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Article

Colorimetric and Reverse Fluorescence Dual-Signal Readout Immunochromatographic Assay for the Sensitive Determination of Sibutramine

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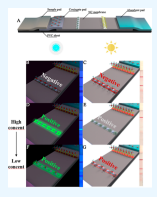
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ABSTRACT: Later flow immunochromatographic assay has been widely used in clinical, environmental, and other diagnostic applications owing to its high sensitivity and throughput. However, most immunoassays operate in the "turn-off" mode for detecting targets of low molecular weight. The signal intensity decreases as the analyte concentration increases, which poses a challenge for achieving ultrasensitive detection at low concentrations and is counterintuitive to new users. In this work, a fluorometric immunochromatographic assay (FICA) is developed to simultaneously read "turn-on" fluorescent and "turn-off" colorimetric signals, where ZnCdSe/ZnS quantum dots act as fluorescence donors and gold nanoparticles (AuNPs) act as quenchers. The fluorescent signal (excitation/emission wavelengths of 365/525 nm) is positively correlated with analytes' concentration. Taking sibutramine (SBT) as the analysis target, the visual limit of detection for SBT reached 3.9 ng/mL, and the limit of Quantitation was 5.0 ng/mg in spiked samples. The developed FICA achieves a high sensitivity in SBT detection, which is much lower than that of the colloidal gold-based immunochromatographic assay. This dual-function detection mode has great



potential to be used as a rapid on-site semiquantitative method, providing an alternative mode for the determination of low levels of target analytes.

INTRODUCTION

Sibutramine (SBT) was approved as an antiobesity drug by the U.S. Food and Drug Administration (FDA) in 1997 and was also regarded as an effective drug for treating obesity in European countries. 1,2 As a serotonin-norepinephrine reuptake inhibitor, its weight loss mechanism is achieved by inhibiting and activating the satiety center and increasing energy expenditure.^{3,4} However, further studies have shown that ingestion of SBT increases cardiovascular and nonfatal stroke risk, a dangerous side effect that has banned its use in many countries.^{5,6} Therefore, in 2010, the FDA, the European Commission, and the Ministry of Health of the People's Republic of China banned SBT and recommended that its production, sale, and use be stopped.^{5,7} However, due to its tremendous weight loss effects, some weight loss supplements still illegally add SBT to increase weight loss. SBT has been detected in many so-called antiobesity and natural weight-loss products in the market. This adulteration of products can present a real risk to consumers and damage the credibility of legitimate products. For both customers and regulators, a rapid, sensitive, affordable, and self-service-accessible approach

At present, instrumental-based approaches for the detection of SBT have undergone substantial development and have achieved extensive utilization; for example, electrospray ionization mass spectrometry (ESI-MS), high-performance

liquid chromatography (HPLC),⁹ gas chromatography (GC),¹⁰ Fourier transform infrared spectroscopy (FTIR),¹¹ and other instrumental methods have been developed. These instrumental analytical methods generally possess high capacity for trace component detection and excellent accuracy, but superior sensitivity comes with a high equipment and maintenance cost, as well as the need for highly trained professionals. In recent years, some innovative analytical approaches offering good sensitivity and convenience were proposed, such as surface-enhanced Raman scattering senor (SERS)¹² and glassy carbon electrode (GCE),^{13,14} but they remain equipment intensive. These methods are inconvenient for testing personnel to screen suspicious samples rapidly under field conditions.

Lateral flow immunochromatographic analysis (LFIA) is a rapid, low-cost, and convenient detection method, which has excellent performance in food safety detection and on-site diagnosis. ^{15,16} Currently, LFIA has been used for the

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qualitative and semiquantitative detection of SBT based on the antigen-antibody biochemical interaction. 17,18 In these systems, a signal reporter of gold nanoparticles offers excellent stability and optical properties while fluorescent quantum dots (QDs) offer high fluorescence quantum yield and tunable emission wavelength. Yet background signal intensity limits ultrasensitive detection. 19,20 For micromolecular analyte detection, current LFIA presents a competitive mode called "turn-off", and there is a negative correlation between the measured signal intensity and target concentration. 21,22 The limit of detection (LOD) of LFIA in "turn-off" mode is defined as the corresponding concentration of analytes causing the apparent signal to completely disappear.²³ For the "turn-off" mode, the intensity of the visualized signal decreases rather than disappears when low concentrations of analyte are present, making it difficult to determine whether a low concentration of analyte is present.

