Needle-shaped electrode for speciation analysis of copper in seawater

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ABSTRACT

Here, a needle-shaped electrode based on commercial acupuncture needle was fabricated and functionalized with dendritic gold nanostructures (AuNDs) and ion-exchange polymer (Nafion) for voltammetric speciation analysis of copper (Cu) in seawater. To achieve the enhanced performance for voltammetric determination of Cu in seawater, the acupuncture needle electrode (ANE) was functionalized with AuNDs via electrochemical deposition and covered with Nafion. AuNDs facilitate the electrochemical responses of Cu due to their excellent catalytic and electroconductive properties, whereas Nafion can stabilize the AuNDs deposited on the surface of the needle tip and enhance the voltammetric responses through its specific cation-exchange ability with Cu2+. The so-designed AuNDs/Nafion/ANE showed excellent performance for the determination of Cu in complex matrix like seawater. Under the optimal conditions, the response of Cu at this functional needle-shaped electrode was linear in the concentration range from 0.05 to 1 nmol L−1 with an ultralow detection limit of 15.4 pmol L−1. Additionally, the functional needle-shaped electrode has been successfully used for the direct determination of different Cu species in seawater via anodic stripping voltammetry.

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1. Introduction

In the past decades, global economic and industrial developments have resulted in a large number of pollutants to be dumped into the marine environment, which led to an increased concern for both public and environmental health [1]. The marine ecosystems are becoming increasingly degraded due to the anthropogenic inputs of heavy metals, such as copper (Cu), cadmium (Cd), and lead (Pb) [2,3]. Although Cu is a trace element indispensable for maintaining seawater biological activities, an excess amount of bioavailable Cu is toxic [4,5]. Indeed, Cu not only has a high impact in the heavy-metal toxicity issues in the marine organisms, but also threatens human health by bioaccumulation [6].

Therefore, the effect of Cu in the marine environment has received considerable attention because of its dangerous consequences on the ecosystem and human health. Cu also plays a crucial role in promoting plankton growth and regulating biogeochemical processes [7]. However, the total concentration of Cu is not a parameter sufficient to efficiently assess its bioavailability, behavior, and toxicity [8]; indeed, the relevance of Cu for the environment does not depend on its concentration but on its aqueous speciation. Therefore, to further study the ecological function, biological impact, and toxic metabolism of Cu, it is essential to determine the occurrence and relative abundance of different Cu species in seawater. So speciation-selective analytical methods should be developed to achieve the speciation analysis of Cu.

Several methods have been used to conduct the speciation analysis of Cu and other metal elements in seawater. A conventional approach for determining the species of metals like Cu combines the separation techniques (including gas chromatography [9], supercritical-fluid chromatography [10], and capillary zone electrophoresis [11]) with detection methods (including atomic absorption spectrometry [12], atomic emission spectrometry [13],...
atomic fluorescence spectrometry [14], and inductively coupled plasma mass spectrometry [15]). Although these methods have many advantages, a significant limitation is their requirement for a series of elaborate pre-treatments, which are time and work expensive. In other words, there is an urgent need to develop more effective and simple methods to conduct the speciation analysis of Cu and other metals.

Anodic stripping voltammetry (ASV) is an efficient and simple electroanalytical technique for speciation analysis due to its high selectivity and sensitivity, excellent performance with saline matrices, such as seawater, as well as the rapid and straightforward analytical processes required [16]. Annibaldi et al. surveyed the total and dissolved Cu, Pb, Cd in the estuary and coastal systems of Italy via square wave anodic stripping voltammetry (SWASV) [17,18]. The mercury electrode has frequently been used as the working electrode for the speciation analysis of Cu and other metals due to its high sensitivity and stability [17–20]. However, the mercury electrode has significant drawbacks including toxicity and difficulties in handling process [21]. Although several chemically-modified electrodes have been developed to replace the mercury electrode, most of them cannot be applied to the determination of dissolved active Cu, one of the Cu species, because of its relatively low in seawater with seawater [6]. Therefore, there is a compelling need to fabricate an ultrasensitive electrode for the direct determination of Cu species in seawater.

