

ARTICLE

Boronate Affinity-assisted Electrochemically Controlled ATRP for Ultrasensitive Electrochemical Aptasensing of Carcinoembryonic Antigen

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Abstract

As an acidic glycoprotein, carcinoembryonic antigen (CEA) is of great value as a broad-spectrum tumor marker in the differential diagnosis and surveillance of malignant tumors. In this work, we report an electrochemical aptasensor for the ultrasensitive and highly selective detection of CEA, taking advantage of the dual amplification by the boronate affinity-assisted electrochemically controlled atom transfer radical polymerization (BA-eATRP). Specifically, the BA-eATRP-based electrochemical aptasensing of CEA involves the capture of target antigens by nucleic acid aptamers, the covalent crosslinking of ATRP initiators to CEA antigens via the selective interactions between the phenylboronic acid (PBA) group and the *cis*-diol group of the monosaccharide residues, and the collection of the ferrocene (Fc) reporters via the eATRP of ferrocenylmethyl methacrylate (FcMMA). As CEA is decorated with hundreds of *cis*-diol groups, the BA-based crosslinking can result in the labeling of each CEA with hundreds of ATRP initiators; furthermore, the eATRP of FcMMA results in the surface-initiated growth of long-chain ferrocenyl polymers, leading to the tethering of each ATRP initiator-conjugated site with hundreds to thousands of Fc reporters. Such that, the BA-eATRP can result in the efficient labeling of each CEA with a plenty of Fc reporters. Under the optimized conditions, the BA-eATRP-based strategy enables the highly selective aptasensing of CEA at a concentration as low as $0.34 \text{ pg} \cdot \text{mL}^{-1}$, with a linear range of $1.0\text{--}1000 \text{ pg} \cdot \text{mL}^{-1}$. Besides, this aptasensor has been successfully applied to the quantitative analysis of CEA in human serum. The BA-eATRP-based electrochemical aptasensor is cost-effective and simple in operation, holding broad application prospect in the ultrasensitive and highly selective detection of CEA.

Keywords: Boronate affinity; Atom transfer radical polymerization; Electrochemical aptasensor; Carcinoembryonic antigen; Tumor marker; Signal amplification

1. Introduction

Carcinoembryonic antigen (CEA) is a cell adhesion-related acidic glycoprotein [1]. As a oncofetal antigen, CEA is normally secreted by the gastrointestinal tissue during fetal development and occurs at very low levels in the serum of healthy adults [1]. However, the level of CEA is often elevated in the serum of individuals suffering from various neoplastic conditions (e.g., colorectal cancer, gastric cancer, pancreatic cancer, lung cancer, breast cancer, and medullary thyroid cancer) [2] and some non-neoplastic

conditions (e.g., cirrhosis, pancreatitis, ulcerative colitis, hypothyroidism, and Crohn's disease) [3]. Due to the poor sensitivity and specificity, CEA is unable to act as a tumor marker for the diagnosis and screening of a specific cancer. However, it is of great value as a broad-spectrum tumor marker in the differential diagnosis and surveillance of malignant tumors [1,4]. As the cutoff value of CEA for clinical application is $5 \text{ ng} \cdot \text{mL}^{-1}$ [5], sensitive and selective detection of CEA is highly desired.

In recent decades, a diversity of methods have already been established for CEA detection, such as fluorescent [6,7], photoelectrochemical (PEC)

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[8–10], surface-enhanced Raman scattering (SERS) [11,12], colorimetric [13,14], electrochemiluminescent [15–18], and electrochemical biosensors [19–21], using such as antibodies [22,23], imprinted polymers [24,25], and aptamers [26–28] as the capture elements. Among them, electrochemical aptasensing of CEA has attracted much attention by virtue of the advantages of low detection cost, minimal matrix/background interference, simple instrument, and high sensitivity and selectivity [29,30]. Nevertheless, most of the previously developed electrochemical CEA aptasensors necessitated the use of sophisticated enzymatic reactions (e.g., DNAzymes) [31,32] or nanocomposites (e.g., nanoclusters) [33,34] for signal amplification, which is not practical for clinical applications.