Gold nanoparticles possess large extinction coefficients and wide characteristic absorption peaks in the visible range that overlap with the emission spectrum of fluorescent materials.² Quantum dots (QDs) have outstanding properties such as high quantum yield, tunable photoluminescence emission, high molar extinction coefficients, narrow and symmetric fluorescence spectra, large Stokes shift, high photochemical stability, chemical stability, and other advantages. 25,26 By exploiting the phenomenon of quenching in the presence of spectrally overlapping fluorescence signals from these two materials, they were shown to be excellent quenchers and donors for the preparation of "turn-on" type fluorescence immunochromatographic assays (FICAs). Sheng et al.²⁷ developed a fluorometric lateral flow immunoassay in "turnon" mode to detect tetracycline based on QDs fluorescence quenching by gold nanoparticles. Chen et al. $\overline{^{28}}$ constructed a high fluorescence quenching probe-based reverse fluorescence enhancement lateral flow test strip (rLFTS) to realize the detection of the relevant nucleic acid of Parkinson disease.

In this study, a FICA was constructed based on a combination of QDs as fluorescent signal source and gold nanoparticle as fluorescence quencher. As the SBT content increases, the fluorescence signal gradually increases, and the colorimetric signal gradually weakens. It not only retains the "turn-off" mode of LFIA but also proposes a "turn-on"mode for fluorescence signal interpretation through ultraviolet light (UV) to achieve visual detection of low-concentration SBT.

MATERIALS AND METHODS

Materials and Instruments. Sibutramine hydrochloride, ovalbumin (OVA), 1-ethyl-3-[3-(dimethylamino) propyl] carbodiimide (EDC), and chloroauric acid (HAuCl₄) were purchased from Sigma-Aldrich Chemical Co, Ltd. (St. Louis, MO, USA). Carboxyl functional ZnCdSe/ZnS quantum dots (QDs) (emission maximum at 525 nm) were obtained from Jiayuan Quantum Dots Co, Ltd. (Wuhan, China). Trisodium citrate dihydrate (SC) was purchased from Sinopharm Chemical Reagent Co, Ltd. (Shanghai, China). Anti-SBT monoclonal antibody (mAb) and SBT coating antigen (SBT-OVA) were produced in our laboratory. Goat antimouse antibody was obtained from Solarbio Biotechnology Co, Ltd. (Beijing, China). Enzyme-linked immunosorbent assay (ELISA) PLATE, poly(vinyl chloride) (PVC) sheet, absorbent pad, conjugate pad, sample pad, and nitrocellulose (NC) membrane were purchased from Thermo Fisher Scientific (Waltham, MA, USA). Automatic guillotine cutter (CM4000)

and dispensing platform (XYZ3050) were purchased from BioDot Co, Ltd. (Irvine, CA, USA). The immunochromatographic assay signal reader was obtained from Helmence Instrument Co, Ltd. (Suzhou, China). A portable UV lamp was purchased from LICHEN Scientific Co, Ltd. (Qingdao, China).

Preparation of QDs-OVA. The conjugation of OVA and QDs was performed by previously described methods with slight modifications as described below. ^{29,30} In brief, 100 μ L of carboxy-surface QDs (8 μ mol, dissolved in 50 mmol borate buffer) were diluted with 10 mmol sodium borate (pH 8.6) to a final concentration of 1 μ mol. The pH of the diluted QDs solution was adjusted to 5.5 using a citric acid—sodium citrate buffer (0.1 M). Then, 15 μ L of EDC solution (10 mg/mL) was added to the above solution and stirred for 4 h in the dark. Subsequently, 1.35 mg of OVA was added to continue the reaction for 12 h. Finally, the resulting mixture was centrifuged for 15 min (13,000 rpm, 4 °C), the supernatant was discarded, and the pellet was redissolved in 100 μ L of PB (0.01 M, pH 7.4) and stored at 4 °C.

