Rapid and quantitative detection of C-reactive protein based on quantum dots and immunofiltration assay

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Abstract: Convenient and rapid immunofiltration assays (IFAs) enable on-site “yes” or “no” determination of disease markers. However, traditional IFAs are commonly qualitative or semi-quantitative and are very limited for the efficient testing of samples in field diagnostics. Here, we overcome these limitations by developing a quantum dots (QDs)-based fluorescent IFA for the quantitative detection of C-reactive proteins (CRP). CRP, the well-known diagnostic marker for acute viral and bacterial infections, was used as a model analyte to demonstrate performance and sensitivity of our developed QDs-based IFA. QDs capped with both polyethylene glycol (PEG) and glutathione were used as fluorescent labels for our IFAs. The presence of the surface PEG layer, which reduced the non-specific protein interactions, in conjunction with the inherent optical properties of QDs, resulted in lower background signal, increased sensitivity, and ability to detect CRP down to 0.79 mg/L with only 5 µL serum sample. In addition, the developed assay is simple, fast and can quantitatively detect CRP with a detection limit up to 200 mg/L. Clinical test results of our QD-based IFA are well correlated with the traditional latex enhance immune-agglutination aggregation. The proposed QD-based fluorescent IFA is very promising, and potentially will be adopted for multiplexed immunoassay and in field point-of-care test.

Keywords: C-reactive proteins, point-of-care test, Glutathione capped QDs, PEGylation

Introduction

Serum C-reactive protein (CRP) is a routinely used biomarker for clinical detection of inflammation and rapid discrimination between bacterial and viral infections in humans.1–3 The normal concentration of CRP in the serum of a healthy human is usually lower than 10 mg/L, while higher levels of CRP are usually related to different health conditions. CRP levels increase to 10–40 mg/L with the incidence of viral infections and mild inflammation, 40–200 mg/L with the incidence of bacterial infection and active inflammation, and >200 mg/L with the incidence of severe bacterial infections and burns.4

Recent clinical research revealed that increased CRP levels, even within the range previously considered normal, strongly predict future coronary events.1,4,5 The American Heart Association and US Centers for Disease Control and Prevention6 have associated CRP levels over 1.0 mg/L with an increased risk of cardiovascular disease. However, conventional CRP detection methods for the clinical diagnosis of acute inflammation are not sensitive enough to detect cardiovascular risks according to this criteria. To measure the trace amount of CRP with high sensitivity, previous studies have reported using magnetic immunosassays with detection limits as low as 0.12 µg/mL.7,8 Fluorescent immunosorbent assays were also developed for sensitive...
Materials and methods

Materials

Cadmium oxide (CdO, 99.99%), trialkylphosphine oxide (TOPO, 90%), trialkylphosphine (TOP, 90%), octadecylamine (ODA, 90%), N-hydroxysuccinimide (NHS, 98%), diisopropylcarbodiimide (DIC, 99%), and bovine serum albumin (BSA) were purchased from Sigma-Aldrich Co. (St Louis, MO, USA). Sulfo-N-hydroxysuccinimide (sulfo-NHS, 98.5%) and 1-ethyl-3-(3-di-methylaminopropyl) carbodiimide hydrochloride (EDC, 99%) were purchased from Thermo Fisher Scientific (Waltham, MA, USA). Selenium (Se, 99.99%), stearic acid (SA, 99%), sulfur powder (99.9%), zinc oxide (ZnO, 99%), oleic acid (OA, 99%), liquid paraffin, and reduced GSH (90%) were obtained from Sinopharm Chemical Reagent Co., Ltd (Shanghai, People’s Republic of China). Carboxyl-PEG-carboxyl (Mn =2,000 Da) was purchased from Jiaxing Biomatrix and Biotechnology Inc. (Jiaxing, People’s Republic of China) Standard CRP and monoclonal mouse-anti-CRP antibody were provided by Shanghai Upper Bio-Tech Pharma Co., Ltd (Shanghai, People’s Republic of China). CRP free serum obtained from HyTest Ltd (Turku, Finland) were used for preparation of standard CRP samples. All the reagents were used as they were received and without any further purification.

