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Noble metal nanomaterial-based aptasensors for microbial toxin detection

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Abstract

Microbial toxins generated by bacteria, fungi and algae cause serious food-safety problems due to the frequent contamination of foodstuffs and their poisonous nature. Becoming acquainted with the contamination condition of foodstuffs is highly dependent on developing sensitive, specific, and accurate methods for targeting microbial toxins. Aptamers, obtained from systematic evolution of ligands by exponential enrichment (SELEX), have significant advantages for microbial toxin analysis, such as small size, reproducible chemical synthesis, and modification, as well as high binding affinity, specificity, and stability. Besides, aptamers have a predictable structure and can be tailored using biomolecular tools (e.g., ligase, endonuclease, exonuclease, polymerase, and so on), which is conducive to the development of flexible and variable amplification methods. Recent studies revealed that the combination of aptamers and noble metal nanomaterials offers unprecedented opportunities for microbial toxin detection. Noble metal nanomaterials with outstanding physical and chemical properties facilitate the detection process and improve the sensitivity and specificity. In this review, we discuss current progress in the development of various noble metal nanomaterial-based aptasensors for microbial toxin detection. These noble metal nanomaterials include gold nanoparticles, gold nanorods, gold nanoclusters, silver nanoparticles, silver nanoclusters, and bimetallic nanomaterials. Aptasensors based on noble metal nanomaterials exhibiting high selectivity and sensitivity represent a promising tool for microbial toxin detection.

Keywords: Algal toxin, Aptamer, Bacterial toxin, Mycotoxin, Noble metal nanomaterial

1. Introduction

Food contaminations caused by microbial toxins (such as bacterial toxin, mycotoxin, and algal toxin) are becoming serious problems to human health [1]. Among them, bacterial toxins have the highest significance with respect to public health. Bacterial toxins are generally classified into endotoxins and exotoxins. The most common bacterial toxins are Staphylococcal toxins, Clostridia botulinum and Clostridium difficile toxins, Bacillus anthracis toxin, and Shiga toxins. Through the diet, bacterial toxins can interact with and damage host cells by disturbing the signaling pathways and structural integrity of the host cells, leading to various types of disease symptoms, such as nausea, vomiting, and diarrhea. Mycotoxins, being secondary toxic metabolites, are primarily derived from fungi, particularly Aspergillus, Penicillium, Fusarium, and Alternaria. They include hundreds of chemically toxic compounds, such as aflatoxins (AFs), ochratoxin A (OTA),

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trichothecene mycotoxin (T-2 toxin), zearalenone (ZEN), fumonisins (F), patulin (PAT), and so on. Toxicological research studies revealed that a long-lasting exposure to a low dose of mycotoxin is capable of causing strong carcinogenic, teratogenic, and mutagenic effects. Algal toxins are generated by algae and are found in both marine and freshwater environments. The consumption of seafood or water contaminated with algal toxins can cause serious disease symptoms, including tingling, nausea, vomiting, paralysis, and diarrhea. To anticipate potential risks and ensure food safety, it is highly demanded and crucial to develop selective, sensitive, rapid, low-cost and on-site detection techniques for microbial toxins.

To ensure the required detection performance, antibodies have long been the most popular recognition elements used for the binding of target analytes. Unfortunately, antibodies suffer from short shelf-life, high costs of manufacturing, relatively low stability and non-negligible batch-to-batch variations. Aptamers, an emerging class of molecules that rival antibodies have become the focus of attention over the past few decades. They are short, chemically synthesized, single-stranded oligonucleotides (DNA or RNA), possessing unique tertiary structures. Aptamers with high affinity and specificity to identify target molecules can be selected via a reliable and efficient procedure known as systematic evolution of ligands by exponential enrichment (SELEX). Collectively, aptamers are less expensive and tedious to manufacture, more stable in long-term storage and easier to synthesize, manipulate, and functionalize compared with antibodies [2]. Moreover, to ensure highly sensitive detection, noble metal nanomaterials offer an unprecedented potential in terms of signal transduction. Noble metal nanomaterials with unique physical and chemical properties (e.g., surface plasmonic effects or luminescence properties) can markedly improve the sensitivity and specificity for their applications in biosensors. With the integration of aptamers as the recognition elements, these biosensing nanoplatforms allow a promising way for cost-effective, rapid and sensitive on-site detection of microbial toxin. Some of these studies have been reviewed elsewhere, with a focus on the fabrication electrochemical aptasensors for the detection of toxins based on gold nanomaterials, or as subclasses in a more general overview of the nanomaterial-based aptasensors [3,4]. In this review, we will focus on the recent development of noble metal nanomaterial-based aptasensors for microbial toxin detection and discuss some major achievements in this field over the past few years, with the aim of improving their future performances of noble metal nanomaterial-based aptasensors in food safety examination.