Acupuncture needle is a new emerging substrate material for constructing highly sensitive electrodes. Acupuncture is a traditional Chinese technique for relieving pain and performing disease therapies by inserting and manipulating acupuncture needles into the body acupuncture points (or acupoints) [22]. Among multiple acupuncture needles, the stainless steel needle is quite cheap, easy to handle [23]. It has been reported that it is possible to manufacture the acupuncture needle as a needle-shaped electrode for electrochemical and bio-sensing applications [24,25]. Because of the low price, the needle-shaped acupuncture needle electrode (ANE) can be used disposable with no extra polishing process before electrochemical experiments. More importantly, the unique needle-shaped structure provides larger specific surface area for the modification of functional nanomaterials. Thanks to the unique shape and size, the ANE modified with functional nanomaterials can exhibit much superior performance in sensitivity and detection limit than the conventional electrodes. Gold nanomaterials, especially those with dendritic structures, possess plenty of interesting electrochemical properties, such as large surface-to-volume ratio, high conductivity, and excellent electrocatalytic performance, which make them optimal candidates for electrode modification [26]. Moreover, gold nanomaterials are provided with an outstanding catalytic performance for the electrochemical detection of Cu$^{2+}$ and other targets [6,27,28]. Nafion, one of the most widely used cation-exchange polymers, can form a thin film on the surface of nanomaterials and stabilize them due to its high chemical and mechanical stability. Furthermore, Nafion has the remarkable effect in pre-concentrating cation targets based on its excellent ion-exchange capability, which make it widely used in the electrochemical analytical field [29].

This work combined the unique needle-shaped structure and micro-size of ANE with the excellent electrochemical properties of dendritic gold nanostructures (AuNDs) and specific pre-concentration ability of Nafion to fabricate a functional needle-shaped electrode with enhanced performance for the analysis of Cu in seawater. Indeed, an AuNDs/Nafion functionalized ANE (AuNDs/Nafion/ANE) with remarkably improved sensitivity for the voltammetric determination of Cu$^{2+}$ have been designed. The schematic of determination of Cu$^{2+}$ on the so-designed functional needle-shaped electrode can be explained as the adsorption-catalysis model. The experimental conditions and analytical performance of the AuNDs/Nafion/ANE have been systematically assessed. To achieve the speciation analysis of Cu, the dissolved active, dissolved acid extractable, and total acid extractable Cu in seawater were all successfully determined with the functional needle-shaped electrode via anodic stripping voltammetry with excellent results.

2. Experimental

2.1. Reagents and materials

All the reagents were analytical-grade chemicals and used without further purification. The stock standard solution of 1000 mg L$^{-1}$ Cu was supplied by the National Research Centre for Certified Reference Materials (CRMs), Beijing, China. Separate standard solutions were prepared by dilution of the stock solution with 10% (v/v) HNO$_3$ to the specific concentration required. HAuCl$_4$·4H$_2$O was purchased from Sinopharm Chemical Reagent Co. Ltd., and 2 mmol L$^{-1}$ HCl solution was prepared in 0.5 mol L$^{-1}$ H$_2$SO$_4$ for electrodoposition. Nafion (5 wt% in a mixture of lower aliphatic alcohols and water, contains 45% water) was purchased from Sigma-Aldrich Co. Ltd. Acetate buffer solution (pH 4.5) was prepared by 1 mol L$^{-1}$ acetic acid and 0.1 mol L$^{-1}$ sodium acetate. Deionized water (18.2 MΩ cm specific resistance) obtained with a Pall Cascada laboratory water system was used throughout.

Stainless steel acupuncture needles (Size 0.25 × 60 mm, Lot No. 1600082) were purchased from Suzhou Medical Appliance Factory Co. Ltd. (Suzhou, China). PVC tubes were obtained by removing the internal copper from the copper electric wires. Silicone rubber (NANDA no. 704) was purchased from Liyang Kangda Chemical Co. Ltd., China.

2.2. Apparatus

The morphologies of the bare and modified ANEs were characterized via scanning electron microscopy (SEM, Hitachi S-4800 microscope, Japan). Electrochemical Work Station (CHI 660E, Shanghai CHI Instruments, China) was used throughout all the electrochemical experiments. The modified ANE served as the working electrode, with Ag/AgCl (3.0 mol L$^{-1}$ KCl) and platinum foil serving as the reference and counter electrodes, respectively. All potentials were measured with respect to the Ag/AgCl reference electrode.

