Signal-on photoelectrochemical immunoassay for salivary cortisol based on silver nanoclusters-triggered ion-exchange reaction with CdS quantum dots

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Abstract
Nowadays, the epidemic, employment, and academic pressures are seriously affecting our physical and mental health. Herein, we designed a magneto-controlled photoelectrochemical immunosensor for noninvasive monitoring of salivary cortisol regarded as a pressure biomarker. A competitive immunoassay model was established by coupling bovine serum albumin-cortisol modified magnetic beads (MB-BSA-cortisol) with silver nanoclusters (Ag NCs)-labelled anti-cortisol antibody, and quantity analysis was operated by photoelectrochemical measurement of the CdS/Au electrode as an ion-exchange platform. Accompanying the formation of immune complexes, the carried Ag NCs were readily dissolved with nitric acid to produce abundant silver ions, which transferred to the electrode for ion-exchange reaction with CdS quantum dots to produce Ag2S, a new electron–hole capture site, leading to a decrease in the photocurrent intensity. The photocurrent signal gradually recovered with the increase of concentration of target cortisol, acquiring the signal-on mode competitive immunosensing system, which is propitious to the detection of small molecules. Within optimal conditions, this sensor had a satisfactory linear relationship in the range of 0.0001–100 ng mL\(^{-1}\) with favorable repeatability, specificity, and acceptable method accuracy. The detection limit was as low as 0.06 pg mL\(^{-1}\). In addition, this strategy provided new thought for the test of other small-molecule analytes and immunosensor applied in the complex biological system.

Keywords Photoelectrochemical · Competitive immunoassay · Ion-exchange reaction · Small-molecule cortisol

Introduction
During the current coronavirus pandemic, it has been reported that discussion of stress etiology is going on increasingly among health care workers as well as the general public [1–3]. Apparently, developing a simple and sensitive method for the detection of stress biomarkers is of great significance in health protection. Cortisol, a biomarker of stress detection in humans considered commonly, in particular, salivary cortisol, has become highly attractive compared to the assays in blood, sweat, or hair due to simple, noninvasive, and well reproducible sampling [4]. Photoelectrochemical (PEC) sensing has recently gained more and more popularity in the fields of biological and chemical analysis [5, 6], which incorporates the excellent merits of photochemical method and electrochemical technology [7], including rapid detection, low cost, good sensitivity, and wide linearity range [8]. Inspiringly, PEC sensing, in the noninvasive monitoring of small molecules in biological fluids, has been reported in succession and demonstrated tremendous progress [9–11]. However, research on the PEC assay of salivary cortisol is still lacking now.

It is well-known that signal-on biosensors are superior to signal-off ones because of their lower background signal and probability for false-positive results [12, 13]. What’s more, in PEC immunoassay, the quantitative analysis of biomarkers is usually carried out through photocurrent changes after biological recognition. Nevertheless, some of the analytes tend to be low in the early stages of diseases. Therefore, it is considerable to employ signal amplification strategies into the PEC system [14–16]. To date, multifarious
signal amplification strategies have been put forward for PEC immunoassay to enhance sensitivity, such as enzyme-mediated catalytic [17], energy transfer [18], chemical redox cycling [19], nano-material labels [20], and other strategies. Among them, ion-exchange technology has opened new roads for designing immune sensors in recent years because of its rapid reaction with another element at room temperature to form a new photoactive species to amplify the change in photocurrent value [21, 22]. Zhang et al. established a PEC sensing system based on an ion-exchange reaction between copper ions (Cu²⁺) and WO₃–Au–CdS nanoparticles, forming the weak photoactive material CuxS to reduce the photocurrent [23]. Song et al. constructed an excellent biosensor for microRNA detection through the ion-exchange reaction between silver ions and CdSe QDs/TiO₂ composites [24]. Likewise, Zhao et al. proposed a PEC platform via the silver nanoclusters-based ion-exchange reaction with CdTe QDs, which dramatically enlarged the PEC response for ultrasensitive target detection [25]. Therefore, PEC immuno-sensors based on ion exchange have great potential in biological analysis in the future.