2. Noble metal nanomaterial-based aptasensors for microbial toxin detection

Noble metal nanomaterials, such as gold nanoparticles (AuNPs), gold nanorods (AuNRs), gold nanoclusters (AuNCs), silver nanoparticles (AgNPs), silver nanoclusters (AgNCs), and bimetallic nanomaterials, offering fascinating signaling properties have been employed as sensing elements to design novel aptasensors. Other attractive characteristics, such as convenient and controllable synthesis, high biocompatibility, size-dependent optical and electronic properties, and ease of surface modification, also make them extremely suitable for developing various types of aptasensors, including colorimetric, fluorescent, surface-enhanced Raman scattering (SERS), electrochemical, and portable sensors (Table 1). In the following sections, we will introduce representative examples of each of these classes of aptasensors for microbial toxin detection.

2.1. AuNP-based aptasensors

2.1.1. AuNP-based colorimetric aptasensors

It is well known that AuNPs exhibit strong distance-dependent optical properties. Once aggregated, the color of AuNPs changes from red to blue, accompanying with a bathochromic shift of the absorption spectrum. This phenomenon is attributed to the alterations in the localized surface plasmon resonance (LSPR) absorption generated by AuNPs. As AuNPs are generally thermodynamically unstable and tend to flocculate at high ionic strength conditions, surface adsorption of single-stranded DNA (ssDNA) can protect them from salt-mediated aggregation [5]. It has become one appealing strategy extensively adopted for a label-free colorimetric aptasensing. For example, Yang and coworkers have reported a rapid and sensitive aptasensor for OTA detection (Fig. 1A) [6]. When OTA is introduced, the color of the OTA’s aptamer switched from a random coil to a rigid G-quadruplex conformation required for target binding. AuNPs are no longer protected and aggregation occurs accompanied by a distinct color change from red to blue. This strategy has further been extended to the analysis of a variety of toxins, such as Staphylococcal enterotoxin B (SEB) [7], aflatoxin-B1 (AFB1) [8], ZEN [9], and microcystin-LR (MC-LR) [10]. To achieve an effective detection, it is noteworthy that the ratio
Table 1. Noble metal nanomaterial-based aptasensors for microbial toxin detection.