2.3. Preparation of the AuNDs/Nafion/ANE

The procedure for the fabrication of the AuNDs/Nafion/ANE was illustrated in Scheme 1. Firstly, PVC tube was cut to a proper length to insulate the acupuncture needle. The acupuncture needle was, thus, inserted into the PVC tube with the needle tip outside about 3 mm for sensing surface and needle handle outside for connecting with Electrochemical Work Station. Then, the two openings of the PVC tube were sealed with silicone rubber, and the sensing tip of the needle was controlled about 2 mm length. After the silicone rubber was solidification at room temperature, the ANE was fabricated.

Before modification, the ANE was treated with an ultrasonic cleaning process by using water and ethanol for 5 min. The formation of AuNDs on ANE surface was achieved by electrodeposition in 2 mmol L$^{-1}$ HAuCl$_4$ solution at the deposition potential of −0.3 V for 120 s. Then, the sensing tip of the AuNDs modified ANE (AuNDs/ANE) was immersed in 2.5 wt% Nafion solution for 1–3 s to cover the Nafion film on the surface of AuNDs. It was found that the immersing time had little effect on the electrochemical response of Cu if the concentration of Nafion was constant (Fig. S1). Eventually, the AuNDs/Nafion/ANE was obtained after the electrode was dried
Seawater samples were collected from three different locations across the Sishili Bay, Northern Yellow Sea, China in November 2017. A 2.5 L sample collector was used to collect seawater into polypropylene containers, which were washed with a solution of 1 mol L\(^{-1}\) HCl and rinsed with deionized water. All the apparatus used were also appropriately cleaned with HNO\(_3\), and then thoroughly washed using deionized water before starting the experiments. Seawater samples were immediately processed after collection, as summarized in Fig. S2. Each sample was divided into three aliquots; the first was filtered (0.45 μm membrane filters) and kept to determine the total acid extractable Cu concentration. The second aliquot was filtered, acidified with ultrapure HCl (pH 1.8) for 48 h, and kept in a refrigerator at 4 °C to determine the dissolved active Cu concentration. The last aliquot was acidified for 48 h, filtered, and kept to determine the total acid extractable Cu concentration. The dissolved active, dissolved acid extractable, and total acid extractable Cu concentrations were determined directly with the AuNDs/Na\(_m\) electrodes (Fig. 2A). No redox peak could be observed in the voltammograms of these three electrodes, as well as the combined effects of AuNDs and Na\(_m\), the typical reduction peak of Na\(_m\) on the surface of AuNDs/ANE, the decrease of the typical peak current and background current might be ascribed to the improved detection performance for Cu\(^{2+}\).

The electrochemical properties of the AuNDs/Na\(_m\)/ANE were further investigated via cyclic voltammetry. The cyclic voltammograms of the bare ANE, AuNDs/ANE, and AuNDs/Na\(_m\)/ANE in 0.1 mol L\(^{-1}\) acetate buffer solutions from 0.2 to 1.3 V have been recorded to characterize the gold nanomaterials modified on the electrodes (Fig. 2A). No redox peak could be observed in the voltammogram of bare ANE, whereas a sharp reduction peak at 0.62 V and an oxidation peak at 1.0 V caused by the redox of AuNDs can be observed for AuNDs/ANE. Interestingly, after the modification of Na\(_m\) on the surface of AuNDs/ANE, the typical reduction peak of AuNDs becomes smaller. The cyclic voltammograms of these three electrodes in acetate buffer from −0.2 to 0.6 V have also been conducted (Fig. 2B). It can be seen that the background current of AuNDs/ANE is considerably larger than that of the bare ANE. However, similar to typical peak of AuNDs, the background current of AuNDs/Na\(_m\)/ANE is smaller than that of AuNDs/ANE. It can be concluded that the background current enhancement of AuNDs/ANE is due to the excellent ability of Na\(_m\) in facilitating electron transfer. For the AuNDs/Na\(_m\)/ANE, the decrease of the typical peak current and background current might be ascribed to the relatively weak electron transfer capability (electro-conductivity) of Na\(_m\).

