRNA analysis is high sequence homology among family members. The let-7 family is one of the well-studied miRNA families, which consists of 9 highly homologous miRNAs (let-7a to let-7g, let-7i, and mir-98). Because their sequences are similar to each other and nucleotide variations exist at several positions, it is difficult to exclusively discriminate let-7 family members, which interferes with our deep understanding of the specific function of particular let-7 miRNAs. Therefore, it is of great importance to develop some simple, sensitive, and specific methods for miRNA detection in complex biological samples. The electrochemical technique offers attractive advantages in high sensitivity, inherent simplicity, and low cost, and has been widely used in bioassay. Various signal amplification strategies, such as gold nanoparticle-based assay, enzymatic assay, rolling circle amplification, and polymerization-based assay, have been used in developing sensitive electrochemical genosensors for the detection of low-abundance miRNAs. However, single amplification could still not satisfy the requirements of ultrasensitive detection of miRNAs, thus multiple signal amplifications were employed. An enzyme amplification technology coupling with a redox-cycling reaction to acquire ultrasensitive detection of miRNAs was reported. He et al. presented a label-free and highly sensitive electrochemical genosensor for miRNAs detection with the triple signal amplification of gold nanoparticles (AuNPs), alkaline phosphatase (ALP) and p-aminophenol (p-AP) redox cycling. Capturer DNA was directly anchored on the work electrode that may disturb the electrochemical signals output.

Herein, we designed a novel electrochemical biosensor with triple signal amplification for ultrasensitive let-7b detection in...
two steps. Firstly, electrochemical signal source is obtained by enzyme amplification technology. Secondly, Pt/Pd/RGO modified screen-printed gold electrode (Pt/Pd/RGO/SPGE) was used to detect and enhance the electrochemical signal with a redox-cycling reaction. The detection limit of 3.55 fM is similar to electrochemical detection of microRNAs based on triple signal amplification of multifunctional gold nanoparticles, enzymes and redox-cycling reaction with a detection of 3 fM, but the linear range of our sensor from 10 fM to 0.1 nM is wider. This method significantly increases the analysis sensitivity and provides a new concept in the development of high selective microRNA biosensor for gene expression profiling and molecular diagnostics.

2 Experimental

2.1 Materials

Dopamine hydrochloride, Tris (2-carboxyethyl) phosphine hydrochloride (TCEP) and 1-naphthyl phosphatedisodium salt (p-NPP) were purchased from Sigma-Aldrich. Tris (hydroxymethyl) aminomethane, iron trichloridehexahydrate (FeCl₃·6H₂O), anhydrous NaAc, ethylene glycol (EG), and 1,6-diaminohexane were purchased from Tianjin Chemical Research Institute (Tianjin, China). All chemical reagents were of analytical grade and used as received without further purification. Water was purified with a Millipore Milli-Q water system to achieve a final resistance of 18.2 MΩ. All solutions were prepared using ultrapure water treated with 0.1% DEPC.

The used oligonucleotides were purchased from Shanghai Sangon Biological Engineering Technology & Services Co. (Shanghai, China). Let-7a, let-7b, let-7c, let-7d, and miR-21 were obtained from Shanghai GenePharma Co., Ltd. (Shanghai, China). The sequences are illustrated in Table S1.

2.2 Apparatus

UV-vis spectra were recorded on a UV-3600 spectrophotometer (Shimadzu, Kyoto, Japan). Fluorescence spectra were recorded from a RF-5301PC spectrofluorophotometer (Shimadzu). Transmission electron micrographs (TEM) were measured on a JEOL JEM 200CX transmission electron microscope, using an accelerating voltage of 200 kV. Scanning electron micrographs (SEM) were obtained with a Hitachi S4800 scanning electron microscope. X-ray-photoelectron spectroscopy (XPS) analysis were recorded with an ESCALAB 250 spectrometer (Thermo-VG Scientific, USA). Cyclic voltammetry (CV) and differential pulse voltammetry (DPV) were performed on CHI 660C electrochemical work station.