| Nanomaterials | Method                        | Target     | Strategy                                        | LOD            | Ref. |
|---------------|-------------------------------|------------|------------------------------------------------|----------------|------|
| AuNPs         | Colorimetric                  | OTA        | Target-induced aggregation of AuNPs            | 20 nM          | [6]  |
|                | Colorimetric                  | SEB        | Target-induced aggregation of AuNPs            | 10 ng/mL       | [7]  |
|                | Colorimetric                  | AFB1       | Target-induced aggregation of AuNPs            | 0.025 ng/mL    | [8]  |
|                | Colorimetric                  | ZEN        | Target-induced aggregation of AuNPs            | 12.5 nM        | [9]  |
|                | Colorimetric                  | MC-LR      | Target-induced aggregation of AuNPs            | 0.37 nM        | [10] |
|                | Colorimetric                  | MC-LR      | Target-induced disassembly of AuNPs            | 0.05 nM        | [51] |
|                | Colorimetric                  | OTA        | Enzyme-induced aggregation of AuNPs            | 5.0 nM         | [11] |
|                | Colorimetric                  | AFB1       | Catalytic DNA circuit amplification strategy   | 10 pM          | [12] |
|                | Colorimetric                  | SEB        | Target-regulated growth of AuNPs               | 1 nM           | [13] |
|                | Colorimetric                  | OTA        | Target-regulated growth of AuNPs               | 1 nM           | [14] |
|                | Lateral Flow Test Strip       | OTA        | Competition between OTA and cDNA with aptamer  | 1 ng/mL        | [15] |
|                | Lateral Flow Test Strip       | ZEN        | Competition between ZEN and cDNA with aptamer  | 20 ng/mL       | [16] |
|                | Fluorescent method            | OTA        | FRET between FAM-aptamer and AuNP              | 5 nM           | [17] |
|                | Electrochemical               | CDTA       | Electrode was modified AuNPs-capture DNA; aptamer-HRP was used to amplify the detection signal |                |      |
| AuNRs          | LSPR-based portable method    | OTA        | Aptamer-labeled AuNRs were coated on the optical fiber; aptamer-OTA binding events caused the LSPR peak of AuNR red shift. | 12 pM          | [25] |
|                | Multicolor colorimetric method| OTA        | G-quadruplex-hemin DNAzyme-mediated etching of AuNRs | 30 nM          | [26] |
|                | Multicolor colorimetric method| OTA        | ALP-mediated metalization of AuNRs             | 9 nM           | [27] |
|                | SERS                          | SEB        | Aptamer modified on the MNPs to capture SEB from the food matrix; Raman reporter and aptamer immobilized AuNRs form a sandwich-like detection model | 2.2 × 10⁻⁶ M   | [28] |
|                | Electrochemical               | OTA        | AuNRs modified-Au electrode; DNA tetrahedral nanostructures | 0.26 pg/mL     | [29] |
| AuNCs          | Fluorometric                  | AFB1 and ZEN| FRET between dual-color AuNCs and WS₂ nanosheets | AFB1: 0.34 pg/mL; ZEN: 0.53 pg/mL | [32] |
| AgNPs          | Fluorometric                  | AFB1       | FRET between fluorescent polymer dots and AgNPs | 0.3 pg/mL      | [33] |
|                | SERS                          | OTA        | Target-mediated Raman reporter Cy5 labelled aptamer desorption from the surface of AgNPs | 0.1 nM         | [34] |
|                | Electrochemical               | OTA        | Gold electrode covered with neutral red and aptamer-AgNPs | 0.05 nM        | [35] |
|                | Electrochemical               | OTA        | Aptamer was immobilized on the gold electrode; AgNPs/PDANSs were used as electrochemical tracer | 0.57 pm        | [36] |
| AgNCs          | Fluorometric                  | SEA        | FRET between AgNCs and PPNPs                  | 0.3393 ng/mL   | [38] |
|                | Fluorometric                  | T-2 toxin  | FRET between AgNCs and MoS₂                   | 0.93 pg/mL     | [39] |
|                | Fluorometric                  | OTA and AFB1| Magnetic separation and DNA-template AgNCs as the fluorophore | OTA: 0.2 pg/mL; AFB1: 0.3 pg/mL | [41] |
|                | Electrochemical               | OTA        | Aptamer was modified on the surface of SPGE; AgNCs metallization on OTA/aptamer complex | 0.7 pg/mL      | [42] |
between aptamers and AuNPs requires careful adjustments. Typically, a least excess of free aptamers is needed to ensure an adequate protection of AuNPs against non-specific aggregation, but without jeopardizing the detection sensitivity. Recently, a sensitive colorimetric aptasensor was developed for the detection of OTA by He and coworkers based on enzyme-induced AuNP aggregation [11]. When OTA was absent, the target-binding aptamer pre-immobilized on the magnetic nanoparticles (MNPs) was hybridized to its complementary sequence possessing an enzyme at one end (cDNA-ALP). However, as the OTA was present, the duplex structure would switch to an OTA-aptamer complex together with the release of cDNA-ALP. Followed by magnetic separation to remove immobilized enzymes, the liberated enzyme (ALP) can turn over the substrate, initiating the subsequent reactions, leading to salt-mediated AuNP aggregation. The proposed method was successfully applied to determine OTA in red wine and grape juice samples, offering high sensitivity with a detection limit as low as 5.0 nM. In addition, with efficient cascade amplification strategies, enzyme-free DNA circuits could also facilitate the development of ultrasensitive toxin detection. For example, a catalytic DNA circuit was devised by Chen et al. to amplify the colorimetric signal of AuNPs for the determination of AFB1 [12]. The constructed assay is ultrasensitive, allowing the visual detection of AFB1 as low as 10 pM without instrumentation.