Synthesis of QDs

CdSe/ZnSe/ZnS core/shell/shell QDs were used for immunooassay because of their improved QYs and photostability when compared to CdSe/ZnS core/shell QDs. The typical procedure of QDs synthesis follow. CdSe core were synthesized and coated with ZnSe and ZnS multi-shell according to the previously reported method. Briefly, CdSe cores were synthesized by heating a mixture of 1 mmoL CdO powder, 4 mmoL SA, and 10 mL liquid paraffin. The mixture solution was degassed and heated to 200°C with magnetic stirring until CdO was completely dissolved. After cooling to room temperature, 1 g TOPO, 3 g ODA, and 10 mL liquid paraffin were added to the flask, heated to 280°C under nitrogen protection, and then 5 mL of 1 M Se in TOP was rapidly injected into the Cd-containing reaction mixture followed by cooling to room temperature after 30 minutes reaction at 260°C.

For ZnSe-ZnS shell coating process, CdSe cores isolated by repeated precipitations from chloroform with methanol, were heated to 220°C in a mixture of 5 g ODA and 15 mL liquid paraffin under nitrogen protection. Then, aliquots of Zn and Se/S precursor solutions were alternately introduced starting with Zn precursor, waiting 20 minutes between each
addition. After shell coating, QDs were annealed at 260°C for another 20 minutes, and then cooled to room temperature. Finally, core-shell QDs were isolated from the solution with chloroform and methanol via centrifugation, washed three times, and stored in chloroform.

Surface modification of QDs
In a typical procedure, 250 mg of GSH and 200 mg of sodium hydroxide were dissolved in 10 mL of methanol, and mixed with 5 mL of 10 µM ODA-coated CdSe/ZnSe/ZnS QDs chloroform solution. Afterward, the solvents in the mixture were gradually evaporated with magnetic stirring under room temperature overnight, and then 25 mL of water was added to disperse the precipitates. Subsequently, 1 mL of 1 M ZnCl₂ solution was added dropwise while stirring and heated to 60°C for 10 minutes to enhance the QYs. Then the QDs solution were filtered through a 0.22 µm syringe filter to remove the aggregates, and dialyzed against pure water for 48 hours with a 14 kDa molecular weight cutoff (MWCO) dialysis tubing to remove the excess of unbound GSH. Finally, GSH-QDs aqueous solution was concentrated to approximately 5 µM by rotary evaporation under reduced pressure.

To improve the colloidal and chemical stability of QDs as in our previous report, PEGylated polymer caged QDs were prepared using carboxyl-PEG-carboxyl as crosslinkers. Typically, 5 µmole of NHS and 5 µmole of DIC were added into 10 µmole of carboxyl-PEG-carboxyl in 5 mL dimethyl sulfoxide (DMSO) solution, and allowed to react for 30 minutes at room temperature to activate the carboxyl groups. Afterward, DTT was added into the solution to quench the excess DIC, and GSH-QDs were then added to the solution, and the reaction continued for another 4 hours. The resultant mixture was dialyzed against pure water for 48 hours to carefully remove the unbound PEG and other small molecules using a 14 kDa MWCO dialysis tubing. The solution was then finally concentrated by rotary evaporation under reduced pressure. During the course of our study, different ratios of DIC, NHS, PEG, and QDs were used to obtain optimal methodologies.