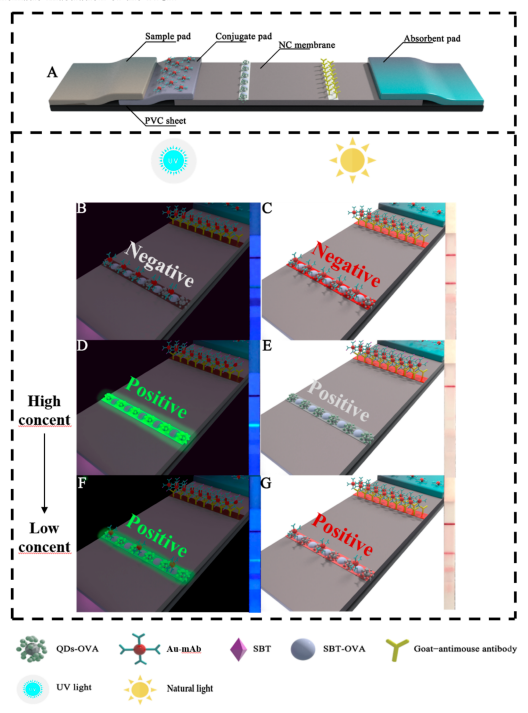
Preparation of AuNPs and Au-mAb Probes. The AuNPs were synthesized on the basis of previous work, 31 and the anti-SBT mAb was obtained previously. 32 0.1 M $\rm K_2CO_3$ was added to the prepared AuNPs solution (1 mL, 0.18 nmol/L) to adjust pH to 7, 8, 9, 10, 11, and 12, then 2.5, 5, 10, and 20 $\mu \rm g$ of anti-SBT mAb was added. Afterward, the reaction mixture was incubated for 1 h at room temperature. To block the uncoupled surface of AuNPs, 10 $\mu \rm L$ volumes of 20% BSA solution and 20% PEG-20,000 were added and incubated for 30 min to avoid nonspecific adsorption. Then, the solution was centrifuged for 30 min (10,000 rpm, 4 °C). Finally, the precipitate was resuspended in 1 mL of ultrapure water, and the Au-mAb probe was obtained.

Fabrication of FICA Strip. The FICA was developed by referring to the work reported and processed as following. ²⁷ A mixture of 50 μ L of QDs-OVA (450 nmol/L) and 50 μ L of SBT-OVA (0.25 mg/mL) were sprayed on the reaction membrane at a dose of 0.6 μ L/cm as the T line. Analogously, goat antimouse antibody (1 mg/mL) was immobilized on NC membrane as C line with a 0.6 μ L/cm dosage. After 4 h of drying at 37 °C, all of the parts mentioned above were cut into 3 mm strips, stored at room temperature, and kept under designation

Optimization. To optimize the pH of the synthesis of the Au-mAb probe, the pH of the AuNP solutions were adjusted to 7, 8, 9, 10, 11, and 12. To optimize the amount of immobilized antibody, 2.5, 5, 7.5, and 10 μ g of anti-SBT mAb was added dropwise to 1 mL of AuNPs solution at final concentrations of 2.5, 5, 7.5, and 10 μ g/mL, respectively. To optimize the SBT-OVA concentration, the T line was sprayed with SBT-OVA at concentrations of 0.1, 0.25, 0.5, 0.75, and 1.0 mg/mL with a spraying speed of 0.6 μ L/cm.

In the above-mentioned experiments, the addition of PBS (0.01 M) buffer containing 0 and 100 ng/mL SBT was used to optimize the key parameters. The pixels per inch (Δ PPI) of colorimetric signal on the T line was monitored as the indicator of optimization. Δ PPI gives the difference value of the T line signal between negative (0.01 M PBS) and positive (100 ng/mL SBT in 0.01 M PBS) controls (Δ PPI = PPI_{negative} – PPI_{positive}). On the basis of the optimized conditions, the optimal quenching efficiency between donors and receptors is established. Final concentrations of 300, 450, and 600 nmol/L QDs-OVA mixed with SBT-OVA were sprayed on the T line,