Recently, a new generation of colorimetric aptasensors has been developed based on the target-regulated growth of AuNPs. According to this principle, an aptasensor was constructed by Zhou and coworkers for the detection of SEB [13]. In the absence of SEB, the guanine (G)-rich DNA oligonucleotides, which were sealed by the SEB aptamer, lost their ability to consume H₂O₂. Hence, AuNPs were synthesized in situ through the reduction of gold ions by H₂O₂, leading to a sensing solution with a red color. However, in the presence of SEB, the structure switching of the aptamer triggered the liberation of G-rich DNA oligonucleotides. The formation of G-quadruplex-based DNAzymes exhibiting a peroxidase-mimicking activity is capable of consuming H₂O₂, which in turn, producing blue-colored solutions. The linear range of this method extended from 0.1 to 500 ppt, allowing for clinical diagnosis. Similarly, another target-regulated-AuNP-growth strategy was developed by Soh et al. for OTA detection [14]. The interaction between aptamer and its target may affect the amount of aptamer adsorbed on the Au NP surface, which consequently generate AuNPs of variable
morphologies. As the OTA concentration increased, the color of the solution changed from blue to red. A visible colorimetric response was archived successfully in red wine samples for the detection of OTA at concentrations as low as 1 nM.

Due to the advantages of simplicity, portability, convenience, and low cost, AuNP-based lateral-flow test strips hold great potential for on-site food-contamination detection. Zhou et al. developed, for the first time, an AuNP-based lateral-flow test strip for the colorimetric detection of OTA using an aptamer as the recognition element [15]. With its high sensitivity and selectivity, a visual limit of detection of 1 ng mL$^{-1}$ of OTA was achieved with in 15 min. Later, the designed aptamer-based strip was also successfully applied to the determination of ZEN in corn samples [16]. The above results indicate that the current technology is able to provide rapid, sensitive and reliable results for numerous food contaminants and multitudinous sample inspection.

2.1.2. AuNP-based fluorescent aptasensors

Due to the strong LSPR and high coefficient extinction, AuNPs can also be used as energy acceptors for constructing aptasensors based on the mechanism of Förster resonance energy transfer (FRET). For example, an aptasensor was reported by Lv et al. for OTA detection with the use of AuNPs as efficient fluorescence quenchers [17]. As shown in Fig. 1B, in the absence of OTA, the carboxyfluorescein (FAM)-labeled aptamer was adsorbed on the surface of the AuNPs, leading to fluorescence quenching of FAM. Upon addition of OTA, the binding event between OTA and the aptamer caused the aptamer to desorb from the surface of the AuNPs, which led to a recovery of the FAM fluorescence. This turn-on aptasensor had a detection limit of 5 nM for OTA and was found to be to be free of interference by the sample matrix of ginger powders.

2.1.3. AuNP-based electrochemical aptasensors

Thanks to the high surface-to-volume ratio, high electron transfer efficiency and excellent biocompatibility, AuNPs has been extensively employed as a solid support for biomolecule immobilization to facilitate the electron transfer between the electrodes and biomolecules, giving the electrochemical devices with high sensitivity. For example, a sensitive AuNP-based electrochemical aptasensor has
been constructed by Luo and coworkers for the detection of type-A *C. difficile* toxins (CDTA) (Fig. 1C) [18]. In the developed device, the electrode was modified by layer-by-layer self-assembly of Nafion–Thionine, AuNPs, capture DNA, and horseradish peroxidase (HRP)-labeled aptamer, respectively. When CDTA was present, HRP-labeled aptamer would dissociate from the electrode surface through the target binding-induced conformational change. As a result, the catalytic activity of HRP was reduced, accompanying by a decrease in the amperometric response. The detection limit was shown to be 1 nM for CDTA with good selectivity, reproducibility, stability, and accuracy. Afterward, Deng et al. has suggested a similar strategy for the detection of SEB [19]. The electrochemical aptasensor, based on a AuNPs/ZrO2/chitosan-modified electrode coupled with a HPR-assisted signal amplification process, was capable of detecting SEB in a wider linear range with a much lower detection limit, providing a versatile tool in clinical diagnosis. To achieve a superior electrochemical response, an ultrasensitive electrochemical aptasensor based on the two-level cascaded signal amplification strategy was described by Yang et al. for OTA detection [20]. In the design, capture DNA, aptamers, and reporter DNA-modified AuNPs were immobilized on the electrode in a sandwiched manner. While AuNPs has been employed as the first-level signal enhancer, a great number of G-rich DNA hybridized on the AuNP surface promoted abundant methylene blue (redox indicator) anchoring, allowing a second-level signal enhancement to further amplify the initial electrochemical response. In the presence of OTA, the binding-induced conformation change enabled the release of G-rich DNA/AuNPs from the electrode surface, thereby reducing the electrochemical response. The proposed method showed promise in the sensitive detection of OTA with a low detection limit of 0.75 pM. Another signal enhancement approach based on two-round enzymatic amplification reactions was adopted by Zheng et al. for trace detection of AFB1 [21]. The aptamer probes were immobilized on the electrode surface and hybridized with the cDNA-modified AuNPs co-tethered with multiple copies of telomerase primers. Telomerase was then added to elongate the primers for abundant methylene blue anchoring, thereby enlarging the initial current signal to achieve the first round of amplification. As the target was present, the probe aptamers would undergo conformation change to release the primer-AuNP-cDNA conjugates from the sensing interface. Exonuclease III was further used to hydrolyze the resulting dsDNA probes to assist AFB1 release and recycling for the second round of amplification. Taking
advantage of the dual-amplification effect, this signal-off aptasensor enabled three orders of magnitude improvement in detection sensitivity, achieving an ultralow detection limit of \(0.6 \times 10^{-14}\) pg mL\(^{-1}\) of OTA.