By virtue of the simplicity, high efficiency, and cost-effectiveness, the grafting of polymer chains through reversible deactivation radical polymerization, such as atom transfer radical polymerization (ATRP) [35,36] and reversible addition-fragmentation chain-transfer (RAFT) polymerization [37,38], has attracted growing attention as an amplification strategy in the sensitive detection of biotargets [39,40]. Among them, ATRP involves the use of the alkyl halide initiators (R-X) and the lower oxidation state transition-metal activators (typically the Cu(I) complexes) [41,42]. Specifically, reduction of the initiators by the activators generates the radical species, which then react with monomers to form long-chain polymers [42]. In the electrochemically controlled ATRP (eATRP), the Cu(I) activators are generated from the electrochemical reduction of the Cu(II) deactivators [43], holding the merits of the improved control over polymer growth, better biocompatibility, and higher O₂ tolerance [40,44].

On the basis of the dual amplification by the boronate affinity (BA)-assisted eATRP (BA-eATRP), we report herein a cost-effective and easy-to-operate electrochemical aptasensor for the ultrasensitive and highly selective detection of CEA. As the capture elements, the DNA aptamers were attached to a gold electrode via the gold-sulfur self-assembly chemistry. After the capture of CEA antigens, the oligosaccharide chains of CEA were leveraged for the covalent crosslinking of ATRP initiators via the selective interactions between the phenylboronic acid (PBA) group and the *cis*-diol group of the monosaccharide residues [45,46]. Further, potentiostatic reduction of the Cu(II) deactivators resulted in the generation of the Cu(I) activators [40,44], triggering the eATRP of ferrocenylmethyl methacrylate (FcMMA) and thus the surface-initiated growth of long-chain ferrocenyl

polymers. As each CEA is decorated with hundreds of *cis*-diol groups for the BA-based cross-linking of ATRP initiators while each ATRP initiator-conjugated site can be tethered with hundreds to thousands of Fc reporters, the BA-eATRP can result in the labeling of each CEA with a plenty of Fc reporters. As a result, the BA-eATRP-based electrochemical CEA aptasensor is highly sensitive and selective. Moreover, it is cost-effective and easy-to-operate, thus holding great promise in clinical applications.

2. Experimental

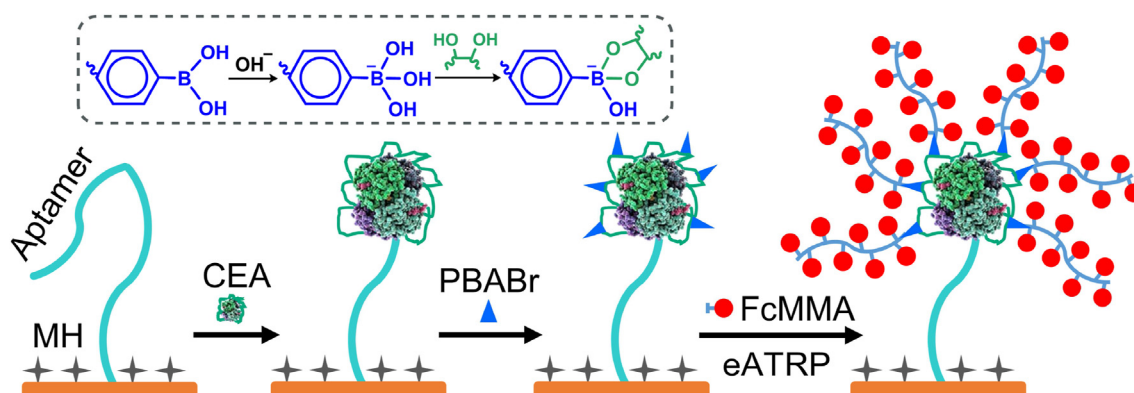
2.1. BA-eATRP-based electrochemical aptasensing of CEA