Since the photocurrent signal arose from the charge separation and migration upon illumination between photoactive material and electrode [8, 26], the active materials play a crucial role in the PEC sensing system. With the continuous development of semiconductor nano-material, they are increasingly widely used in PEC fields, particularly quantum dots (QDs), attributed to their efficient photoelectric conversion, desirable optoelectronic property, adjustable band gap, and unique biocompatibility [27–29]. CdS, an n-type semiconductor with excellent visible light activity, has become one of the intensively studied photoelectric materials, yet pure CdS also suffers from severe photo-corrosion [30]. Fortunately, this drawback can be overcome by doping Au nanoparticles (NPs) into CdS due to the surface plasmon resonance (SPR) of Au NPs, which acts as electron transfer mediator, effectively separates and transfers photogenerated charge carriers [31–33].

Consequently, our motivation is to develop a novel PEC immunosensing platform for small-molecular cortisol in saliva via the ion-exchange reaction between Ag NCs as the labels and Au NPs decorated CdS quantum dots as photoactive receptors. As shown in Scheme 1, this assay was a competitive-immune and split-type detection mode. In a centrifugal tube, bovine serum albumin-cortisol modified with magnetic beads (MB-BSA-cortisol) were used as competitors to compete with the target cortisol for silver nanoclusters labelled anti-cortisol antibody (Ab-Ag NCs). Then, magnetic separation removed both the Ab-Ag NCs bound target and the unbound target cortisol. Subsequently, the carried Ag NCs onto magnetic beads accompanying the antigen–antibody reaction was expected to release numerous Ag⁺ with the aid of nitric acid (HNO₃). Finally, the as-released Ag⁺ could exchange with CdS QDs to form Ag₂S nanocrystals, which mediated the exciton trapping and resulted in a remarkable reduction in the photocurrent of the CdS/Au photoelectrode. The above ion-exchange reaction: [(CdS)n+2n•Ag²⁺(aq)→(Ag₂S)n+n•Cd²⁺(aq)] was feasible at room temperature, accompanied by the color changing from yellow to dark brown, which was on account of the fact that the solubility product constant of Ag₂S (Ksp~6.69×10⁻⁵⁰) was lower than that of CdS (8.00×10⁻²⁷), rendering fractional replacement of Cd²⁺ ions with Ag⁺ ions [34]. MB-BSA-cortisol not only served as competitors to participate in competitive immunoassay that was recognized as the preferred method for small-molecule immunoassay but also facilitated excluding coexisting species from actual samples under external magnetic fields to enhance specificity. Also, as Ag NCs could release a large number of silver ions for ion exchange compared with silver nanoparticles and achieve
double signal amplification, the sensitivity of such a PEC biosensor is doomed to be lower. As expected, the as-designed PEC sensors were capable of monitoring cortisol in saliva samples, and its photocurrent signal is proportional to the target cortisol level with excellent selectivity and sensitivity. Hence, it had potential applications in stress testing.

**Experimental section**

**Chemicals and materials**

Anti-cortisol mouse monoclonal antibody (XM210, Ab) was ordered from Abcam (England). Thiocicic acid (99%, m/m), Cadmium chloride hemi (pentahydrate) (CdCl$_2$·2.5H$_2$O, 98%, m/m), hydrocortisone (cortisol, 98%, m/m), and progesterone (Pro, 98%, m/m) were all obtained from Macklin Biochemical Co., Ltd. (China). 1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide hydrochloride (EDC, 99%, m/m) was purchased from Shanghai Macklin Biochemical Co., Ltd. (China). N-hydroxysuccinimide (NHS, 99.0%, m/m) and bovine serum albumin (BSA) derived from Beijing Solarbio Science Technology Co., Ltd. (China). Silver nitrate (AgNO$_3$, 99.8%, m/m), sodium borohydride (NaBH$_4$, 96.0%, m/m), iron trichloride hexahydrate (FeCl$_3$·6H$_2$O, 99%, m/m), and tetrachloroaaric (III) acid tetrahydrate (HAuCl$_4$·4H$_2$O, 47.8%, m/m) were all obtained from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). Glucose (Glu) was purchased from Chengdu Jinshan Chemical Test Co., Ltd. (China). Sodium sulfide nonahydrate (Na$_2$S·9H$_2$O, 98.0%, m/m) was all obtained from Xilong Scientific Co., Ltd. Nitric acid (HNO$_3$, 65.0–68.0%, v/v) was purchased from Chongqing Chuandong Chemical Co., Ltd. (China). Phosphate buffer solution (PBS) was prepared with the following reagents: sodium phosphate dibasic (Na$_2$HPO$_4$, 99.0%, m/m) and sodium dihydrogen phosphate (NaH$_2$PO$_4$) with monitoring by pHST-310 Ohaus pH meter (USA), which was calibrated via standard pH buffer solutions. Ultrapure water used in all solutions was from tap water purified by a Water Milli-Q system (Merck KGaA, Germany). Human salivary cortisol enzyme-linked immunosorbent assay (ELISA) kit (Cat. No. RXJ103617H; inter-assay CV <15%; sensitivity: 0–0.036 ng mL$^{-1}$; examination area: 1.1–17.4 ng mL$^{-1}$) was purchased from Ruixin Biotech (China).