Moreover, a disposable aptasensing device for label-free detection of fumonisin B1 (FB1) has been reported by Ren et al. In the design, the screen-printed carbon electrode (SPCE) was equipped with a self-designed microcell. A dense decoration of AuNPs on SPCE was accomplished by an in situ electrodeposition, providing abundant binding sites for aptamer anchoring, finally enhancing the impedance signal significantly. The specific binding between FB1 and aptamer inhibited the redox probe \([\text{Fe(CN)}_6]^{3/-4}\) from reaching the electrode surface. Therefore, FB1 can be easily detected by monitoring the electrochemical impedance spectroscopy of the system. The developed aptasensor presents a promising strategy to achieve facile, cost-effective, and highly sensitive detection of FB1, with a limit of detection of 3.4 pg mL\(^{-1}\) [22]. He et al. also reported an aptasensor utilizing ZnO nanorods for higher AuNPs and probe loading to effectively improve the electrode performance toward PAT detection [23].

Upon target binding, the formation of PAT-aptamer complex on the electrode surface would impede the electron transfer between the electrode and a redox indicator, hexacyanoferrate, causing the current to decrease. The prepared aptasensor showed satisfactory results when applied to the determination of PAT in apple juice samples. A signal-on electrochemical aptasensor was developed by Jalaliana et al. for the detection of aflatoxin M1 (AFM1) [24]. In the presence of the AFM1, the hairpin structure of the probe aptamers would undergo conformation change; thereby allowing a target-induced immobilization of AuNPs in close contact to the electrode surface. Upon the addition of redox indicators, the current signal would be increased, enabling a sensitive detection of AFM1 with a detection limit of 0.9 ng L\(^{-1}\). The developed aptasensor has been successfully applied to AFM1 analysis in real samples, including milk and serum samples.

### 2.2. AuNR-based aptasensors

AuNRs, a particular type of anisotropic metal NPs, exhibit two LSPR peaks, namely, a longitudinal and a transverse LSPR peak. The LSPR wavelength, especially for the longitudinal mode can be tuned over hundreds of nanometers in the visible and near-infrared regions by alterations of the aspect ratio (length/width). For this reason, AuNRs are considered as an attractive optical transducer for biosensing, in particular when a specific wavelength is desired.

#### 2.2.1. AuNR-based LSPR aptasensors

Lee et al. constructed a portable LSPR aptasensor based on optical fibers for facile and rapid OTA detection [25]. In the design, the aptamer-labeled AuNRs were coated on the optical fibers and subjected to OTA analysis. As illustrated in Fig. 2A, upon target binding, the aptamer is transformed into a G-quadruplex structure, and thus, leading to a red shift of the LSPR peak. Through the sequential dipping and measurement process, OTA in grape juice can be accurately detected with a limit of detection of 12.0 pM. The developed device is simple to operate and can be applied to in-situ analysis of OTA.