Prior to use, gold electrode (Au) was cleaned as described elsewhere [47,48]. After incubating the Au with 10 μL of 0.5 $\mu\text{mol}\cdot\text{L}^{-1}$ aptamer (Apt, 10 $\text{mmol}\cdot\text{L}^{-1}$ NaH₂PO₄/Na₂HPO₄ (PBS, pH7.4)) at 37 °C for 30 min, the Apt/Au was rinsed with ultrapure H₂O, incubated with 6-mercapto-1-hexanol (MH, 1.0 $\text{mmol}\cdot\text{L}^{-1}$ in 60% ethanol) at 37 °C for 15 min, and rinsed with ethanol and ultrapure H₂O. After incubating the MH/Apt/Au with 10 μL of CEA (10 $\text{mmol}\cdot\text{L}^{-1}$ PBS, pH7.4) at 37 °C for 45 min, the CEA/MH/Apt/Au was rinsed with ultrapure H₂O, incubated with (4-(2-bromo-2-phenylacetylaminophenyl)boronic acid (PBABr, 0.5 $\text{mmol}\cdot\text{L}^{-1}$ in 10 $\text{mmol}\cdot\text{L}^{-1}$ PBS containing 10% DMF (pH9.5)) at 37 °C for 15 min, and rinsed with ethanol and ultrapure H₂O. The PBABr/CEA/MH/Apt/Au was soaked into the polymerization solution (obtained by adding 0.1 mL of FcMMA (10 $\text{mmol}\cdot\text{L}^{-1}$ in DMF), 0.1 mL of Cu^{II}Br/Me₆TREN (10 $\text{mmol}\cdot\text{L}^{-1}$ in H₂O), 7.5 mL of KPF₆ (0.1 $\text{mol}\cdot\text{L}^{-1}$), and 1.0 mL of KBr (1.0 $\text{mol}\cdot\text{L}^{-1}$) into 1.3 mL of DMF), followed by the eATRP of FcMMA at room temperature (ca. 25 °C) for 40 min using *i-t* curve. After removal of the possible Cu(0) residues via the anodic oxidation at 0.19 V (*vs.* saturated calomel electrode (SCE)) for 30 s, the Fc/PBABr/CEA/MH/Apt/Au was tested by square wave voltammetry (SWV; potential increase, 4.0 mV; potential amplitude, 25 mV; frequency, 15 Hz) in 0.5 $\text{mol}\cdot\text{L}^{-1}$ LiClO₄ [47,48].

3. Results and discussion

3.1. Principle of the BA-eATRP-based electrochemical CEA aptasensor

The principle of the electrochemical CEA aptasensor is shown in Scheme 1. The aptamer was used as the recognition element and it was tethered via the Au-S chemistry. As CEA is a glycoprotein



Scheme 1. Principle of the BA-eATRP-based electrochemical CEA aptasensor.

[49], the ATRP initiators for the later polymerization were covalently crosslinked to the aptamer-captured antigens via the facile, selective interactions between the *cis*-diol groups of the monosaccharide residues and the PBA group [50–52]. Afterward, the eATRP of FcMMA resulted in the surface-initiated growth of the Fc-conjugated polymer chains and consequently the collection of Fc reporters [48]. As each CEA is decorated with hundreds of *cis*-diol groups for the BA-based crosslinking of ATRP initiators while each long-chain polymer can be composed of a plenty of Fc reporters, the BA-eATRP enabled the dual-amplification detection. Thus, the BA-eATRP-based electrochemical CEA aptasensor is ultrasensitive and highly selective.

3.2. Effect of eATRP potential

The surface-initiated growth of the Fc-conjugated polymer chains via the eATRP of FcMMA was initiated and sustained through the electrochemical reduction of the Cu(II) deactivators. With this in mind, the redox properties of the Cu(II)

deactivator were investigated. As displayed in Fig. 1A, the cyclic voltammetric (CV) curve of the Cu(II) deactivator at the PBABr/CEA/MH/Apt/Au presented a reduction peak at -0.58 V, corresponding to the reduction of the Cu(II) deactivators into the Cu(I) activators [43,44]. For the $[\text{Cu}^{\text{II}}\text{Br}/\text{L}]/[\text{Cu}^{\text{I}}/\text{L}]$ redox couple ($\text{L} = \text{Me}_6\text{TREN}$), the Nernst equation can be expressed as $E_{\text{app}} = E^{\circ} + (RT/nF) \cdot \ln\{[\text{Cu}^{\text{II}}\text{Br}/\text{L}]/[\text{Cu}^{\text{I}}/\text{L}] \cdot [\text{Br}^-]\}$. It is obvious that the concentration of the Cu(I) activator is highly dependent on the eATRP potential (E_{app}) [43,44]. Specifically, a higher E_{app} would result in a lower concentration of the Cu(I) activator, while a more negative E_{app} would result in the generation of more Cu(I) activators [43]. As the initiating radicals (R^{\bullet}) were generated from the reduction of the ATRP initiators by the Cu(I) activators [43], the rate of the surface-initiated growth of polymer chains was dependent on the E_{app} [43,44]. With this in mind, the effect of eATRP potential on the detection signal was investigated. The peak current corresponding to the anodic oxidation of the Fc reporters, as displayed in Fig. 1B, exhibited the highest value at the eATRP potential of -0.54 V,