Preparation of QD-antibody conjugates
The covalent conjugation of antibody onto QDs was achieved through using the classic strategy of EDC coupling reaction between the carboxyl groups on the surface of QDs and the amino groups in the antibodies. Typically, 100 µL of QDs (5.0 µM) was dispersed in 200 µL phosphate-buffered saline (PBS) buffer (10 mM, pH 6.0) in a tube. Sulfo-NHS (500 nmoL) and EDC (500 nmoL) were added and incubated for 30 minutes at room temperature to activate the carboxyl groups. Over excess activators were removed by centrifugation at 20,000 rpm for 0.5 hour and carefully removing the supernatant. The resultant pellet was dispersed in 500 µL of borate buffer (10 mM, pH 8.0). Following this, anti-CRP labeling antibodies (120 µg) were added into activated QDs and the mixture was incubated at room temperature for 2 hours. In order to block unreacted carboxyl groups on the surface of QDs, BSA (0.5 mg) was added into the tube, and incubated for another 30 minutes. The mixture was purified by ultracentrifugation at 20,000 rpm for 1 hour, and then washed twice with PBST buffer (PBS with 0.05% Tween® 20) to remove free antibodies and other unreacted small molecules. The purified QD-antibody conjugates were finally dispersed in PBS buffer and stored at 4°C.

Characterization of the prepared QDs and QDs-antibody conjugates
Absorption (Ab) spectra were acquired with an ultraviolet-visible (UV-Vis) spectrophotometer (Shimadzu 2450; Shimadzu Corporation, Kyoto, Japan), and fluorescent spectra were recorded on a fluorescence spectrometer (PerkinElmer Inc., Waltham, MA, USA). The morphology and size of QDs were analyzed with a transmission electron microscope (TEM; Hitachi-7650, Hitachi Ltd, Tokyo, Japan) at an accelerating voltage of 100 kV. The hydrodynamic diameters and size distribution of QDs were determined by a NICOMPTM ZLS system (Particle Sizing Systems, Port Richey, FL, USA). The particle size distribution analysis was conducted by the NICOMP number-weighted distribution analysis.

Immunofiltration assay
QDs-based immunoassay was performed using an immunofiltration system. The system was mainly composed of a thin, porous NC membrane on which CRP monoclonal antibody was immobilized. The membrane was positioned over an absorbent paper and sealed in a plastic cassette containing a hole to expose the membrane. The sample was firstly mixed with QD-antibody conjugates, and antigens in the sample were to be captured by QD-antibody conjugates. Subsequently, the assay was typically performed as follows: First, 5 µL of the serum sample was added into the QD-antibody conjugates solution and gently pipetted for a minimum of three times. Second, 120 µL of conjugates solution was added into the hole of immunofiltration device. After the conjugates solution totally soaked in, 200 µL of washing buffer was added to rinse the unbound conjugates. The whole assay can be finished within 5 minutes. The test
results could be read by naked eye under UV light (365 nm) illumination, and the corresponding fluorescent images were captured directly by a Canon digital camera (G12; Canon Inc., Tokyo, Japan). The fluorescent intensities of QDs on test spots were measured by an optical fiber spectroscopy couple with a UV laser (405 nm).

Clinical samples
A total of 50 human serum samples from 43 patients with chronic hepatitis B virus (HBV), and seven patients with hepatitis C virus infections were obtained from Shanghai Changzheng Hospital. All clinical samples were firstly analyzed and the levels of CRP were measured using a commercial latex agglutination kit (Shanghai Kehua Medical Instruments Co., Ltd, Shanghai, People’s Republic of China), along with our developed QDs-based IFA. The study and data analyses were approved by the Ethics Committee of Shanghai Changzheng Hospital. Patient consent was not obtained as all personal identifiers and patient information were delinked from the serum specimens.

Results
Assembly and working principle of the QDs-immunofiltration assay
The developed fluorescent QD-IFA was used for sensitive and rapid detection of the CRP marker. It is mainly comprised of a porous NC membrane modified with CRP-capturing antibody and an absorbent pad assembled in a plastic cassette (Figure 1A); the working principle is schematically shown in Figure 1B. The serum sample is firstly added to the labeling QDs-antibody conjugates. When CRP is present in the sample, it specifically interacts

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**Figure 1** Schematic illustration of the PEGylated QDs-based immunofiltration assay.