#### 2.2.2. AuNR-based multicolor colorimetric aptasensors

In addition to the alternation in local refractive index induced with an aptamer-target binding event, etching or metallization of AuNRs also lead to a significant modulation of the LSPR peak, turning solutions of rainbow-like colors. Based on this principle, several multicolor colorimetric methods have been developed for the visual semiquantitative determination of a variety of targets. Yu et al. established a multicolor colorimetric aptasensor for the visual detection of OTA based on the G-quadruplex–hemin DNAzyme-mediated etching of AuNRs [26]. A DNA sequence that was composed of a hemin aptamer and an OTA aptamer was designed as a recognition probe. In the absence of OTA, this ssDNA probe was cut into fragments by Exo I from the 3’ end. However, in the presence of OTA, the target binding-induced structure switching can halt the Exo I-mediated digestion due to the steric hindrance. At the same time, the retained hemin aptamer can convert to its active form and exerts a peroxidase-like activity to catalyze the oxidation of 3,3’,5,5’-tetramethylbenzidine (TMB) to TMB\(^{3+}\). TMB\(^{3+}\) was capable to etch AuNR by oxidizing Au (0) to Au(I), causing a vivid color change for visual inspection of OTA with the detection limit of 30 nM. Tian et al. reported another multicolor colorimetric approach for OTA detection on the basis of enzyme-mediated metallization of AuNRs (Fig. 2B) [27]. With OTA binding to its aptamer, the structural switching of the aptamer enabled the release of cDNA-ALP. Following the cDNA-ALP-catalyzing reactions, the obtained products, ascorbic acids could reduce Ag\(^{+}\) to Ag on the surface of AuNRs, resulting in a visible multicolor change. The assay was quite sensitive, since approximately 9 nM OTA could be readily detected.
by the naked eye. Collectively, AuNR-based multi-color colorimetric method hold great potential for the on-site visual detection of toxins in resource-limited regions.

2.2.3. AuNR-based SERS aptasensors

Plasmonic nanostructures, particularly AuNPs and AgNPs have been explored extensively as SERS substrates for decades. In general, anisotropic NPs possessing sharp tips, corners and edges give rise to intense local electromagnetic fields, providing a strong Raman enhancement for SERS. SERS with state-of-the-art performance is capable of offering extremely high sensitivity, down to the single-molecule detection level. Temur et al. established an ultrasensitive aptasensor using AuNRs as SERS probes for SEB detection [28]. As shown in Fig. 2C, Au-coated MNPs were functionalized with the capture aptamers to separate SEB from the food matrix. Then, the aptamer-immobilized and Raman reporter-tagged AuNRs can conjunct with the captured targets via a sandwich-like manner, giving rise to a strong SERS signal. The proposed method achieved a detection limit of $2.2 \times 10^{-16} \text{M}$ of SEB and is applicable to complex matrices, such as milk, urine, and serum.

2.2.4. AuNR-based electrochemical aptasensors

Similar to AuNPs, AuNRs have also been widely used in electrochemical biosensors for signal amplification. An ultrasensitive aptasensor has been developed by Wei et al. for OTA detection by utilizing an AuNR-modified electrode to enhance the loading of DNA tetrahedral probe (DTP) and to promote the electron-transfer rate (Fig. 2D) [29]. DTP, as a novel type of probe, was modified with an OTA aptamer on the top vertex and three thiol groups at the other three vertices. Once assembled onto the electrode, DTP exhibited a stronger bonding with gold surfaces than the conventional ssDNA probe with just one anchor point. Moreover, the mechanical rigidity of DTP can force the aptamer to adopt an upwards orientation, thereby providing efficient molecular recognition at the sensing interface with encouraging outcomes [30]. The OTA/aptamer binding event causes a transformation of the aptamer structure into a G-quadruplex. The G-quadruplex binds to hemin to form a DNAzyme with peroxidase-mimicking activity that can catalyze the oxidation of aniline to polyaniline in the presence of $\text{H}_2\text{O}_2$, thereby generating an amplified current signal. The obtained aptasensor presented acceptable reproducibility and repeatability, and had high selectivity toward OTA against other mycotoxin, with the detection limit of 0.26 pg mL$^{-1}$.