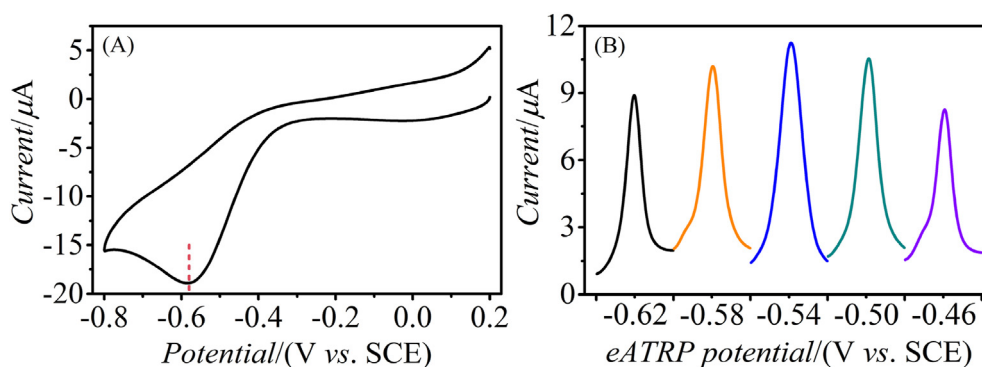


Fig. 1. (A) CV curve of the PBABr/CEA/MH/Apt/Au in the FcMMA-free polymerization solution (the dashed vertical line indicates the reduction peak). (B) Effect of eATRP potential on the detection signal. CEA, $1.0 \text{ ng} \cdot \text{mL}^{-1}$.

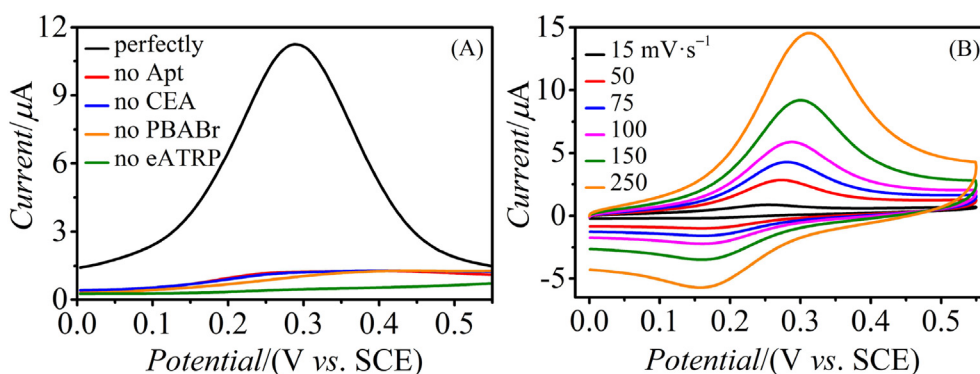


Fig. 2. (A) SWV curves of various electrodes. (B) CV curves of the Fc/PBABr/CEA/MH/Apt/Au at various scan rates. CEA, $1.0 \text{ ng}\cdot\text{mL}^{-1}$.

which was hence adopted in the subsequent experiments.

3.3. Characterizations of the electrochemical CEA aptasensor

The feasibility of the BA-eATRP-based electrochemical CEA aptasensor was investigated based on the SWV curves derived from various electrodes. As seen in Fig. 2A, the perfectly modified electrode (i.e., the Fc/PBABr/CEA/MH/Apt/Au) can be featured by a high oxidation peak at ca. 0.29 V, while no oxidation peak could be observed for the PBABr/CEA/MH/Apt/Au (i.e., without eATRP). This observation validated that the oxidation current observed at ca. 0.29 V was derived from the electrochemical oxidation of the Fc reporters. Likewise, almost no oxidation peak could be observed for the electrodes modified in the absences of aptamers, CEA antigens, or the PBABr agents. The high reliability of the detection signal validated the feasibility of the BA-eATRP-based electrochemical CEA aptasensor. CV of the Fc/PBABr/CEA/MH/Apt/Au was further applied