2.3 Modification of the screen-printed gold electrode (SPGE)

The screen-printed three gold electrodes were purchased from Drop Sens (Spain) including a gold electrode as the working electrode, a platinum electrode as the counter electrode, and a silver electrode as reference electrode. Prior to experiments, the screen-printed gold electrode (SPGE) was first scanned in 0.5 M H₂SO₄ from 0 V to 1.5 V at 100 mV s⁻¹ until a reproducible cyclic voltammogram (CV) was achieved. For electrodeposition of multilayered graphene films, 1.0 mg mL⁻¹ graphene oxide (GO) in carbonate buffer solution (10 mM, pH=9.0) was prepared. The colloidal graphene oxide was electrochemically reduced to yield multilayered graphene films on SPGE surface by cyclic voltammetry from 0.6 V to -1.5 V with a scanning rate of 25 mV s⁻¹ and potential step of 1 mV for 50 rounds. Then a mixture containing 2 mM PtCl₆²⁻ and 2 mM PdCl₂ was dropped onto RGO/SPGE surface to obtain Pt/Pd/RGO/SPGE electrode by a co-electrodeposition using an amperometric i-t curve method from -0.2 V to 0.1 V for 100 s.

2.4 Preparation of Fe₃O₄@PDA

Fe₃O₄ beads were prepared following a previously reported protocol with minor modification. Briefly, 1.0 g of iron trichloridehexahydrate (FeCl₃·6H₂O), 2.0 g of sodium acetate anhydrous, and 6.5 g of 1,6-diaminohexane were added to 30 mL of ethylene glycol with the help of ultrasound at room temperature. Then, the mixture was maintained at 473 K for 6 h in a Teflon-lined stainless steel autoclave (50 mL capacity). The black Fe₃O₄ were separated by magnetic field, washed with deionized water and ethanol alternately several times, and dried at 323 K in a vacuum drying oven. 10 mg of Fe₃O₄ nanospheres was dispersed in 30 mL Tris-HCl (10 mM, pH 8.5) and ultrasound for 30 min. After that, 20 mg of dopamine was added and shaken for 4 h to obtain polydopamine coated Fe₃O₄ (Fe₃O₄@PDA), which was separated by magnetic field, washed with deionized water and ethanol alternately several times. The obtained Fe₃O₄@PDA was dispersed in Tris-HCl buffer (10 mM, pH=9.0) for further use.

2.5 Preparation of Fe₃O₄@PDA-DNA1

The thiol-DNA1 was activated with 10 mM TCEP in pH 5.2 acetate buffer for 1 h to prevent the terminal cysteine from forming disulphide bonds. Afterwards, 420 μL of 10 μM thiol-DNA1 was mixed with 1 mL Fe₃O₄@PDA (1 mg mL⁻¹) and reacted overnight at 25°C, then at 4°C for 12 h. Subsequently, Fe₃O₄@PDA-DNA1 was separated by magnetic field, washed with Tris buffer solution several times. The obtained Fe₃O₄@PDA-DNA1 was dispersed in Tris-HCl buffer (pH=7, 10 mM) for further use.

2.6 RNA Hybridization Assay and Electrochemical Measurements

In a typical miRNA assay, 5μL 1 mg mL⁻¹ Fe₃O₄@PDA-DNA1 was mixed with 5 μL 5 μM DNA2, 1μL let-7b at various assigned concentrations, 1.0 μL 40U μL⁻¹ RNase Inhibitor and 1μL 1000U μL⁻¹ T4 DNA ligase at 37°C for 4 h to perform the hybridization reaction. Fe₃O₄@PDA-DNA1/let-7b/DNA2 was separated by magnetic field, washed with Tris-HCl buffer several times. Then 10 μL10 μg mL⁻¹ SA-ALP was added into the obtained Fe₃O₄@PDA-DNA1/let-7b/DNA2 was separated by magnetic field, washed with Tris-HCl buffer several times. After that, Fe₃O₄@PDA-DNA1/let-7b/DNA2/SA-ALP composite was washed with 1% BSA and then filled with a Tris buffer solution (pH=9.6, 100 μL, containing 10 mM Mg²⁺) containing 5 mM p-NPP. The supernatant was collected and the
differential pulse voltammetric (DPV) response was then recorded after 15 min from -0.2 to +0.5 V using Pt/Pd/RGO/SPGE electrode with a step potential of 4 mV, and a scan rate of 50 mV s⁻¹ in the absence or presence of 1 mM TCEP, respectively. The TCEP solution was freshly prepared for each experiment. All measurements were repeated for a minimum of three times with separate electrodes to obtain statistically meaningful results.

2.7 Polyacrylamide Gel Electrophoresis Analysis

8% polyacrylamide gel was employed to verify the binding of let-7b with DNA1 and DNA2. Firstly, 2 μL DNA1 (10 μM) and 2 μL DNA2 (10 μM) was incubated with 2 μL let-7b (10 μM) at 37 °C for 4 h to form stable DNA hybrid by a base paring hybridization in Tris buffer (10 mM, pH 7.4). Electrophoresis was carried out in TBE buffer (pH 8.0) at 100 V for 45 min at room temperature. After separation, the gel was stained with ethidium bromide and imaged by Bio-Rad imaging system (Hercules, CA).