2.3. AuNC-based aptasensors

When the particle size of noble metals becomes comparable to the Fermi wavelength for the electron, LSPR peaks disappeared. In this ultrasmall size regime, metal nanoclusters (NCs), consisting of several to hundreds of atoms, exhibit unique molecule-like properties including discrete electronic states and size-dependent photoluminescence. So far, fluorescent metal NCs have been extensively studied for biosensing and bioimaging due to their attractive set of features, such as ultrasmall size, good biocompatibility, variable emission wavelength and excellent photostability. As an important class of noble metal NCs, AuNCs are more attractive because of their ease of synthesis and modification [31]. For example, biomolecule-protected AuNCs with tunable fluorescent colors have been prepared based on one-pot strategy via a facile biomineralization-mimicking process [32]. After the covalent modification of AFB1 and ZEN aptamers, respectively, on the surfaces of the blue-(442 nm) and red-emitting (650 nm) AuNCs, these two donor probes were attached to the WS$_2$ nanosheet, as an efficient fluorescence quencher. Binding of the specific targets induced a concentration-dependent fluorescence recovery, allowing this dual-color turn-on nanoplatfor platform for simultaneous detection of AFB1 and ZEN. Under optimal conditions, AFB1 and ZEN can be sensitively detected with detection limits of 0.34 and 0.53 pg mL$^{-1}$, respectively.

The developed aptasensor presenting an accurate and reliable strategy for simultaneous recognition, holds a promising potential in multipurpose biosensing.

2.4. AgNP-based aptasensors

2.4.1. AgNP-based fluorescent aptasensors

Apart from gold, AgNPs have also attracted considerable attention for the development of biosensors. Due to their broad LSPR absorption and high extinction coefficients, particularly in the visible range, it makes them to be good candidates as efficient energy acceptors. For example, Nasirian et al. developed an aptasensor for AFB1 detection based on the FRET process between aptamer-labelled fluorescent polymer dots and AgNPs conjugated with cDNA [33]. In the absence of AFB1, DNA hybridization would bring the FRET pair in
close proximity,witching off the fluorescence. As the aptamer bound to its target, the FRET pair was separated, resulting in recovery of fluorescence. The assay was successfully applied to the detection of AFB1 in wheat flour, with an ultralow detection limit of 0.3 pg mL\(^{-1}\).

2.4.2. AgNP-based SERS aptasensors

In addition, AgNPs have been widely employed as SERS substrates due to the strong electromagnetic field enhancement on and near the surface of AgNPs. Ganbold et al. developed a SERS aptasensor based on AgNPs for OTA detection [34]. The Raman reporter Cy5-labeled aptamer was adsorbed on the surface of the AgNPs, generating high SERS signals. The specific binding between the aptamer and OTA induced the departure of the SERS-tagged aptamer from the substrates, resulting in a decrease in the intensity of the SERS signals. This SERS aptasensor exhibited a low detection limit of 0.1 nM of OTA.

2.4.3. AgNP-based electrochemical aptasensors

Recently, the use of AgNPs on the electrode surface led to an enhanced analytical performance due to an improvement in the regularity of the composition of the electrode surface and an enhancement of the electron transfer between the electrode and the modified biomolecules. Evtugyn et al. developed an electrochemical aptasensor based on AgNPs for OTA detection [35]. In their design, a gold electrode covered with electropolymerized neutral red and AgNPs was covalently attached with multiple copies of thiolated OTA aptamers through Ag–S bonds. The formation of the aptamer-OTA complex resulted in switching the conformation of the aptamer, which caused an increase of the charge transfer resistance and impedimetric signal changes. The applicability of the developed aptasensor was validated in beer samples, with a detection limit of 0.05 nM of OTA. To further improve the detection sensitivity, Zhang et al. developed an ultrasensitive electrochemical aptasensor utilizing polydopamine nanospheres (PDANSs) for signal amplification [36]. Due to the large surface area of the PDANSs, a greater amount of AgNPs was immobilized on their surface. The detection was realized by the electrochemical stripping signal of AgNPs on PDANS. After OTA binding, a target-induced combination change enabled the release of signal tags on the aptasensor surface, thereby reducing the electrochemical response signal. The proposed strategy offered a wide linearity ranges of 4 orders of magnitudes and a low detection limit of 0.57 pM of OTA.