To verify the capability of electron transfer of different electrodes, as well as the combined effects of AuNDs and Na\(_m\), electrochemical impedance spectrum (EIS) was conducted. The semicircle at the high frequency of EIS indicates an electron-transfer-limited process on the electroactive surface. So the...
Fig. 1. SEM images of the so-fabricated ANE (A, B), AuNDs/ANE (C), and AuNDs/Nafion/ANE (D).

Fig. 2. Cyclic voltammograms of the bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE in 0.1 mol L\(^{-1}\) acetate buffer (pH 4.5) solution from 0.2 to 1.3 V (A), −0.2 to 0.6 V (B) with the scan rate 50 mV s\(^{-1}\). Nyquist plots of the bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE in 0.1 mol L\(^{-1}\) KCl solution containing 10 mmol L\(^{-1}\) K\(_3\)[Fe(CN)\(_6\)] (C). Cyclic voltammograms of the bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE in 0.1 mol L\(^{-1}\) KCl solution containing 10 mmol L\(^{-1}\) K\(_3\)[Fe(CN)\(_6\)] from −0.2 to 0.6 V with a scan rate of 25 mV s\(^{-1}\) (D).
electron transfer resistance (Ret) of the electroactive surface can be measured by the semicircle diameter at high frequency. As shown in Fig. 2C, the Nyquist diagrams of bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE were measured in 10 mmol L\(^{-1}\) \(K_3[Fe(CN)]_6\) solution with 0.1 mol L\(^{-1}\) KCl. By comparing the semicircle diameters of different electrodes, it can be observed that the Ret of AuNDs/ANE is significantly smaller than that of bare ANE. This phenomenon is due to the excellent ability of AuNDs in facilitating electron transfer which promotes the \([Fe(CN)]_6^{3-}/[Fe(CN)]_6^{4-}\) redox process. However, the Ret becomes larger after the modification of Nafion on the surface of AuNDs/ANE, which indicates the relatively weak electro-conductivity of Nafion and the inhibition effect between negatively-charged Nafion and \([Fe(CN)]_6^{3-}/[Fe(CN)]_6^{4-}\) probe. These results demonstrate that the AuNDs/Nafion/ANE could be successfully fabricated, and AuNDs have the excellent ability in facilitating electron transfer, while Nafion has relatively weak electro-conductivity. It should be noted that despite its relatively weak electro-conductivity, Nafion can facilitate the adsorption and pre-concentration of Cu\(^{2+}\) due to its outstanding cation-exchange ability.

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\left[Fe(CN)]_6^{3-}-[Fe(CN)]_6^{4-}\text{ redox couple has also been commonly used as the electrochemical probe to calculate the electroactive surface area and study the properties of modified electrodes. Fig. 2D shows the cyclic voltammograms of the bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE using } [FeCN]_6^{3-}/[FeCN]_6^{4-}	ext{ redox couple as an electrochemical probe. Almost no redox peak can be observed in the voltammogram of bare ANE, whereas the AuNDs/ANE shows the highest current response. The visible symmetrical redox reaction of } K_3[Fe(CN)]_6\text{ on the AuNDs/ANE was proven as a reversible process. The relatively weak electro-conductivity of Nafion might be presented as an obvious explanation for the current decrease of AuNDs/Nafion/ANE. Moreover, Nafion is negatively charged and could hinder the electron transfer between the probe and electrode, which would also decreases the current response. Considering that no other free ions were added in the [FeCN]_6^{3-}/[FeCN]_6^{4-} solution, the difference between reduction and oxidation peak on intensity and shape might be resulted from the hindrance role of Nafion. The electroactive surface areas of the modified electrodes were estimated according to the Randles-Sevcik equation:
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Ip = 2.69 \times 10^5 n^{3/2} A D^{1/2} v^{1/2} c_0 \quad (1)
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where \(Ip\) is the anodic peak current, \(n\) is the electron transfer number, \(A\) is the electroactive surface area of the electrode, \(D\) is the diffusion coefficient, \(v\) is the scan rate, and \(c_0\) is the concentration of \(K_3[Fe(CN)]_6\). For 10 mmol L\(^{-1}\) \(K_3[Fe(CN)]_6\) in 0.1 mol L\(^{-1}\) KCl electrolyte, \(n\) is 1 and \(D\) is \(7.6 \times 10^{-6}\) cm\(^2\) s\(^{-1}\). The active surface areas could be calculated from the slope of the \(Ip\) versus \(v^{1/2}\), which was shown in Fig. 3. Five different scan rates (10, 25, 50, 75, and 100 mV s\(^{-1}\)) were used to set up the calibration curves for determination of electroactive surface areas. Thus, the electroactive surface areas of bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE were \(2.35 \times 10^{-6}\), \(3.36 \times 10^{-3}\) and \(1.93 \times 10^{-3}\) cm\(^2\), respectively. The results show that the presence of AuNDs can increase the electroactive surface significantly compared with the bare ANE. Although the existence of Nafion decreases the electroactive surface area due to its relatively weak electro-conductivity and inhibition effect, it can enhance the electrochemical response of Cu\(^{2+}\) due to its excellent ability in adsorption and pre-concentration.