Scheme 1. Schematic Illustration of the FICA^a



 $^{a}(A)$ construction of FICA strip, negative sample detection under (B) UV light and (C) natural light, positive sample detection under (D, F) UV light and (E, G) natural light.

and 0, 0.25, 0.5, 0.75, 1.0, and 2.0 μ L of Au-mAb was dried on conjugation pad. Then 0, 0.98, 1.95, 3.90, 7.81, 15.63, 31.25, 62.5, 125, and 250 ng/mg of SBT-containing PBS (0.01 M, pH 7.4) buffer was added for detection according to each of the above-mentioned protocols. The best correlation was found, responding to the lowest concentration of SBT. All experiments were performed in triplicate.

Method Validation. Two types of dietary fiber soft capsules and weight loss supplement tablets in e-commerce stores were randomly purchased as samples and were

confirmed by free LC–MS/MS. For analysis, accurately weighed 895.3 mg of ground and homogenized capsule contents or tablets were spiked with SBT standards to final concentrations of 2, 5, 100, and 250 ng/mg. The samples were dissolved in 30 mL of methanol, vortexed for 10 min, subjected to ultrasonic extraction for 15 min (at a water bath temperature of 40 °C), and then centrifuged for 5 min (4000 rpm, 25 °C). The supernatant was collected and dried under a N_2 stream, then redissolved in 50 mL of PBS containing 5% methanol, vortexed for 30 s, and filtered

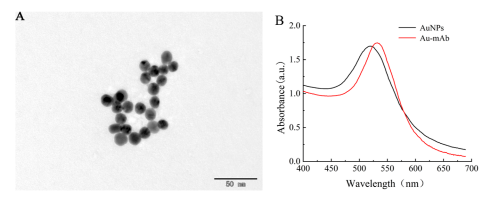


Figure 1. Characterization of AuNPs and Au-mAb. (A) TEM images of AuNPs. (B) UV-vis absorption spectra of AuNPs and Au-mAb.

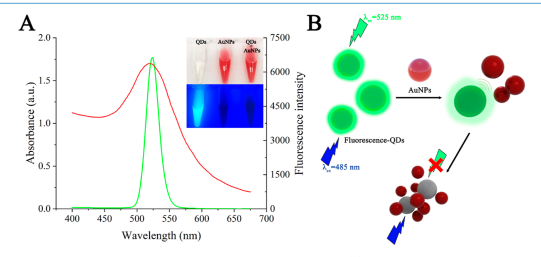


Figure 2. (A) UV—vis absorption spectrum of AuNPs and fluorescence spectrum of QDs. (B) Process representation of the fluorescence quenching mechanism by AuNPs.

through a 0.22 μ m Teflon membrane. The extracts were further diluted 16-fold in PBS containing 5% methanol to suppress the interference of matrix and then ready for the analysis of FITC and LC–MS/MS.

Performance of FICA. As illustrated in Scheme 1A, QDs were labeled on the OVA as fluorescent emitters, and the conjugate of QDs-OVA was sprayed evenly on the T line. The mAb-labeled AuNPs as probes and quenchers were precoated on the conjugate pad and kept dry. In the absence of SBT, AumAb and SBT-OVA formed a sandwich structure, resulting in a close adjacency of AuNPs and QDs. Then, concomitant arresting fluorescence quenching occurred on the T line (Scheme 1B), and no fluorescence signal would be observed under UV light. Meanwhile, AuNPs aggregation induced color emergence, which allowed naked eye readout (Scheme 1C). In contrast, when the sample contained a high concentration of SBT, the Au-mAb complex was unable to bind with SBT-OVA on the T line, and the fluorescence was restored; thus, a strong green fluorescence signal was observed (Scheme 1D), but no colorimetric signal was detected (Scheme 1E). However, a relatively low target concentration might cause a visible weak colorimetric signal to appear (Scheme 1G), resulting in a falsenegative judgment by the naked eye. Under this circumstances, a green fluorescence signal still appeared in UV light (Scheme 1F), which ensures the identification of low concentrations of SBT. Thus, the developed FICA with simultaneous "turn-on" and "turn-off" modes achieved a dual-signal readout,

supporting ultrasensitive semiquantitative and rapid largescale screening.