**Apparatus**

X-ray diffraction (XRD) analysis was implemented on a Shimadzu XRD-6000 diffractometer (Shimadzu, Shimadzu, Japan). High resolution transmission electron microscopy (HRTEM) characterization was completed on a JEM-2100F microscope (JEOL, Tokyo, Japan). The characterization of elements was carried out by a Thermo Scientific Escalab 250Xi X-ray photoelectron spectroscopy (Thermo, MMAS, USA). The UV–vis absorption spectra were tested on a UV-5500PC UV–visible spectrophotometer (Metash, Shanghai, China). F-4700 spectrophotometer was used to perform the fluorescence spectra (Hitachi, Tokyo, Japan). The zeta potential was estimated by NanoBrook Omni Brookhaven Instruments (BIC, Brookhaven, USA). The FT-IR characteristic measurement was carried out by FT-1600400 Spectrum Two infrared spectrometer (PerkinElmer, Llantrisant, UK). Electrochemical impedance spectroscopy (EIS) and cyclic voltammetry (CV) were carried out at the CHI 660E electrochemical workstation (Shanghai Chenhua Instrument, Shanghai, China).

**Preparation of Ab-Ag NCs and MB-BSA-cortisol**

The synthesis process of Ag NCs and MB was described in the electronic supplementary information (ESM). Like the reported method, the as-prepared Ag NC was used to label anti-cortisol antibodies [35]. First, 1 mL Tris–EDTA buffer solution contained 3 mM glutathione was added to a brown Eppendorf tube, subsequently joining anti-cortisol antibody (100 μL, 100 μg/mL) together with 188 μL of Ag NCs and 1 mL solution containing EDC and NHS (0.01 M). The above solution was mixed evenly on the vortex apparatus and then shaken gently on a shaker for 2 h at room temperature. Finally, purification (100 kDa pore size, Amicon ultra centrifugal filter units) was performed using the following process: a first cycle of 5500 rpm for 10 min and two ultrapure water cleaning cycles at 5500 rpm for 5 min. The Ag NCs-labelled anti-cortisol antibody (Ab-Ag NCs) in the filter was dispersed in PBS (2 mL, 0.1 M, pH = 7.4) and was stored in the fridge.

According to the previous method [36], the magnetic beads need to carboxylate before biological coupling. In the first, 1.0 g of bare magnetic beads was added to a round-bottomed flask containing 100 mL ultrapure water followed by ultrasonic dispersion. Then, under nitrogen protection, 2.5 g of citric acid was dissolved in the suspension and kept for 30 min, and heated the suspension to 70 °C for refluxing 1 h. At the end, using a permanent magnet to collect carboxylated magnetic beads after thorough washing with ultrapure water, dried in a vacuum. The carboxylated magnetic beads and BSA-cortisol were conjugated by a classical carbodiimide coupling referring to the previous report with slight modifications [11]. In brief, 1.0 mL of the carboxylated magnetic beads (5.0 mg/mL) was dispersed in 1.0 mL PBS solution (0.1 M, pH 7.4) with 100 mM NHS and 400 mM EDC. The resulting solution was then lightly shaken for 60 min before being magnetically separated and washed three times with ultrapure water to take out superfluous EDC and NHS. Afterwards, the active magnetic beads
were dispersed in a BSA-cortisol solution (5 mg mL⁻¹) and shaken gently overnight. Finally, the obtained BSA-cortisol modified with magnetic beads (MB-BSA-cortisol) was dispersed in 2.0 mL PBS and stored in the refrigerator.