### 3.2. Mechanism of Cu\(^{2+}\) determination on the AuNDs/Nafion/ANE

SWASV responses of bare ANE, AuNDs/ANE and AuNDs/Nafion/ANE to 50 nmol L\(^{-1}\) Cu\(^{2+}\) in 0.1 mol L\(^{-1}\) acetate buffer supporting electrolyte were shown in Fig. 3. No anodic signal can be detected for the bare ANE with 50 nmol L\(^{-1}\) Cu\(^{2+}\), while an oxidation peak at about 0.3 V can be observed for the AuNDs/ANE due to the excellent electrochemical performance of AuNDs. As expected, the current response of AuNDs/Nafion/ANE is much larger than that of AuNDs/ANE. There is no doubt that AuNDs/Nafion/ANE shows the best performance for Cu\(^{2+}\) detection after the combination of AuNDs and Nafion. To explain the superior performance of AuNDs/Nafion/ANE, mechanism of Cu\(^{2+}\) determination on the electrode has been investigated.

For anodic stripping voltammetric analysis, it is very important for the effective adsorption of target on the electrode surface. At this stage, a large amount of Cu\(^{2+}\) will be pre-concentrated on the surface of Nafion due to its outstanding adsorption ability through ion-exchange reaction. And after the adsorption of Cu\(^{2+}\), the AuNDs on the surface of ANE will serve as the electrochemical reaction interfaces. On the interfaces, the reactions of Cu\(^{2+}\) reduced to Cu\(^0\) during the accumulation process, and Cu\(^0\) oxidised to Cu\(^{2+}\) during the stripping process will take place. It is well known that the AuNDs have superior properties in facilitating electron transfer and excellent catalytic performance for electrochemical detection. Despite Nafion could decrease the current response due to its relatively weak electro-conductivity, it enhances the detection performance by facilitating the cation-exchange reaction for the pre-concentration of Cu\(^{2+}\). Therefore, in the competition between the two effects provided by Nafion, the pre-concentration effect dominated in the voltammetric detection of Cu\(^{2+}\). So the improved electrochemical response of the AuNDs/Nafion/ANE for Cu\(^{2+}\) could be explained as a combined effects of the adsorption ability of Nafion and excellent electrocatalytic ability of AuNDs [30,31]. The schematic of voltammetric detection of Cu\(^{2+}\) on the AuNDs/Nafion/ANE based on the adsorption-catalysis model was shown in Scheme 2.