■ RESULTS AND DISCUSSION

Characterization of AuNPs and Au-mAb. The morphology and size of AuNPs are illustrated via the transmission electron microscopy (TEM) image in Figure 1A. It revealed that the size of AuNPs was 20 ± 0.7 nm with a uniform dispersion. In the UV-vis absorption spectrum (Figure 1B), the maximum absorption peak of Au NPs was 519 nm, the narrow peak shape could be observed, indicating a stable state of dispersion, and the small red shift was attributed to the coupling of antibody with AuNPs.

Reaction Mechanism Exploration. AuNPs were highly efficient fluorescence quenchers of molecular excitation energy (up to 99.8%) over a long distance (about 40 nm) because of their extremely strong absorption. The ZnCdSe/ZnS QDs were chosen as the donors to construct luminophore-AuNPs composites due to their slower diffusion of excitons to dissociative defects, leading to a relatively brighter emission. Figure 2A showed the absorption spectrum of the AuNPs (surface plasmon maximum at 519 nm), which had a large degree of overlap with the emission spectrum of the QDs (emission maximum at 525 nm). Figure 2B illustrates that the fluorescence was gradually extinguished with the narrowing distance between QDs and AuNPs. Fluorescence quenching modes include the "on—off" method, which can be summarized as follows: (a) by reducing the light emission

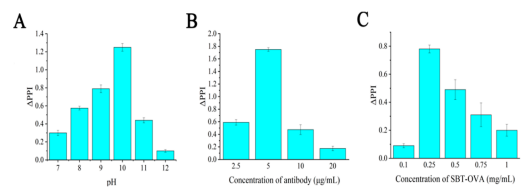


Figure 3. Optimization of key parameters. Δ PPI of the T line at different (A) pH values, (B) antibody concentrations, and (C) SBT-OVA concentration.

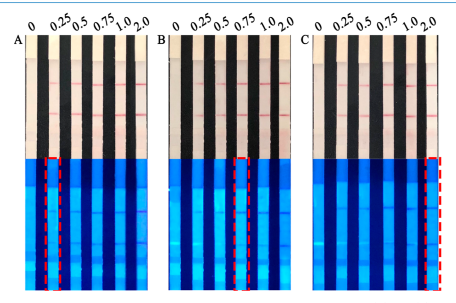


Figure 4. Fluorescence quenching relationship between 0, 0.25, 0.5, 0.75, 1.0, and 2.0 μ L of Au-mAb and (A) 300, (B) 450, and (C) 600 nmol/L of QDs-OVA.

intensity of fluorescent nanoprobes. (b) "On-off-on" approach by breaking the bond between the fluorescent nanoprobe and the analyte. (c) "Postfunctionalization" method, chemically modifying the surface of prepared fluorescent nanoparticles to introduce specific receptor molecules or other molecular functional groups so that the probe has specific biological recognition or chemical response characteristics.³⁶ Obviously, FICA in this study belongs to the third mode. In this detection process, the vast majority of quenching mechanisms relies on fluorescence resonance energy transfer (FRET), 30,37 inner filter effect (IFE), 37,38 and nanometal surface energy transfer (NEST).³⁸ In this study, the emission spectrum of the QDs overlapped with the UV absorption spectrum of the AuNPs, which met the requirements of fluorescent material (donor) and quencher (acceptor) in FRET and IFE. The prerequisite for FRET is that the distance between fluorescence donor and acceptor needs to be less than 10 nm.³⁹ The mixture of QDs-OVA (donor) and SBT-OVA was sprayed on the T line and the proximity between AuNPs (acceptor) and QDs-OVA relied on the specific binding of antigen and antibody, but it is hard to ensure the ideal distance between QDs and AuNP to generate FRET.²⁷ IFE becomes the crucial factor for fluorescence quenching. IFE needs to meet the requirements of only spectral overlap for the quenching phenomenon to

occur. It does not have strict requirements on the distance between the donor and the acceptor, which becomes a key factor leading to fluorescence quenching in this study.³⁶ Some research suggested that the primary cause of fluorescence quenching in LFIA was attributed to the IFE and the selfabsorption of fluorescence. 40,41 Besides, Nebu et al. 38 and Yao et al.³⁴ considered that the system followed NEST, in which interaction between donor and acceptor was generated through an electromagnetic field by the fluorescent material dipole and the conduction band charge of AuNPs. Here, either the IFE or NEST could lead to fluorescence quenching. Therefore, the developed FICA with a dual-signal readout is anticipated to offer enhanced sensitivity compared to that of the single-signal-based AuNPs IFCA, serving as an on-site rapid diagnostic strategy for the accurate detection of trace analytes.