**Photoelectrode preparation and PEC immunoassay**

The synthesis process of CdS QDs and Au NPs is shown in ESM. The fluorine-tin-oxide electrode (FTO) electrode was cleaned by sonication with acetone, ethanol, and ultrapure water in sequence and dried at 60 °C before use. Then, the waterproof transparent tape punched with a hole (r=0.28 cm, S=0.25 cm²) was stuck on the surface of the electrode. After that, 30 μL of Au NPs colloids was dropped into the hole of the FTO electrode. After drying naturally at room temperature, 20 μL CdS QDs solution was also coated into the hole of the FTO electrode. After that, 30 μL of Au NPs colloids was dropped into the hole of the FTO electrode. After drying naturally at room temperature, 20 μL CdS QDs solution was also coated on the surface of the above electrode and dried at 40 °C.

Scheme 1 describes the schematic illustration of the PEC sensing system for target cortisol based on Ag NCs-aided ion-exchange reaction with CdS quantum dots. The competitive reaction was carried out in a 500 μL centrifugal tube containing 100 μL of the as-synthesized MB-BSA-cortisol. First, 50 μL of target cortisol standard/sample together with 50 μL of Ab-Ag NCs (certain concentration as prepared) were injected into the tube and incubated at 37 °C for 50 min on a shaker. The resulting suspension was magnetically separated and washed twice with PBS (0.01 M, pH = 7.4) to remove unbound antibodies and target cortisol. Following that, 30 μL HNO₃ (0.1 mM) was added into the tube and reacted for 15 min to release abundant Ag⁺ from the competed Ag NCs labels. Afterwards, with the aid of magnetic, 20 μL of supernatant droplet containing Ag⁺ was deposited on the CdS/Au-modified FTO electrode, then kept at room temperature for 20 min to execute the ion-exchange between Ag⁺ and CdS QDs. At last, the resulting electrode was washed with ultrapure water for subsequent PEC detection. The photocurrent was tested in 0.1 M Na₂SO₄ solution (10 mL) with a traditional three-electrode system using the obtained electrode as the working electrode, a platinum wire electrode as the counter electrode, and a saturated Ag/AgCl electrode as the reference electrode. The electrochemical workstation configured with a 500 W Meze lamp (Beijing NBET Technology, China) as the excitation light source, which switched on and off every 10 s.

**Results and discussion**

**Materials characterization**

In practical application, it was necessary to confirm whether the prepared CdS QDs can be exchanged by Ag⁺. For this purpose, we used different techniques to characterize CdS QDs before and after exchange with certain Ag⁺ ions. Figure 1A showed a HRTEM image of the as-prepared CdS QDs with an average size of approximately 8 nm. When CdS QDs reacted with excessive Ag⁺, we dimly saw from Fig. 1B that the shape and lattice of the resulting crystal were diverse those of alone CdS QDs (morphologies of other magnifications in Fig. S1). In addition, by comparing curve “black” vs curve “red” in Fig. 1C, we observed clearly the XPS characteristics of the Ag 3d double peak for the resulting Ag⁺, indicating that silver element exists in nanocrystals. For better understanding, we had also carried out the core-level spectra of Cd and Ag, as exhibited in Fig. S3A and S3B. It was evident that the intensity of Cd 3d₅/₂ and 3d₃/₂ decreased, while it was seen that of Ag 3d₅/₂ and 3d₃/₂ increased correspondingly. Furthermore, the crystal structures of the materials successively modified on the electrodes were characterized by XRD in Fig. 1D. Except for the diffraction peaks of the FTO substrate, several typical characteristic peaks of CdS QDs located at 2θ=26.5°, 43.8°, and 51.8° were indexed to the (111), (220), and (311) crystal planes, which coincided with the phase of CdS QDs (JCPDS 80–0019) [37]. And, the characteristic peak of Ag₂S generated by ion exchange on the electrode was not obvious, which could be caused by overlapping with that of CdS QDs. Nevertheless, it varied significantly at a peak magnification of about 37.5° in Fig. S3C, as done that of the Au NPs.