### 3.3. Optimization for the response of Cu on the AuNDs/Nafion/ANE

The deposition time of the AuNDs at ANE surface was investigated ranging from 30 to 210 s. As shown in Fig. 4A, the anodic peak current (Ip) gradually increases from 30 to 120 s and, then, remains almost constant from 120 to 210 s. The maximum Ip is at the deposition time from 120 to 210 s. It has been illustrated that AuNDs increase the electroactive surface area significantly and show excellent catalytic performance for the electrochemical

![Fig. 3. Square wave voltammograms of the bare ANE, AuNDs/ANE, and AuNDs/Nafion/ANE in 0.1 mol L\(^{-1}\) acetate buffer (pH 4.5) solution containing 50 nmol L\(^{-1}\) Cu\(^{2+}\).](image-url)
At the first 120 s of the deposition time, more AuNDs will be deposited on the surface of ANE and provide more electroactive sites for Cu$^{2+}$ detection. As a result, the Ip gradually increases with the deposition time at this stage. However, when the deposition time is more than 120s, the electroactive surface area will not increase continually due to the aggregation of AuNDs, and sequentially no enhancement of Ip obtained. Therefore, 120 s was selected as the optimal time for the deposition of AuNDs.

The effect of the amount of Na$^{+}$ on the AuNDs/ANE surface could be investigated at the concentrations ranging from 0.1% to 5% (Fig. 4B). The anodic Ip gradually increases from 0.1% to 5% of the Nafion solution. It should be noted that excess Nafion could hinder the electron transfer between the analyte and modified electrode. However, the decrease of Ip with high concentrations of Nafion has not been observed. The reason might be that only a small quantity of Nafion has been covered on the AuNDs/ANE considering the modification has been conducted via an immersion method. In this work, 2.5% of the Nafion solution was adopted which was enough for the determination of the dissolved active Cu.

The effect of the accumulation potential on the current response was also investigated with the potential varying from −0.7 to 0.1 V. As shown in Fig. 4C, the maximum Ip can be observed at the

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**Scheme 2.** Schematic of voltammetric determination of Cu$^{2+}$ on AuNDs/Nafion/ANE based on the adsorption of Nafion and the fast electron transfer of AuNDs.
potential of −0.3 V. The Ip gradually increases with decreasing potential from 0.1 to −0.3 V due to the full reduction of Cu²⁺ by extremely negative potentials. However, the Ip decreases from −0.3 to −0.7 V, which is probably caused by the competitive reduction of oxygen. Obviously, the optimum accumulation potential was chosen to be −0.3 V, and this value has been used in the following measurements.

The influence of the accumulation time on the stripping response was studied in the range from 30 to 330 s by fixing the deposition potential at −0.3 V. As shown in Fig. 4D, the Ip increases gradually as the accumulation time increases to 240 s and then tends to be steady because of the adsorption equilibrium of Cu²⁺ between the electrode surface and the solution. An accumulation time of 120 s is enough for carrying out the speciation analysis of Cu. However, higher sensitivity could be obtained by applying an accumulation time longer than 120 s.

### 3.4. Analytical performance of the AuNDs/NaFion/ANE for Cu

Fig. 5 shows the calibration curve and voltammograms obtained at the AuNDs/NaFion/ANE under the optimal conditions, which are 120 s of the deposition time, 2.5% of the NaFion concentration, −0.3 V of the accumulation potential, and 120 s of the accumulation time. The linear regression equation for Cu²⁺ determination derived from the SWASV curves is \( Ip = 18.98C + 4.47 \) (\( R^2 = 0.997 \)). The Ip is linear when the concentrations of Cu²⁺ range from 0.05 to 1 nmol L⁻¹. The sensitivity and detection limit of the AuNDs/NaFion/ANE for Cu²⁺ determination were 18.98 ± 0.89 nA/nmol L⁻¹ and 15.4 ± 0.7 pmol L⁻¹ (s/n = 3), respectively. Additionally, comparison of the proposed method with other analytical approaches for determination of Cu²⁺ is reported in Table 1. Compared with other electrodes, the AuNDs/NaFion/ANE provided here has the excellent performance for Cu²⁺ detection, which is essential to the direct analysis of dissolved active Cu in seawater. It can be concluded that the AuNDs/NaFion/ANE would be a promising choice for Cu²⁺ determination in the real seawater samples with higher sensitivity.