**Notes:** (A) The structure illustration of the immunofiltration pad. (B) The detection process includes an initial step of serum sample mixing with QD-antibody conjugate followed by sample loading and washing with buffer. (C) The qualitative detection is monitored by naked eye under UV light illumination, while the quantitative detection of the resulted fluorescent signals is carried out using an optical fiber spectroscopy and 405 nm laser excitation.

**Abbreviations:** QD, quantum dot; UV, ultraviolet; NC, nitrocellulose; PEG, polyethylene glycol.
with the labeling conjugates. The formed complex is then specifically captured by antibodies previously immobilized on the NC membrane. After a simple washing step and under UV light illumination, the formed three-component sandwich immunocomplexes can be directly visualized by the naked eye for qualitative detection of CRP or quantitatively measured using an optical fiber spectroscopy (Figure 1C). The measured fluorescence intensity was observed to significantly increase with the gradual accumulation of QD conjugates on the antibody spot indicating the potentiality of the developed QD-IFA assay for accurate quantitative detection of CRP.

Characterization of the prepared QDs and QD-antibody conjugates

High quality multi-shell CdSe/ZnSe/ZnS QDs coated with ODA/TOPO were synthesized for the preparation of labeling probes. Hydrophobic QDs with a QY over 50% were solubilized in water by replacing the ODA/TOPO ligands from their surface with GSH. The obtained water-soluble QDs displayed a high QY around 30% and an ultra-small hydrodynamic diameter of less than 10 nm (Figures S1 to S4). PEG derivatives with terminal COOH groups were grafted on the surface of GSH-QDs with controlled ratio (Figure 2A). Identical absorbance

Figure 2 Synthesis and characterization of PEGylated QDs and QD-antibody conjugates.

Notes: (A) Schematic of QD and QD-antibody structure applied for CRP detection using IFA system. Identical fluorescence emission (B) and UV-Vis absorbance (C) spectra of GSH-QDs (black circle) and PEGylated GSH-QDs (red rhombus) dissolved in PBS buffer solution (0.01 M, pH 7.2). (D) Agarose gel electrophoresis of QDs and QD-antibody conjugates, running at 80 V for 20 minutes.

Abbreviations: Ab, antibody; QD, quantum dot; UV-Vis, ultraviolet-visible; IFA, immunofiltration assay; CRP, C-reactive protein; GSH, glutathione; PEG, polyethylene glycol; PL, photoluminescence; PBS, phosphate-buffered saline.
and fluorescent spectra of GSH-QDs and PEG/GSH-QDs indicate that PEG modification have minor influence on the optical properties of QDs (Figure 2B and C). While in agarose gel electrophoresis, the migration rate of PEG/GSH-QDs is decreased due to the PEG derivative conjugation on GSH-QDs (Figure 2D). In addition, the obtained colloidal solution of PEG/GSH-QDs remained stable in the pH buffer and crosslinker chemical solution, and displayed a compact hydrodynamic diameter of 15.1±2.1 nm (Figure S4).

CRP-specific monoclonal antibodies were coupled with the surface PEG layer of QDs via carboxyl and amine groups using carbodiimide chemistry. Successful conjugation of antibodies was evidenced using the agarose gel electrophoresis technique (Figure 2D). The addition of CRP antibody with large molecular weight and slightly positive charge resulted in hindered QD-antibody conjugate migration in gel electrophoresis.