The structural and optical characteristics of the prepared Ag NCs were characterized by UV–vis, fluorescence spectra, and TEM. Figure 2A demonstrated three characteristic absorption spectra of the Ag NCs (curve a), centered at 329, 427, and 497 nm respectively which was in accordance with previous reports [38]. Then, the Ag NCs were functionalized with anti-cortisol antibodies to prepare signal nanoprobes and perform spectral characterization. As shown in Fig. 2A, the spectrum of Ab-Ag NCs (curve c) consisted of the characteristic peak of antibody at about 260 nm (curve b) and that of Ag NCs, indicating that the functionalization was successful. Besides, from Fig. 2C, for zeta potential, the negative charges of Ag NCs reduced after modification with Ab, which also validated the successful conjugation of Ab onto Ag NCs. The emission spectra of the Ag NCs present a maximum of about 670 nm in Fig. 2B. However, the fluorescence characteristics disappeared under HNO₃ treatment attributed to generating ions, which served as a source of Ag⁺ for ion-exchange reactions. Moreover, the size and shape of Ag NCs were characterized through TEM. As illustrated in Fig. S2A, the Ag NCs were uniformly distributed with a mean diameter of 2–5 nm and could be seen a distinct lattice plane with the inter-fringe distance of 0.22 nm. In addition, the biological coupling of magnetic beads is also characterized by UV–vis shown in Fig. 2D, and the MB-BSA-cortisol had both the absorption spectrum of MB and BSA-cortisol, demonstrating the successful fabrication.
Furthermore, the FT-IR spectra further confirmed this result, as shown in Fig. S2B, and the absorption peaks of the synthesized MB-COOH are consistent with the literature [39]. Compared with the MB-COOH, the MB-BSA-cortisol presented newly absorbance peaks located at 1540 cm\(^{-1}\) and 1630 cm\(^{-1}\), which are ascribed to C=C bonds from aromatic rings of the cortisol molecule [40]. In addition, a broad peak of -OH at 3200 cm\(^{-1}\) shifted to lower wave numbers, which may be due to the hydrogen bond association between hydroxyl group on cortisol surface and water.

**Investigation of PEC performance and feasibility**

It is crucial to evaluate the PEC property of the prepared CdS/Au electrode, which directly determines the monitoring performance of the analyte. As depicted in Fig. 3A, the photocurrent response of Au NPs modified electrode was very weak, almost the same as that of bare FTO electrode. After the assembly in turn of Au NPs and CdS QDs, the photocurrent intensity of CdS/Au, as expected, was much stronger than that of CdS alone; also, the photocurrent was stable and reliable (in Fig. S4A). The enhanced PEC response was mainly due to the modification of Au NPs on CdS QDs, which greatly promoted the separation and transfer of photogenerated electron–hole pair at the CdS/Au Schottky junction interface. EIS measurements were also conducted for the different modified FTO electrodes. In the Nyquist diagram, there are two main arcs in the high-frequency region (semicircle, \(R_{ct}\)) and the low-frequency region (line segment, \(R_{w}\)), respectively, representing the charge transport resistance and the recombination resistance, where the recombination resistance is inversely proportional to the recombination rate. As shown in Fig. 3B, it was clear that \(R_{ct}\) of CdS/Au was much lower than that of CdS alone, while \(R_{w}\) was slightly higher than that of CdS, namely, the recombination rate of CdS/Au electrode was lower. In addition, it also could be verified that the introduction of Au NPs could improve conductivity and promote electron transport from the CV curves in the inset. Taken together, all of the above results reveal the excellent PEC performance of the CdS/
Au electrode. In addition, we also performed the complete electrochemical characterizations of the sensor (please see ESM and Fig. S5).