to investigate the redox behavior of the Fc reporters (Fig. 2B). As the Fc reporters were surface-tethered, the peak currents showed well-defined linear correlation with the scan rate (Figure S1), confirming the surface-confined charge-transfer process [47,48]. Fig. 3 clearly shows that there is a high density of polymer chains on the Fc/PBABr/CEA/MH/Apt/Au. Apart from the P elements observed for the DNA aptamer in the energy dispersive X-ray spectrum (EDS) mapping as shown in Fig. 4A, the EDS mappings of the B (Fig. 4B) and Br elements (Fig. 4C) proved the presence of PBABr, while that of the Fe element (Fig. 4D) unequivocally demonstrated the presence of the Fc reporters.

The preparation of the Fc/PBABr/CEA/MH/Apt/Au was monitored by electrochemical impedance spectroscopy (EIS, Fig. 5A) and CV (Fig. 5B). For the Au, the charge-transfer resistance (R_{ct}) was the smallest (Fig. 5A,a), while the redox currents were the highest (Fig. 5B,a). Owing to the steric hindrance of the aptamer layer, and the electrostatic repulsion between the phosphate groups and the $[\text{Fe}(\text{CN})_6]^{3-/4-}$ [44], the modification of aptamers (i.e., the Apt/Au) resulted in the increase of the R_{ct} (Fig. 5A and b) and the decrease of the redox currents (Fig. 5B,b). As the blocking of the residual sites would cause an increase in the steric hindrance [43,47], the MH/Apt/Au showed a higher R_{ct} (Fig. 5A,c) and the lower redox currents (Fig. 5B and c) relative to the Apt/Au. As displayed, the capture of CEA antigens by aptamers (i.e., the CEA/MH/Apt/Au) resulted in a slightly increased R_{ct} (Fig. 5A,d) and slightly decreased redox currents (Fig. 5B,d) [45,46]. As the affinity crosslinking between the *cis*-diol groups and the PBA groups resulted in the formation of the negatively-charged phenylborates [53,54], the PBABr/CEA/MH/Apt/Au featured a significantly increased R_{ct} (Fig. 5A,e) and significantly decreased redox currents (Fig. 5B,e) relative to the CEA/MH/Apt/Au [45,46]. As the densely packed Fc reporters could

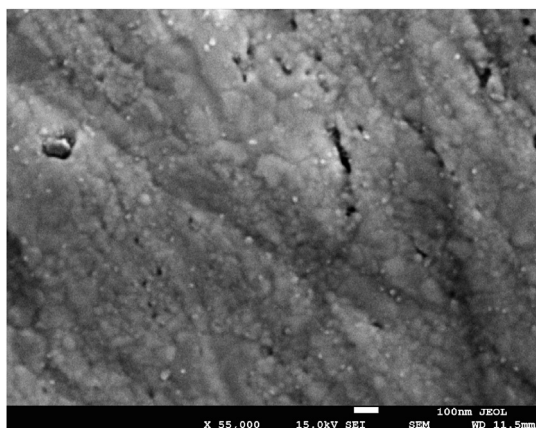


Fig. 3. SEM image of the Fc/PBABr/CEA/MH/Apt/Au. CEA, $1.0 \text{ ng}\cdot\text{mL}^{-1}$.