**Effect of QDs PEGylation on the performance of QDs-immunofiltration assay**

The immunoassay performance of QD conjugates obtained from GSH-QDs and PEG/GSH-QDs was compared to elucidate the effect of PEG layer on the performance of IFA. Fluorescent images of IFA pads tested using GSH-QDs and PEG/GSH-QDs are presented in Figure 3. There was a strong non-specific binding between QDs conjugates made of GSH-QDs when compared with that made of PEG/GSH-QDs. As shown in Figure S5, this non-specific binding was obvious even though there was no capturing antibody immobilized on the surface of NC membrane. Whereas, the addition of PEG to QDs helped largely to suppress most of the non-specific interactions in the tested CRP concentration range of 0–25 mg/L. Moreover, immunoassays using PEG/GSH-QDs conjugates are more sensitive and have reliable test results.

**Optimization of experimental conditions**

In addition to QDs surface chemistry, many other factors are expected to influence the detection sensitivity and dynamic range of the developed assay, including the concentration of the QD-antibody probe, the dilution of the target sample, and these factors need to be optimized. The influence of...
QD conjugate concentrations on the assay performance was investigated. Different concentrations of QD conjugates were tested for detection along with both negative and positive control samples. As shown in Figure 4, both fluorescent images and quantitative test results indicated that high concentrations of QD conjugates (≥40 nM) resulted in an intense background fluorescence signals. In contrast, concentrations of QD probes that are too low can eventually reduce the fluorescent signal responses as observed on the antibody spot tested with positive samples. Therefore, the signal/noise (S/N) ratio was calculated and optimized based on the fluorescence intensity ratio of positive to negative samples. As shown in the lower inset of Figure 4B, the optimum S/N ratio can be achieved when the concentration of QD conjugates is approximately 20 nM.

Our test is an immunoassay and its dynamic range of detection is largely expected to be affected by the dilution of original samples. We evaluated the dynamics of the assay and the extent of high-dose hook effect using different dilutions of samples. For this part of the process, 5 µL of serum samples were diluted to 81- and 161-fold using 400 µL and 800 µL of QD conjugates, respectively, and 120 µL aliquot was loaded on the IFA pad for detection (Figure 5). As shown in Figure 6, the dynamic range of the assay was extend from 1.0 to 200 mg/L with 161-fold dilution of serum samples.

At the same time, the sensitivity of the assay was also slightly reduced because of the increased dilution of samples in the QD conjugate solution. According to clinical reference concentration range of CRP in serum (0.5–200 mg/L), we selected 800 µL of QD conjugate for further analysis.

The analytical performance and sensitivity of QDs-immunofiltration assay

Under optimal experimental conditions, the analytical performance of the assay was evaluated with standard CRP samples. Each sample was measured in triplicate and the average results are shown in Figure 7A and B. For the qualitative detection of CRP, Figure 7A displays the fluorescent images of the testing spot on the immunofiltration pad after the performance of assay. The results can be clearly and easily observed with the naked eye under UV lamp illumination. This is mainly owing to the high fluorescence intensity and photostability of QDs. Also, the ascending increase in the concentration of CRP causes more QD conjugates to be captured on the test spot and directly intensifies the fluorescence signals generated from the test spot. Interestingly, the fluorescence intensity of the test spot in the negative CRP samples is very weak and can be easily distinguished from the positive sample even at very low CRP concentration of 0.19 mg/L, which can be considered as

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**Figure 5** Fluorescent images (upper) and spectra (lower) of immunofiltration pad spot tested with a series of CRP samples.

**Notes:** 5 µL of CRP samples in serum respectively added to 400 µL (A) and 800 µL (B) of 20 nM QD conjugates solution, then 120 µL aliquot was loaded into IFa pad.

**Abbreviations:** QD, quantum dot; IFA, immunofiltration assay; CRP, C-reactive protein; PL, photoluminescence.
the limit of detection (LOD) for visual qualitative detection using our assay. On the other hand, the test results were also quantified by an optical fiber spectroscopy under excitation with a 405 nm laser. The standard curve for this QDs-based IFA was constructed by drawing the values of fluorescence spectra peak area against the concentration of CRP (Figure 7B). From this figure, the LOD of our assay using fiber spectroscopy can be 0.79 mg/L, when LOD is defined as the concentration corresponding to a signal that is 3 times SD (standard deviation) above the negative samples. The total results are cumulatively confirming that the developed test can be potentially considered for the rapid qualitative and quantitative testing of CRP serum samples in 5 minutes, depending on the conditions and requirements.