Furthermore, when Ag$^+$ was introduced into the system, the photocurrent of the CdS/Au electrode decreased significantly than that of the pure CdS electrode (Fig. 3C, “II” vs “I”), accompanying different color changes in the inset, respectively. Significantly, the photocurrent signal decreases successively with the increase of Ag$^+$ concentration in Fig. S4B. According to the results obtained, it could be concluded that the CdS/Au electrode could be used to develop a competitive immunoassay using the Ag NCs-labelled strategy. As demonstrated in Fig. 3D, curve “a” displayed the photocurrent response of CdS/Au-modified FTO. When the modified electrode was applied to the immune system, the photocurrent decreased significantly without target (curve “b”). In comparison, curve “c” exhibited the photocurrent recovered with target cortisol (1.0 ng mL$^{-1}$) intervention. The results revealed that silver ions produced by the immune complex after acid treatment resulted in a significant reduction of photocurrent signal; meanwhile, the photocurrent was proportional to the target concentration. In order to understand why the photocurrent decreased, we study the electronic band structures of CdS QDs by the Mott–Schottky (M-S) analyses in Fig. S6, where, please see ESI for detailed calculation of $E_g$, $E_{cb}$, and $E_{vb}$ of CdS QDs. Notably, the positive slopes of the tangents line in the M-S plots for CdS QDs by the Mott–Schottky (M-S) analyses in Fig. 3.4B. According to the results obtained, it could be concluded that the CdS/Au electrode could be used to develop a competitive immunoassay using the Ag NCs-labelled strategy. As demonstrated in Fig. 3D, curve “a” displayed the photocurrent response of CdS/Au-modified FTO. When the modified electrode was applied to the immune system, the photocurrent decreased significantly without target (curve “b”). In comparison, curve “c” exhibited the}

**Fig. 2** A UV–vis absorption of (a) Ag NCs, (b) antibody, and (c) Ab-Ag NCs. B The fluorescence of Ag NC disappeared after acid treatment. Inset: photos of the Ag NCs under visible light (A) and 365 nm UV light irradiation (B); C change of zeta potential after modification of antibody onto Ag NCs. D the absorption spectra of MB, BSA-cortisol, and MB-BSA-cortisol (Inset: the superparamagnetic behavior of as-prepared MB under an external magnetic field).
Ag$_2$S (+0.92 eV), thus forming the trapping sites in Fig. S7. The above demonstrated that the signal-on and competitive-type PEC immunesensor was successfully constructed based on ion-exchange reaction on the CdS/Au electrode.

**Analytical performance of PEC immunoassay**

To obtain the best performance of the PEC biosensor, it was necessary to optimize its potential, competitive immune time and ion-exchange reaction time. First, as indicated in Fig. S8A, a lower potential would minish the photocurrent of the CdS/Au electrode, while a higher one also did not significantly increase the photocurrent. Therefore, 0 V was selected for PEC measurement in this work. Next, as can be seen from Fig. S8B, with the extension of reaction time, the photocurrent gradually reduced, reaching a stable value at 50 min and no longer decreasing. So, the reaction time of 50 min is the best time for a competitive immune system. Certainly, it also greatly influenced the photocurrent of the system in the ion-exchange time between CdS QDs and the immuno-reaction product (Ag$^+$). As shown in Fig. S8C, with the increase of reaction time, the photocurrent of the CdS/Au electrode gradually decreased and levelled off after 15 min. In order to minimize the determination time, it was appropriate that the ion-exchange reaction is 15 min. Under the above best experimental parameters, the competitive-type immunosensing system was used for subseuent detection. Figure 4A presents the photocurrents of the PEC biosensor toward different concentrations of the target cortisol. It was observed that the photocurrent signals gradually increased with increasing cortisol concentration. As observed from Fig. 4B, the photocurrent had a well-fitted linear dependence with the logarithm
of cortisol concentration within the dynamic range of 0.0001–100 ng mL\(^{-1}\). The linear regression equation was \(y = -0.50 \log x + 4.67\) \((R^2 = 0.998)\) with a limit of detection (LOD) of 0.06 pg mL\(^{-1}\). Compared to other immunoassays, as seen from Table S1, our assay offered a wider detection range and a lower detection limit. To investigate the reproducibility of the method, eight independent measurements of 1 ng mL\(^{-1}\) cortisol were performed in Fig. 4C, respectively, and the RSD values of the results were 1.20%, indicating that the method had good reproducibility. Furthermore, the selectivity of this PEC immunosensor toward cortisol (1 ng mL\(^{-1}\)) was also evaluated by non-target analytes, including glucose (Glu), progesterone (Pro), and BSA, and the result is shown in Fig. 4D. The system had little response to a high concentration (100 ng mL\(^{-1}\)) of non-target analytes. On the contrary, the photocurrent signal of the mixture containing (1 ng mL\(^{-1}\)) cortisol and the non-target analyte resembled that of cortisol alone, suggesting that the PEC biosensor was extremely specific for cortisol.