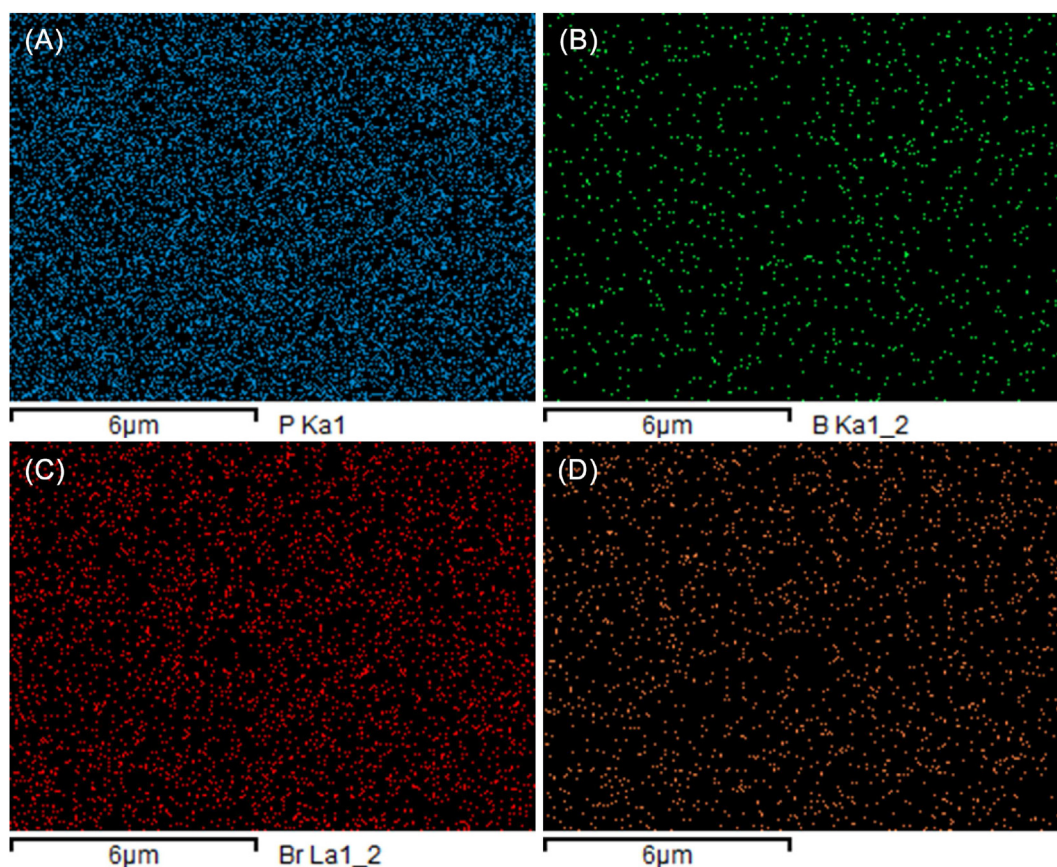


Fig. 4. EDS mappings of (A) P, (B) B, (C) Br, and (D) Fe elements. CEA, $1.0 \text{ ng}\cdot\text{mL}^{-1}$.

significantly facilitate the charge transfer between the electrode and the $[\text{Fe}(\text{CN})_6]^{3-/4-}$ [38,44,47], the Fc/PBABr/CEA/MH/Apt/Au showed a significantly decreased R_{ct} (Fig. 5A,f) and the significantly increased redox currents (Fig. 5B,f) when compared with the PBABr/CEA/MH/Apt/Au. As the increase of the R_{ct} was accompanied by the decrease of the redox currents, while the decrease of the R_{ct} was accompanied by the increase of the redox currents [48], the EIS results agreed well with the CV results [45,46], which validated the

successful preparation of the Fc/PBABr/CEA/MH/Apt/Au.

3.4. Analytical performance of the electrochemical CEA aptasensor

Under the optimal conditions, the analytical performance of the BA-eATRP-based electrochemical CEA aptasensor was investigated. From Fig. 6, it can be seen that the peak current increased with the increase of CEA concentration, indicating

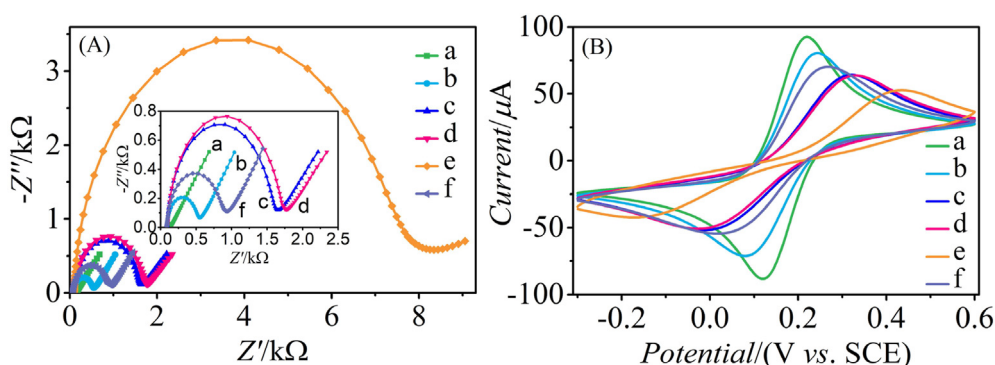


Fig. 5. (A) Nyquist plots and (B) CV curves of the Au (a), Apt/Au (b), MH/Apt/Au (c), CEA/MH/Apt/Au (d), PBABr/CEA/MH/Apt/Au (e), and Fc/PBABr/CEA/MH/Apt/Au (f). CEA, $1.0 \text{ ng}\cdot\text{mL}^{-1}$.