2.8 Cell Culture and Cell Lysis

Human breast adenocarcinoma cell line (MCF-7), human cervical cancer cell (HeLa cells) and human lung carcinoma cells (A549), were grown at 37 °C in DMEM (Dulbecco’s Modified Eagle Medium) supplemented with 10% (v/v) fetal bovine serum (FBS), 100 U/mL penicillin, and 100 μg/mL streptomycin in a 5% CO₂ environment. After growing to 90% confluence, the cells were washed three times with PBS and the cell number was estimated by a hemocytometer. A 1.0 mL confluence, the cells were washed three times with PBS and the cell number was estimated by a hemocytometer. A 1.0 mL amount of ~5 x10⁵ suspended cells was added into 6-well plate for 12 h. Total RNA from these cells was extracted using TRIzol reagent according to the manufacturer’s recommended protocol. RNA concentration was determined by UV-vis absorption at 260 nm.

2.9 Let-7b detection by qRT-PCR

The total cellular RNAs were extracted from HeLa, NIH3T3 and HL-60 cells using Trizol reagent (Invitrogen) according to manufacturer’s protocols. The first-strand cDNA was generated using AMV reverse transcriptase (Takara Bio, Shiga, Japan) and a stem-loop RT primer (Invitrogen). Quantitative PCR was carried with an ABI 7300 Sequence Detection System (Applied Biosystems) using SYBR Green PCR Master Mix. For analysis of the let-7b level, we used 5'-ACACTCCAGCTGGGTGAGG TAGTAGTTGT-3’ as the sense primer and 5'-CCTCAACTGG TGTCGTTGAGTCCGAAATTCAGTTGAGAACACAC-3’ as the antisense primer, as well as 5'-CTGCCGTTGCGCAGC ACA-3’ and 5’-AACGCTTTACCGATTGCGT-3’ for U6. The relative expression of let-7b was calculated using the 2-ΔCT method, in which ΔCT=CTlet-7b-CTU6. All RT-PCR reactions were performed in triplicate.

3 Results and discussion

3.1 The mechanism and characterization of the capture probe

Self-polymerization of dopamine (DA) easily produced a surface-adjacent polydopamine (PDA) nanolayer deposited on the surface of magnetic Fe₃O₄ nanoparticles (Fe₃O₄@PDA) at weak alkaline pH 24,25 (Fig. 1A). It is simple to separate and wash Fe₃O₄ carriers under magnetic field in the decoration process. Transmission electron microscope was used to characterize the modification procedure of Fe₃O₄@PDA. PDA shell was observed on the surface of Fe₃O₄ nanoparticles (Fig.1B and C). PDA contained the oxidized quinone form of catechol that can undergo reactions with various functional groups, including thiol, amine, and quinone itself26. Therefore, Fe₃O₄@PDA can be further modified by thiol-terminated DNA1 via Michael addition. UV-vis techniques were used to characterize the modification procedure of Fe₃O₄@PDA-DNA1. After the conjugation of DNA1, an obvious UV emission peak around 260 nm could be observed, which was attributed to the characteristic absorbance band of the nucleic acid, indicating the successful attachment of DNA1 to Fe₃O₄@PDA (Fig.S1).

Scheme 1 illustrates the design of the genosensor for let-7b via sandwich-type hybridization assay. Two stem-loop structured DNAs were used as capturers; the stem-loop structured hairpin DNA with its excellent specificity has been widely explored for discrimination testing since stem-loop probes not only prevented their self-ligation and nonspecific
hybridization with precursor miRNAs but also allowed precise
distinction of a single mutant in the miRNA sequence27,28.
DNA1 has 11 bases at the 3’-end complementary to half part of
let-7b as shown in Table S1. Upon hybridization with the target
let-7b, the hairpin structure of DNA1 was unfolded, and
meanwhile the other half 11 bases of let-7b was further
hybridized with probe 2 (DNA2) that modified by biotin at the
3’ end. In the process, T4 DNA ligase is used to efficiently
catalyze the joining between 3’-hydroxyl group of DNA1 and
5’-phosphomonoester group of DNA2 in the presence of let-7b
like a tack29. The hybridization reaction between the target and
the capture probes was analyzed by polyacrylamide gel
electrophoresis. As shown in Fig.S2, no band appeared in Lane
2 because the sequence of single-stranded let-7b is short and
difficult to be dyed by EB. Two bands in Lane 3 assigned to
DNA1 and disulphide DNA1, one band in Lane 4 assigned to
DNA2. In the presence of let-7b and T4 DNA ligase (Lane 8),
the emission band that is higher than 300bp indicated the
formation of DNA1/let-7b/DNA2/T4 complex because T4
DNA ligase is a polypeptide with high molecular weight so the
DNA1/let7b/DNA2/T4 complex exhibited a wide emission
band. Then, alkaline phosphatase-labeled streptavidin (SA-ALP)
was attached onto DNA2 via the specific affinity between
streptavidin and biotin. Alkaline phosphatase converted 1-naphthyl phosphate disodium salt (p-NPP) into electroactive 1-naphthyl (p-NPP).