Detection of real samples

To further evaluate the potential application and feasibility of Ag NCs-labelled PEC immunosensor, we examined cortisol in human saliva samples, which range from 0.6 to 12.85 ng mL\(^{-1}\) in healthy people [42, 43]. Firstly, saliva from four volunteers was collected twice in the morning and afternoon and detected by this method and ELISA after certain sample treatment (see ESM for saliva collection in details). As shown in Table 1, the detection results of the prepared PEC biosensor were close to that of ELISA. Importantly, all Student’s \(t\)-test values of these samples were under \(t_{\text{crit}}\) \((t_{\text{crit}}[0.05, 4] = 2.78)\), indicating excellent reliability of this system. Surprisingly, we found vaguely that cortisol levels in the morning slightly more than in the afternoon.
suggesting that people secreted higher cortisol in the morning, consistent with previous reports.

Conclusions

In summary, we have developed a signal-on photoelectrochemical immunoassay for salivary cortisol based on silver nanoclusters-triggered ion-exchange reaction with CdS quantum dots. As markers of detection antibody in competitive immunocomplexes, Ag NCs released thousands of Ag⁺ ions with acidic conditions, and Ag⁺ ions were transferred to the PEC electrode for ion-exchange reaction with CdS QDs, decreasing the photocurrent and achieving signal-on detection for photoelectrochemical detection of cortisol is as low as 0.06 pg mL⁻¹. The highlights of our strategy lie in the following issues: (i) the bio-recognition of this PEC sensor was performed in the liquid phase instead of on the solid–liquid interfaces, avoiding the time-consuming and complex defects of assembling photoelectric beacons, identification elements, and targets on the electrodes, boosting the stability and reproducibility; (ii) the hole-exciton capture was easily mediated by Ag₂S generated by the ion-exchange reaction between the undissolved Ag⁺ and CdS QDs, thus led to significant amplification in photocurrent intensity; (iii) magnetic bead labelling could effectively separate other interferences and improve selectivity. In addition, the proposed PEC biosensor was used to test human salivary cortisol samples, displaying satisfying feasibility and accuracy compared to a commercial ELISA kit. Therefore, this work provided an alternative for the detection of pressure markers as well as other small-molecule and even PEC immunosen sor applied in the complex biological system.

Table 1 Comparison of the measured results between the as-prepared PEC biosensor and the human salivary cortisol ELISA kit

| Time       | Sample no | Method; level (mean ± SD, ng ml⁻¹, n = 3) | t_exp |
|------------|-----------|------------------------------------------|-------|
|            |           | PEC immunoassay                           | ELISA kit |
| Morning    | 1         | 1.05 ± 0.09                              | 0.90 ± 0.06 | 2.37 |
|            | 2         | 2.91 ± 0.11                              | 2.81 ± 0.16 | 0.90 |
|            | 3         | 1.10 ± 0.02                              | 0.98 ± 0.11 | 1.98 |
|            | 4         | 10.43 ± 0.43                             | 9.36 ± 0.75 | 2.15 |
| Afternoon  | 1         | 1.06 ± 0.06                              | 0.97 ± 0.12 | 1.13 |
|            | 2         | 2.39 ± 0.15                              | 2.09 ± 0.16 | 2.30 |
|            | 3         | 0.97 ± 0.05                              | 0.98 ± 0.10 | 0.20 |
|            | 4         | 9.35 ± 0.15                              | 9.20 ± 0.28 | 0.81 |

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Declarations

The authors declare no competing interests. The use of human salivary was granted by the Ethics Committees of Guizhou University (Guizhou, China) (approval number: EAE-GZU-2021-T044). The source of human salivary was obtained from healthy volunteers, which were students in the College of Pharmacy, Guizhou University.

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