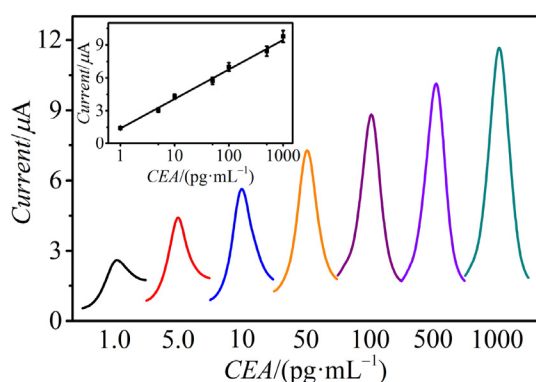


Fig. 6. SWV curves toward various CEA concentrations and the calibration plot ($n = 5$, the inset).

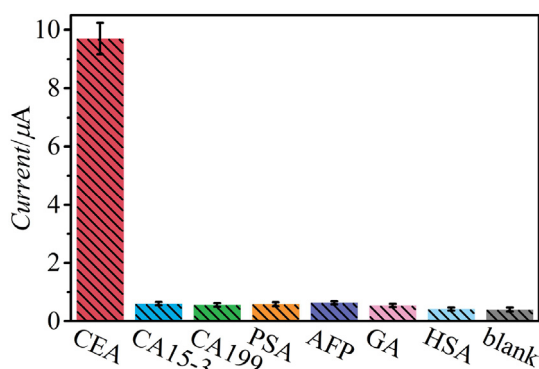


Fig. 7. Peak currents toward $1.0 \text{ ng} \cdot \text{mL}^{-1}$ CEA or $10 \text{ ng} \cdot \text{mL}^{-1}$ interferences ($n = 5$).

the signal-on detection. The inset shows that there was a good linearity between the peak current and the logarithm of CEA concentration over the range from 1.0 to $1000 \text{ pg} \cdot \text{mL}^{-1}$ ($R^2 = 0.995$). The regression equation of the calibration plot can be expressed as $y = 1.395 + 2.688 \lg x$, in which y represents the peak current in μA and x represents the concentration of CEA in $\text{pg} \cdot \text{mL}^{-1}$. The detection limit was calculated to be $0.34 \text{ pg} \cdot \text{mL}^{-1}$, which was satisfactory when compared with some of the previous methods (Table S1). The high detection sensitivity was a result of the dual amplification by the BA-eATRP-based strategy. The relative standard deviation (RSD) of the peak current toward the intraassay of $1.0 \text{ ng} \cdot \text{mL}^{-1}$ CEA was 5.3% ($n = 5$) while for the interassay it was 5.7% ($n = 5$), clearly showing that the reproducibility of the electrochemical CEA aptasensor was acceptable. As a proof of concept, the storage of the Fc/PBABr/CEA/MH/Apt/Au at 4°C for two weeks resulted in the

loss of the peak current by about 5.4% , showing the high storage stability of the perfectly modified electrode.

3.5. Selectivity of the electrochemical CEA aptasensor

The selectivity of the BA-eATRP-based electrochemical CEA aptasensor was investigated based on the detection of $10 \text{ ng} \cdot \text{mL}^{-1}$ CA15-3, CA199, prostate-specific antigen (PSA), alpha-fetoprotein (AFP), glycated albumin (GA), and human serum albumin (HSA), and $1.0 \text{ ng} \cdot \text{mL}^{-1}$ CEA. From Fig. 7, it can be seen that the presence of $1.0 \text{ ng} \cdot \text{mL}^{-1}$ CEA could result in the generation of a high peak current, while the presence of $10 \text{ ng} \cdot \text{mL}^{-1}$ other component resulted in no significant difference in the intensity of the peak current relative to the blank sample. The results indicated that the BA-eATRP-based electrochemical CEA aptasensor is highly selective. The high selectivity was a result of the aptamer-based recognition [45,46,55–57].