3.2 The characterization of the electrode
The supernatant containing p-NP was collected and dropped
onto Pt/Pd/RGO/SPGE to obtain electrochemical signal.
Pt/Pd/RGO/SPGE was fabricated via the electrodeposition of
RGO, Pt and Pd on SPGE (Fig.2A). The detailed modification
procedure was described in the experiment section. The
preparation process was investigated by scanning electron
microscope (SEM). In comparison with Fig. 2B, graphene film
(RGO) could be observed on the surface of SPGE (Fig. 2C).
After deposition of Pt and Pd, bimetallic Pt/Pd nanoparticles
were attached on the RGO sheets as shown in Fig. 2D. The size
of Pt/Pd nanoparticles is about ~70nm as Fig.S3A showed. The
EDS pattern (Fig. S3B–E) displayed the typical diffractions of
C, Pt and Pd. XPS was further used to confirm the formation of
bimetallic Pt/Pd/RGO modified SPGE (Figure S4). The three

![Figure 2](image-url) (A) Preparation process of SPGE and the detection of miRNA. SEM images of SPGE(B), RGO/SPGE(C), Pt/Pd/RGO/SPGE(D).

![Figure 3](image-url) (A) Cyclic voltammograms of SPGE (a), RGO/SPGE (b), Pt/Pd/RGO/SPGE (c) in 1 mM K3Fe(CN)6, at a scan rate of 100 mV s⁻¹. (B) Cyclic voltammograms of p-NP in 5 mM p-NPP solution containing 10 μL, 0.1 mg mL⁻¹ ALP using SPGE(a), RGO/SPGE (b) and Pt/Pd/RGO/SPGE (c) electrode respectively. (C) The peak values of the current in Fig.3B. (a) 0.138V, (b) 0.113V, and (c) 0.086V.

peaks at 284.2, 70.7, and 334.9 eV shown in the XPS spectrum of
the electrode can be attributed to C1s, Pt4f, and Pd3d, respectively. The atomic percentages of C, Pt, and Pd were
79.11%, 12.52% and 8.36%, revealing the successful deposition
of RGO, Pt, and Pd onto RGO modified SPGE greatly improved
the conductivity of the electrochemical genosensor. As shown
in Fig. 3A, the peak potential difference (ΔEp) for the
Pt/Pd/RGO/SPGE (57 mV) was found to be smaller than that for
both the RGO/SPGE (70 mV) and SPGE (135 mV), while the
peak current is higher than the other two. Moreover, a highest
signal of p-NP could be observed using Pt/Pd/RGO/SPGE
(Fig.3B and C). The peak values of SPGE, RGO/SPGE and
Pt/Pd/RGO/SPGE are 0.138, 0.113, and 0.086V, respectively.
The peak potential decreased because the modification of RGO
and Pt/Pd lead to the decrease of the overpotential. These
results indicated that Pt/Pd/RGO/SPGE can enhance the
sensing performance by promoting electron transfer on the