Application of QDs-immunofiltration assay for detecting CRP in human serum

Finally, 50 clinical serum samples were collected to evaluate the developed method for field detection of CRP in human serum. All samples were parallel tested by a standard assay of CRP using a commercial latex agglutination kit, and our optimized QDs-IFA. Linear regression analysis revealed good correlation between our proposed method and this widely used commercial immunoassay kit. The equation of the regression curve was $y=0.9169x + 2.3095$, and correlation coefficient was $r=0.9805$, which implied that our QD-base assay can be applied for clinical determination of CRP in human serum.

Discussion

CRP is known to be synthesized by the liver and frequently secreted into plasma in response to different levels of inflammations, various types of microbial infections, and cancer. The change of CRP concentration in blood serum is frequently correlated to different health conditions. Thus, CRP is currently gaining more importance as a differential

Figure 6 The quantitative dynamic range of the developed QDs-based immunofiltration assay using different volume of QDs conjugates.

Abbreviations: QD, quantum dot; CRP, C-reactive protein; PL, photoluminescence.

Figure 7 Application of the developed QD-based immunofiltration assay for CRP detection.

Notes: (A) Typical fluorescent images of detection results carried out with twofold serially diluted CRP calibrator and a negative control (0 mg/L CRP serum sample) 5 µL of CRP samples in serum respectively added to 800 µL of 20 nM QD conjugates solution, then 120 µL aliquot was loaded into IFA pad. (B) The calibration curve of the quantitative detection by the developed QD-based immunofiltration assay. (C) Correlation between the results of QD-based immunofiltration assay and latex enhanced immune-agglutination assay for 50 human serum samples.

Abbreviations: QD, quantum dot; IFA, immunofiltration assay; CRP, C-reactive protein; PL, photoluminescence.
biomarker in clinical diagnosis. However, this potential is not yet fully realized in field applications, mainly due to the absence of efficient and sensitive assays for the quantitative in field detection of CRP.

We have demonstrated quantitative and sensitive detection of CRP in serum using QD-antibody conjugates based IFA. Following our glutathione and PEG coating approach, the prepared QD-antibody conjugates have negligible non-specific binding and high stability against aggregation which are especially favorable for quantitative protein detection using IFAs.

It is possible to envision that the non-specific binding in IFAs is mainly caused by the hydrophobic and electrostatic interactions taking place between the loaded QD-antibody conjugates and NC membrane. Surface modification of QDs with PEG can shield the negative charge of both the core QD nanocrystals and loaded antibodies. This will directly change the electrical double layer surrounding the QDs and increase the total hydrophilicity of QDs-antibody labeling conjugates. This will subsequently reduce the possibility of hydrophobic and electrostatic binding of labeling conjugates with NC membrane, allowing lower background signal and better detection performance and sensitivity. This was further confirmed by studying the effect of surface chemistry on the detection sensitivity of IFA. IFA employing PEGylated GSH-QDs conjugates as labels demonstrated higher detection sensitivity than that rely on GSH-QDs conjugates. In addition, the presence of PEG layer facilitates the efficient loading and conjugation of antibodies to the surface of QDs without significant reduction in the optical properties of the fluorescent core usually described by our peers. This can also be attributed to the enhanced hydrophilicity and reduced steric hindrance caused by PEG, which prevent the nanoparticles aggregation in the antibody conjugation process and increase the stability of final conjugates. Moreover, the flexible PEG motif on the surface of nanoparticles could maintain the natural conformation of the antibody, and thus improve the affinity of the QD-antibody conjugates.