3.6. Analysis of CEA in human serum

The analysis of CEA in human serum was applied to investigate the potential practicability of the BA-eATRP-based electrochemical aptasensor. Due to the lack of clinical specimens, the proof-of-concept experiments were conducted using the commercially supplied NHS. The NHS was filtered using a Millipore Amicon® Ultra-4 centrifugal filter unit (cutoff, 100 kDa) to remove the endogenous CEA antigens, and then diluted by 20 times using $10 \text{ mmol} \cdot \text{L}^{-1}$ PBS (pH 7.4). As seen in Table 1, the recoveries were within the range from 94.3% to 105.5% , while the RSDs were within the range from 5.2% to 5.8% . The results of the standard addition experiments indicated that the BA-eATRP-based electrochemical aptasensor is applicable to the quantitative analysis of CEA in serum samples.

4. Conclusions

In summary, a cost-effective and easy-to-operate electrochemical aptasensor has been reported for the ultrasensitive and highly selective detection of CEA, taking advantage of the dual amplification by the BA-eATRP. The covalent crosslinking between the PBA group and the *cis*-diol groups of the monosaccharide residues allows for the site-specific labeling of each CEA antigen with hundreds of ATRP initiators, while the surface-initiated growth of long-chain polymers through eATRP results in the tethering of each ATRP initiator-conjugated site with hundreds to thousands of Fc

Table 1. Analysis of spiked CEA in human serum.

Sample	Added ($\text{pg} \cdot \text{mL}^{-1}$)	Found ($\text{pg} \cdot \text{mL}^{-1}$)	Recovery	RSD ($n = 5$)
1	1000	943.0	94.3%	5.2%
2	100	103.5	103.5%	5.4%
3	10	10.55	105.5%	5.8%

reporters. Thus, the BA-eATRP enables the labeling of each CEA antigen with a plenty of Fc reporters, leading to the dual-amplification detection. The BA-eATRP-based amplification strategy is free from the use of the sophisticated enzymatic reactions or nanocomposites, offering the benefits of cost-effectiveness and simple operation to the electrochemical aptasensing of CEA. Taking together, the BA-eATRP-based electrochemical CEA aptasensor holds great promise in clinical applications.

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基于硼酸盐亲和辅助电化学调控 ATRP 的癌胚抗原超灵敏电化学适体传感研究

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摘要

癌胚抗原 (CEA) 是一种酸性糖蛋白, 其作为一种广谱肿瘤标志物在恶性肿瘤的鉴别诊断与监测等方面具有重要价值。在此, 借助于硼酸盐亲和辅助电化学调控原子转移自由基聚合 (BA-eATRP) 的双重信号放大作用, 我们报道了一种电化学适体传感器, 用于 CEA 的超灵敏、高选择性检测。基于 BA-eATRP 的电化学 CEA 适体传感的基本原理为: 待核酸适体捕获 CEA 抗原后, 借助于苯硼酸 (PBA) 基团与单糖残基上的顺式二醇基团间的选择性亲和相互作用将 ATRP 引发剂位点靶向性地共价偶联到 CEA 抗原上; 随后, 以二茂铁甲基丙烯酸甲酯 (FcMMA) 作为单体, 借助于 eATRP 将二茂铁 (Fc) 探针引入电极表面。由于 CEA 上含有数百个顺式二醇基团, 基于硼酸盐亲和的交联反应可使得在每个 CEA 抗原上标记数百个 ATRP 引发剂分子。此外, 通过 eATRP 反应, 可以在电极表面接枝长的二茂铁基聚合物链, 使得每个标记有 ATRP 引发剂的位点均能连接上成百上千个 Fc 探针。因此, BA-eATRP 可使得每个 CEA 抗原上标记上大量的 Fc 探针。在最佳条件下, 基于 BA-eATRP 的电化学适体传感器能够实现浓度低至 $0.34 \text{ pg} \cdot \text{mL}^{-1}$ 的 CEA 的高选择性检测, 其线性范围为 $1.0\text{--}1000 \text{ pg} \cdot \text{mL}^{-1}$ 。而且, 该适体传感器可用于人血清中 CEA 的定量分析。基于 BA-eATRP 的电化学适体传感器具有成本低廉、操作简便等优良特性, 在 CEA 的超灵敏、高选择性检测方面具有广阔的应用前景。

关键词: 硼酸盐亲和; 原子转移自由基聚合; 电化学适体传感器; 癌胚抗原; 肿瘤标志物; 信号放大