Optimization. Before matching the QD fluorescence emitter with the amount of AuNPs quencher, we optimized key parameters of the FICA assay to achieve the best detection performance. In order to improve the detection sensitivity of SBT, we employed Δ PPI as our analytical metric for result assessment. The magnitude of fluorescent restoration is directly related to the amount of Au-mAb combined on the T line, which is positively correlated with the amount of

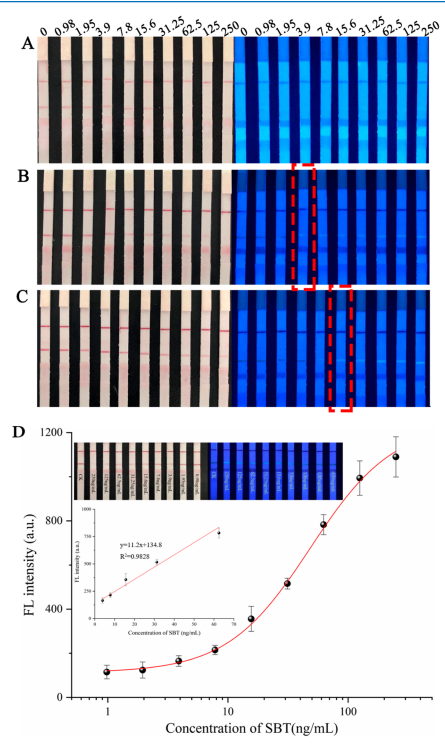


Figure 5. Detection of SBT in PBS at 0, 0.98, 1.95, 3.9, 7.8, 15.6, 31.25, 62.5, 125, and 250 ng/mL for (A) 0.25 μ L of Au-mAb and 300 nmol/L QDs-OVA, (B) 0.75 μ L of Au-mAb and 450 nmol/L QDs-OVA, (C) 2 μ L of Au-mAb and 600 nmol/L QDs-OVA, and (D) standard curve for quantitative analysis of SBT through changes in T-line fluorescence intensity. The inset shows the linear relationship in the low concentration range and the actual plot.

analytes in the sample. Therefore, the higher the Δ PPI, the higher the detection sensitivity.

Figure 3A shows the effect of the pH on PPI. Δ PPI reached its maximum value at pH = 10 and then decreased with the increase of pH, indicating that pH = 10 was the optimum pH condition for the coupling process. Figure 3B shows the effect of the antibody addition on Δ PPI. Δ PPI peaked when 5 μ g of antibody was added to 1 mL of AuNPs solution for probe

coupling. As the amount of antibody added increased from 5 to 20 μ g, Δ PPI decreased due to supersaturation of the antibody adsorbed to the probe on the T-line. Therefore, 5 μ g was selected as the optimal antibody addition amount. Figure 3C shows the effect of SBT-OVA on Δ PPI. As the concentration increased from 0.1 to 1 mg/mL, Δ PPI peaked and gradually decreased. In order to obtain a higher Δ PPI, the optimum

spraying concentration of SBT-OVA on the T line was 0.25 mg/mL (spraying speed 0.6 μ L/cm).