Through employing these rationally designed PEG/GSH-QDs antibody conjugates in IFA, we were able to detect CRP levels ranging from 0.79 to 200 mg/L. The test results can be checked with the naked eye under UV illumination for qualitative detection in emergence conditions, while quantitative results are able to be performed using an optical fiber spectroscopy. The spectra obtained from the test spot can be eventually analyzed for more differentials and can potentially be used for multiplexed detection using multicolor QDs. Interestingly, this dual mode of detection could support a wider range of emergency and central lab applications, and further indicates the rationality of our developed QD-based IFAs.

Compared with the previously reported QDs-based assay and traditional ELISA method, the advantages and disadvantages are briefly summarized in Table 1. Even though the developed QD-based IFA test is not as sensitive as microplate and magnetic beads based assays, the sensitivity and dynamic range of this method provide satisfactory analytical characteristics for clinical CRP detection. Importantly, proposed QD-based assay has simpler and faster testing procedures, and requires less sample volume, which is more suitable for on-site testing in an emergency situation. The quantitative point-of-care testing kit may allow more accurate targeting of appropriate therapy and result in considerable cost saving.

**Conclusion**

A QDs-based IFA was successfully developed for rapid (within 5 minutes), sensitive, and quantitative detection of CRP in human serum using small sample volumes of 5 µL. Using surface PEGylation of QDs, the immunoassay performance of QDs-antibody conjugate was largely enhanced, and under optimal experimental conditions, the proposed QDs-based IFA was able to qualitatively detect concentrations as low as 0.19 mg/L CRP in the serum sample. The detection range of this assay is 0.09–200 mg/L with a quantitative detection sensitivity of 0.79 mg/L using serum samples. Moreover, the obtained results were well correlated (r=0.9805) with the data detected by a commercial latex

| Methods | QD-based IFA | QD-based lateral flow immunoassay | QD labeled microplate immunoassay | QD labeled magnetic beads immunoassay | Traditional ELISA |
|---------|-------------|---------------------------------|-----------------------------------|--------------------------------------|-------------------|
| Detection limit | 0.79 mg/L | 0.25 mg/L | 0.06 µg/L | 1 µg/L | 0.16 mg/L |
| Detection time | <5 minutes | 15 minutes | 1–1.5 hours | 1–3 hours | Over 12 hours |
| Sample required | 5 µL | 10 µL | 100 µL | 50–100 µL | 5 µL |
| Test procedure | One step | One step | Two steps | Three steps | Three steps |

Abbreviations: QD, quantum dot; IFA, immunofiltration assay; CRP, C-reactive protein; ELISA, enzyme-linked immunosorbent assay.
agglutination assay, which confirms the potentiality of our QDs-based IFA for the development of rapid, quantitative point-of-care diagnostics.

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Disclosure
The authors report no conflicts of interest in this work.

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Supplementary materials

Figure S1 UV-Vis absorbance and fluorescent spectra of CdSe core and CdSe/ZnSe/ZnS core/shell/shell QDs.
Abbreviations: UV, ultraviolet; UV-Vis, ultraviolet-visible; PL, photoluminescence; QDs, quantum dots.

Figure S2 TEM graphs of CdSe core (A) and CdSe/ZnSe/ZnS core/shell/shell (B) QDs.
Abbreviations: TEM, transmission electron microscopy; QDs, quantum dots.

Figure S3 Dynamic light scattering analysis of GSH-QDs.
Abbreviations: QDs, quantum dots; GSH, glutathione.
Figure S4 Dynamic light scattering analysis of PEG/GSH-QDs.

Abbreviations: QDs, quantum dots; GSH, glutathione; PEG, polyethylene glycol.

Figure S5 Comparison of non-specific binding of GSH-QDs and PEG/GSH-QDs conjugates with NC membrane without capturing antibody.

Abbreviations: QD, quantum dot; GSH, glutathione; PEG, polyethylene glycol; NC, nitrocellulose.