### 3.5. Reproducibility, repeatability and selectivity

The repeatability and reproducibility of the newly developed functional needle-shaped electrode were also assessed. The repeatability was evaluated by detecting 50 nmol L⁻¹ Cu²⁺ at the same electrode for ten measurements and the estimated relative standard deviation (RSD) was 2.9%. The reproducibility of the AuNDs/NaFion/ANE was investigated by six independently modified electrodes prepared in the same method in 50 nmol L⁻¹ Cu²⁺ and the RSD was 4.9%. Hence, the AuNDs/NaFion/ANE shows good reproducibility and repeatability for the determination of Cu²⁺. The modified electrode was stored in air condition and tested by determining Cu²⁺ for five measurements in each day, the current response remained 95.2% of the initial value after six days, reflecting the excellent stability of the AuNDs/NaFion/ANE.

Additionally, the effects of possible interfering species for Cu²⁺ determination were investigated by adding foreign species into the 0.1 mol L⁻¹ acetate buffer containing 50 nmol L⁻¹ Cu²⁺ under the optimal working conditions. Square wave voltammograms obtained on the AuNDs/NaFion/ANE for the determination of Cu²⁺ with typical interfering species (Ca²⁺ and Fe³⁺) were shown in Fig. 4F. Our results showed that the 100-fold K⁺, Ca²⁺, Mg²⁺, Ni²⁺, Co²⁺, Cr³⁺, NO₃⁻, SO₄²⁻, 50-fold Cd²⁺, Bi³⁺, 40-fold Fe³⁺, 10-fold Pb²⁺ did not affect the determination of Cu²⁺ (<5% of response current change). These findings indicated that the AuNDs/NaFion/ANE had an excellent selectivity for the determination of Cu²⁺. The excellent selectivity of AuNDs/NaFion/ANE was mainly due to the rapid

![Fig. 5. Calibration curve of Cu²⁺ at the AuNDs/NaFion/ANE, where the concentrations of Cu²⁺ are from 0.05 to 10 nmol L⁻¹ in 0.1 mol L⁻¹ acetate buffer (pH 4.3). Inset shows the voltammograms with baseline correction obtained on the AuNDs/NaFion/ANE with successive addition of 0.05, 0.07, 0.1, 0.3, 0.5, 0.7, 1, 3, 5, 7, 10 nmol L⁻¹ Cu²⁺.](image_url)

Table 1

| Methods | Electrodes | Linear range/nmol L⁻¹ | Detection limit/nmol L⁻¹ | Ref. |
|---------|------------|------------------------|--------------------------|------|
| ED-XRF | No | 800–23000 | 200 | [32] |
| FAAS | No | 120–1200 | 10 | [33] |
| GFAAS | No | 130–700 | 1.4 | [34] |
| SWASV | Hg/MWCNTs/CHIT/SPE | 40–200 | 12 | [35] |
| SWASV | Si/AuNP/gold microelectrode | 80–8000 | 30 | [36] |
| DPASV | rGO/Bi/CPE | 300–1600 | 400 | [37] |
| DPASV | L-Cys-rGO/GCE | 400–2000 | 40 | [38] |
| LSSV | CNT/NaFion/GCE | 30–1000 | 50 | [39] |
| OSW | Ph-GO/gold electrode | 300–1600 | 27 | [40] |
| DPASV | Na/SWNTs/rGO/GCE | 200–2000 | 6 | [41] |
| DPASV | rGO/AuNP/NaFion/gold electrode | 20–1000 | 4 | [6] |
| SWASV | Gold microwire electrode | | 0.025 | [42] |
| SWASV | AuNDs/NaFion/ANE | 0.05–1 | 0.015 | this work |

a) ED-XRF, energy dispersive X-ray fluorescence; FAAS, Flame atomic absorption spectrometry; GFAAS, graphite furnace atomic absorption spectrometry; LSSV, linear sweep stripping voltammetry; OSW, Osteryoung square wave voltammetry.

b) Electrode: MWCNTs, multi-walled carbon nanotubes; CHIT, chitosan; CPE, carbon paste electrode; SPE, screen printed electrode; rGO, (reduced) graphene oxide; L-Cys, l-Cysteine; GCE, glassy carbon electrode; CNT, carbon nanotubes; Ph, phenyl group; SA, sodium alginate; SWNTs, single-walled carbon nanotubes.
The electrochemical response of Cu$^{2+}$ and the variety of redox potentials of different electroactive substances.