Visual Sensitivity Assessment of the FICA. The visual limit of detection (vLOD) of FICA in this work was defined as the lowest concentration of SBT at which the colorimetric signal began to decrease under natural light, and at the same time a green fluorescent signal appeared under 385 nm UV light. It was the key to finding an appropriate match between QDs-OVA and Au-mAb to make the FICA sensitively respond to the change of analytes amounts, and the fluorescence could be completely quenched in the negative condition. Under optimized conditions, 0, 0.25, 0.5, 0.75, 1.0, and 2.0 μ L volumes of Au-mAb (0.18 nmol/L) probes were dried on the conjugate pad, and optimized SBT-OVA (0.25 mg/mL, spraying speed 0.6 μ L/cm) was mixed with final concentrations of 300, 450, and 600 nmol/L QDs-OVA and sprayed on the T line. Through detection of blank samples (PBS), the minimum volume quenching fluorescence was deemed to be the optimal addition of Au-mAb. As shown in Figure 4, 0.25, 0.75, and 2.0 μ L of Au-mAb (0.18 nmol/L) were the minimum volumes for 300, 450, and 600 nmol/L QDs-OVA (0.25 mg/ mL, spraying speed 0.6 μ L/cm) fluorescence quenching, respectively. Then, a series of concentrations of SBT standard solution (0, 0.98, 1.95, 3.9, 7.8, 15.6, 31.25, 62.5, 125, and 250 ng/mL) were detected in the 0.25 μ L-300 nmol/L (Figure 4A), 0.75 μ L-450 nmol/L (Figure 4B), and 2.0 μ L-600 nmol/ L schemes (Figure 4C).

The results are shown in Figure 5, and as the concentration of SBT increases, the fluorescence intensity on the T line increases gradually. However, Figure 5A showed that the green fluorescence signal is indecipherable to recognize for 300 nmol/L QDs-OVA (0.25 mg/mL, spraying speed 0.6 μ L/cm) with related 0.25 μ L of Au-mAb (0.18 nmol/L). As shown in Figure 5B, when the concentration of SBT reached 3.9 ng/mL under the conditions of 0.75 μ L Au-mAb and 450 nmol/L QDs-OVA, a bright green fluorescence signal was observed. As the SBT concentration increases to 250 ng/mL, the fluorescence signal on the T line is fully turned on, and the colorimetric signal is turned off. By comparison, Figure 5C shows that when the concentration of SBT reaches 15.6 ng/ mL in the 2 μ L of Au-mAb and 600 nmol/L QDs-OVA solutions, the fluorescence signal turns on. Therefore, 0.75 μL of Au-mAb and 450 nmol/L QDs-OVA were selected as the optimal experimental conditions.

Under the previous optimal experimental protocol, the sensitivity of the developed FICA was evaluated by a series of concentrations of SBT (0, 0.98, 1.95, 3.9, 7.8, 15.6, 31.25, 62.5, 125, and 250 ng/mL) in PBS (0.01 M, pH 7.4). Based on the changes in SBT concentration and T-line fluorescence intensity, a quantitative analysis curve was established, as shown in Figure 5D. The regression equation is y = 11.2x + 134.8 ($R^2 = 0.9829$), and the linear range is 3.9-62.5 ng/mL, LOD (defined as the T-line fluorescence intensity of the negative sample plus the lowest SBT concentration of 3 times the standard deviation) was 1.128 ng/mL, and when the concentration of SBT was 3.9 ng/mL, an obvious fluorescence signal was observed. Therefore, the vLOD was set to 3.9 ng/mL

For comparison, previous reports for the determination of the SBT are summarized in Table 1. Among these methods, electrochemical sensors and SERS display extremely high sensitivity and can provide quantitative data. But these methods require relatively complicated operations, such as

Table 1. Overview of Recently Reported Methods for the Detection of SBT

method	qualitative/quantitative	lowest LOD	references
LFIA based on gold nanoparticle	qualitative	500 ng/mL	18
FICA based on visual up conversion nanoparticle	qualitative	20 ng/mL	17
colorimetric detection based on aggregation of AuNPs	quantitative	1.15 μ mol/L	42
paper device for distance-based	quantitative	0.22 mmol/L	1
SERS	quantitative	10^{-9} mol/L	7
carbon screen-printed electrode	quantitative	$0.3~\mu\mathrm{mol/L}$	14
FICA based on "turn-on" mode	qualitative	3.9 ng/mL	this work

the process of adsorptive stripping pulse differential voltammetry (AdSDPV), preparation of the SERS substrate, and requirement of a device to record. By contrast, the FICA developed in this study exhibited a better LOD with other existing qualitative detection systems and even with other quantitative analysis procedures.