![Figure 4](image-url) (A) Differential pulse voltammetric (DPV) responses for different target
let-7b concentrations: (a) 10⁻⁶ M, (b) 10⁻⁵ M, (c) 10⁻⁴ M, (d) 10⁻³ M, (e) 10⁻² M, (f) 10⁻¹ M, (g) 10⁻⁰ M, (h) 10² M, and (i) 10³ M. Inset is the relationship between
the peak current and the logarithm of the target RNA concentration. (B) The
linear relationship between the peak current and the logarithm of the target
dNA concentration from 10⁻³⁰ M to 10⁻¹⁴ M. ΔI was calculated by I-I0, where I and I0
are the peak current observed in the presence and absence of let-7b, respectively.
(C) DPV signals in presence of let-7a, let-7b let-7c let-7d and miR-21 (all at a
concentration of 10⁻⁶ M). Mix1: the mixture of let-7b (10⁻⁶ M) and let-7c (10⁻⁷ M),
Mix2: the mixture of let-7b (10⁻⁷ M) and let-7c (2×10⁻⁷ M), Mix3: the mixture of
let-7b (10⁻⁸ M) and let-7c (10⁻⁸ M). Each value is the average of three
measurements. (D) Relative expression level of let-7b in various cells obtained by
qRT-PCR and electrochemical biosensor (EC).
electrode/electrolyte interface. Meanwhile tris(2-carboxyethyl)phosphine (TCEP), which can reduce electroactive organic materials such as dehydroascorbic acid and quinone at a fast rate, was added to regenerated p-NP, realizing a redox-cycling reaction (Fig. 2A). The optimal concentrations of p-NPP and TCEP were 5mM and 1mM, respectively (Fig.S5 and Fig.S6). As we predicted, in the presence of TCEP, the peak current is about one time higher than that in the absence of TCEP, indicating that the constructed Pt/Pd/RGO/SPGE had strong signal amplification, which is valuable for the design of new types of sensitive electrochemical biosensors.

3.3 Quantitative detection of Let-7b

The sensitivity of the biosensor was investigated by varying the target miRNA concentrations. The current signals of p-NP after the sensor hybridized with different concentrations of let-7b were measured via differential pulse voltammetry (DPV). As shown in Fig. 4A, the absolute current value increased with the increased concentrations of let-7b in the concentration range of 10 fM–1 μM, and showed a good linear correlation with the logarithm of target miRNA concentrations in the range of 10 fM–0.1nM (Fig. 4B) The regression equation is as followed: ΔI (I_0-I_μA)=8.35+0.57×logC (M), R^2=0.975. The detection limit (S/N=3) was calculated to be 3.55 fM.

3.4 Specificity

To examine the selectivity of the biosensor, a comparison assay on mismatch targets and perfect complementary target, including the perfect complementary target (let-7b) and single-base mismatch target (let-7c), two-base mismatch target (let-7a) and four-base mismatch target (let-7d), and non-complementary sequence (miR-21) at a concentration of 10 nM, were performed respectively. As shown in Fig. 4C, the complementarity target let-7b leads to a current increase which is approximately two times higher than that for let-7c and five times higher than let-7a, while the let-7d and miR-21 caused only a small increase in the electrochemical signal, demonstrating that this biosensor has an excellent discrimination for similar miRNAs. Mixtures of let-7b and let-7c with different concentrations were detected as well. Let-7b/let-7c concentration ratios are 10:1, 5:1 and 1:1, respectively. It can be seen when the let-7b/let-7c concentration ratio decreased to 1:1, the electrochemical signal dramatically increased, demonstrating that let-7c will obstruct the signal at high concentration, while at low concentration there is no high influence to the current signal.

3.5 MicroRNA Detection in Cell Lysates

The sensor was also directly used to analyze miRNAs in real samples, the content of let-7b extracted from HeLa cells, MCF-7 cells and A549 cells. As shown in Fig. 4D, the order of the relative expression level is MCF-7 > A549 > HeLa using electrochemical method (EC), in compliance with the results obtained by qRT-PCR, which demonstrated the validity of as-proposed method. These results confirmed that the biosensor could be used for the determination of let-7b in real samples.

Moreover, for qRT-PCR, it is necessary to reverse-transcribe miRNA into cDNA. Then cDNA hybridized with two primers to realize amplification and finally probes were added to produce fluorescence signals. The process is complicated and skilled technicians are needed. Our proposed electrochemical biosensor can be used to detect miRNA directly, and easy to operate, providing a new, simple platform for the quantification of miRNAs and having a potential application for the clinical diagnosis.

4 Conclusions

In summary, we have developed an ultrasensitive electrochemical method for the detection of let-7b by integrating phosphatase amplification with redox-cycling amplification. Moreover, Pt/Pd/ RGO fabricated SPGE of excellent conductivity was used to enhance the electrochemical signal. A proportional relationship was observed between the logarithm of target let-7b concentration and the DPV peak currents in a linear range from 10 fM to 0.1 nM with a detection limit of 3.55 fM. At the same time, this proposed biosensor is sufficiently selective to discriminate the target miRNA from homologous miRNAs and non-complementary miRNAs due to the loop-stem structure of capture probes and T4 DNA ligase. Subsequently, the proposed strategy has successfully achieved the detection of human miRNAs from cancer cell-specific total RNA extracts. Therefore, this biosensor is an attractive candidate for the development of an accurate, selective, and ultrasensitive method for miRNA expression profiling and clinical diagnostics.

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Notes and references

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