### 3.6. Speciation analysis of Cu in seawater

The primary purpose of the fabrication of this electrochemical platform based on AuNDs/Nafion/ANE was to investigate the concentrations of different species of Cu in seawater. Standard addition method was adopted to determine the concentrations of different Cu species. Fig. 6A shows the typical SWASV curves of the directly filtered seawater (aliquot 1), and the inset shows the corresponding linear regression equation. It should be noted that the current response obtained without addition of Cu$^{2+}$ in this aliquot was much small, and baseline correction was used for each recorded voltammogram. When dealing with the baseline correction, the voltammogram obtained without accumulation process was used as the background, and the voltammograms obtained with successive addition of Cu$^{2+}$ were presented after the background subtraction. The dissolved active Cu concentration has been calculated to be 1.90 nmol L$^{-1}$. It should be noted that it is very difficult to achieve the determination of dissolved active Cu in seawater due to its low concentration and complex matrix interference. Thanks to the excellent performance for Cu$^{2+}$ detection of the AuNDs/Nafion/ANE, the dissolved active Cu in seawater can be determined directly without any pre-concentration treatments or any additional reagents. Similarly, Fig. 6B and C determined the dissolved acid extractable and total extractable Cu concentrations. Considering that the current responses obtained without addition of Cu$^{2+}$ were remarkable in Fig. 6B and C, there was no need to deal with the background signals. All results of different Cu species of the seawater samples collected from the Sishili Bay were listed in Table 2. The results indicated that our proposed AuNDs/Nafion/ANE was reliable and suitable for the determination of the dissolved active, dissolved acid extractable, and total acid extractable Cu in seawater.

### 4. Conclusions

In summary, a novel and efficient needle-shaped electrode functionalized with AuNDs/Nafion was fabricated for the ultra-sensitive speciation analysis of Cu in seawater samples collected from Sishili Bay, Northern Yellow Sea, China. Thanks to the combined effects of the unique needle-shaped structure of ANE, the excellent electrochemical properties of AuNDs and the specific pre-concentration ability of Nafion, the AuNDs/Nafion/ANE showed excellent performance for the electrochemical determination of Cu$^{2+}$ in seawater. The schematic of voltammetric determination of Cu$^{2+}$ on the so-designed functional needle-shaped electrode was explained as the adsorption-catalysis model. The dissolved active, dissolved acid extractable and total acid extractable Cu concentrations were determined directly at the AuNDs/Nafion/ANE via SWASV. This so-fabricated AuNDs/Nafion/ANE has several advantages, such as easy fabrication procedure, high sensitivity, excellent repeatability and reproducibility, high storage stability, which can be of great benefit for the speciation analysis of Cu and other metals in seawater. Furthermore, the speciation and distribution of other

![Fig. 6. Voltammetric determination of dissolved active Cu (A), dissolved acid extractable Cu (B), and total acid extractable Cu (C) in seawater by standard addition method. The concentrations of Cu$^{2+}$ added are (A) 0, 2, 4, and 6 nmol L$^{-1}$; (B) 0, 5, 10, and 15 nmol L$^{-1}$; and (C) 0, 20, 40, and 60 nmol L$^{-1}$ (from bottom to top). The insets show the corresponding linear regressions respectively.](image)

| Seawater samples | Dissolved active Cu/nmol L$^{-1}$ | Dissolved acid extractable Cu/nmol L$^{-1}$ | Total acid extractable Cu/nmol L$^{-1}$ | Ratio of dissolved active Cu |
|------------------|---------------------------------|-------------------------------------------|----------------------------------------|-----------------------------|
| Location 1       | $1.90 \pm 0.05$                 | $5.76 \pm 0.17$                          | $25.83 \pm 0.78$                       | $7.4 \pm 0.4\%$            |
| Location 2       | $0.99 \pm 0.04$                 | $12.60 \pm 0.34$                         | $45.29 \pm 1.37$                       | $2.2 \pm 0.2\%$            |
| Location 3       | $1.92 \pm 0.06$                 | $6.39 \pm 0.18$                          | $36.42 \pm 1.12$                       | $5.3 \pm 0.3\%$            |
metal elements in different water samples can be assessed by the functional needle-shaped electrode as well.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.electacta.2018.08.057.

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