Specificity of the FICA. To investigate the specificity of the FICA, the bioactive metabolites of SBT, N-desmethylsibutramine (M_1) and N,N-bidesmethylsibutramine (M_2) were tested by the FICA strips. Besides, sildenafil (SDF) was also regarded as a potential interfering substance because of its unexpected addition to slimming supplements. As shown in Figure S2, no fluorescence signal could be observed even when M2 and SDF standards were added at a concentration of 1 μ g/mL. Cross reactivity has been observed for the detection of M_1 and SBT standards at 10 ng/mL. We further evaluated the FICA developed for the detection of M1. The concentration of M1 was detected to be the same as that of SBT, and a standard curve was established (Figure S3). The regression equation was y = 11.8x + 160.8 ($R^2 = 0.9449$), the LOD was 2.087 ng/mL, and the vLOD was 7.8 ng/mL.

Method Validation. All sample preparation and extraction were as described previously. To remove the matrix effects, the finial extract was diluted 2, 4, 8, and 16-fold with 0.01 M PBS. Without SBT, the interference of the matrix was analyzed by the FICA assay using the pretreated blank sample, and the least dilution multiple could be ascertained when the fluorescence on the T line was quenched completely. The results of Figure S4 showed that the samples (both of the matrixes, dietary fiber soft gel, and tablet) needed to be diluted 16-fold to eliminate the matrix effects. Then, the sensitivity was evaluated by analyzing the two matrixes spiked at the different concentrations of SBT (0, 2, 5, 100, and 250 ng/mg) in the actual samples. Established LC-MS/MS for the detection of SBT was used to verify the accuracy of the FICA, with three replicate analyses in triplicate (Figures S5, S6). The lowest spiked concentration of 5 ng/mg was detected in dietary fiber soft gels and tablets, respectively. Qualitatively, all results of FICA and LC-MS/MS methods were consistent, and the corresponding quantitative results are shown in Table 2, which demonstrated the developed FICA possessing acceptable reliability and potential for future practical application.

Table 2. Verification of the Spiked Slimming Supplements Using LC-MS/MS and the Developed FICA in This Study

Sample	Spiked (ng/mg)	LC-MS/MS detected (ng/mg)	Recovery (%)	RSD (%)	FICA- judgement ^a
Dietary fiber soft gel	0	ND^b	ND	ND	
	2	ND	ND	ND	
	5	4.65	93	6.38	++
	100	94.66	94.66	6.01	++
	250	234.45	93.8	4.74	+++++
Tablet	0	ND	ND	ND	
	2	ND	ND	ND	
	5	5.21	104.2	5.04	++
	100	96.73	96.73	4.85	+++
	250	234.11	93.64	6.12	+++++

"Visual assessment: "— (red)" represents the negative result under natural light, "+ (red)" represents the positive result under natural light, "+ (green)" represents the negative result under UV light, and "+ (green)" represents the positive result under UV light. ^bNot detected.

CONCLUSIONS

In this work, based on the fluorescence quenching effect of AuNPs on QD-OVA, a dual-signal FICA combining "turn-on" and "turn-off" modes is proposed for ultrasensitive on-site detection of SBT. Under the optimal detection condition, the lowest visual LOD and LOQ of the developed FICA for SBT was 3.9 ng/mL, 5.0 ng/mg. Its sensitivity was much higher than that of the traditional LFIA based on the "turn-off" mode. The developed FICA detection system has the advantages of fast detection speed, low cost, and high sensitivity, and the obtained results can be semiquantitative to the naked eye, demonstrating the potential application for the determination of sibutramine in weight loss supplements. However, before its commercial application, it requires validation for the stability and reproducibility of large-scale synthesis. In summary, the developed FICA detection system is expected to be a method suitable for rapid on-site screening.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.3c09050.

Chemical structure and synthetic route of hapten-SBT; specificity analysis of FICA; chemical structure of SBT, SDF, M1, and M2; elimination of matrix effects; and detection of SBT in spiked samples (PDF)

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Notes

The authors declare no competing financial interest.

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