2.5. AgNC-based aptasensors

2.5.1. AgNC-based fluorescent aptasensors

As inspired by the nature process of biomineralization, the cytosine-rich oligonucleotide has been successfully utilized as a scaffold for the synthesis of

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Fig. 3. Schematic representation of an aptasensor based on AgNCs for the simultaneous detection of OTA and AFB1. Adapted with permission from Ref. [41]. Copyright Elsevier, 2016.
fluorescent AgNCs [37]. These DNA-templated AgNCs have received great attention as novel nanotools in bioapplications, due to their strong fluorescence, excellent light stability, and good biocompatibility. It is therefore, highly promising for the construction of label-free fluorescent aptasensors based on AgNCs. By designing a chimeric oligonucleotide, which contains a domain with a 12-mer cytosine for AgNC formation and a domain with an aptamer for target recognition, Zhang et al. developed a label-free aptasensor for Staphylococcal enterotoxin A (SEA) detection [38]. In their design, polypyrrole nanoparticles (PPNPs) were used as the quencher. The specific binding between SEA and the aptamer caused the aptamer–DNA–AgNCs complex to move away from the PPNPs, leading to an effective fluorescence recovery. This label-free aptasensor is very simple and cost effective, with a low detection limit of 0.3393 ng mL$^{-1}$. A similar strategy has also been reported by Khan et al. for the detection of T-2 toxin [39]. Based on the use of AgNC and MoS$_2$ as a FRET pair, a sensitive detection of T-2 toxin in maize and wheat samples was achieved with a low detection limit of 0.93 pg mL$^{-1}$.

Simultaneous monitoring of various mycotoxins is essential for food safety because agricultural products and foodstuff are often contaminated with multiple mycotoxins. Delightfully, the fluorescent emission of AgNCs are finely tunable by adjusting the sequence and length of the DNA template [40]. Therefore, AgNCs hold great potential for the simultaneous detection of different mycotoxins.

Based on this principle, Zhang et al. developed a fluorescent aptasensor for the simultaneous detection of OTA and AFB$_1$ [41]. As shown in Fig. 3, an OTA aptamer and an AFB$_1$ aptamer were immobilized on the surface of MNPs. Signal probes 1 and 2 were designed not only to be complementary to the OTA and AFB$_1$ aptamers, respectively, but also to be used as templates to form AgNCs with different maximum emission wavelengths. Moreover, the addition of Zn$^{2+}$ was found to increase the fluorescence intensity of the DNA-scaffolded AgNCs, and thereby improving the detection sensitivity. The detection limits of OTA and AFB$_1$ was approximately 0.2 and 0.3 pg mL$^{-1}$, respectively.

2.5.2. AgNC-based electrochemical aptasensors

Based on the fact that catalytic silver deposition (termed as the silver enhancement) can provide further signal enhance, silver-metallized aptamers have also been devoted to the design of electrochemical aptasensor for the ultrasensitive detection of OTA [42]. As a result of target binding, aptamers on the electrode surface would self-assembled into a G-quadruplex structures. After the introduction of Exo I to digest the randomly coiled unbound aptamers, chemical and electrochemical reduction processes were initiated to generate silver metallized aptamers for voltammetric detection. The combination of Exo-I-mediated background-signal suppression and AgNC-mediated signal amplification leads to a label-free, sensitive, specific, and “turn-on” electrochemical aptasensor for the precise determination of OTA at trace levels.
2.6. Bimetallic nanomaterial-based aptasensors

2.6.1. Bimetallic nanomaterial-based SERS aptasensors

As an ultrasensitive analytical technique, SERS has been extensively applied for trace microbial toxin detection to ensure food safety. In general, an average enhancement factor ranges between $10^4$ and $10^6$ was reported for metallic NPs made of gold, silver, and copper [43]. More recently, it has been found that gold–silver bimetallic nanomaterials have the ability to further improve the SERS signals compared to their mono-metallic gold or silver counterparts. This phenomenon is attributed to the tunability of the SPR band and the synergistic effect of these metals. For instance, Li et al. reported the use of heterogeneous gold nanostar (AuNS) core–AgNP satellite assemblies as SERS-active sensors for ultrasensitive detection of AFB1 [44]. The Raman reporter was modified on the surface of AgNP satellites. As shown in Fig. 4, in the absence of AFB1, hybridization occurred between the aptamer-modified AgNP satellites and cDNA-modified AuNS. Formation of the core–satellite assemblies give rise to an enhanced SERS signal. However, the specific binding of AFB1 to the aptamer caused the release of Ag satellites, resulting in a decreased SERS signal. The developed SERS aptasensor showed a high sensitivity for AFB1 with a detection limit of 0.48 pg mL$^{-1}$. A variety of core–satellite assemblies have also been fabricated for ultrasensitive detection; for example, e.g., an aptasensor based on Au nanoflower core–AgNP satellites was developed for the detection of MC-LR [45]. Similarly, the SERS-active assemblies exhibited amplified SERS signals, enabling a sensitive detection of trace level of MC-LR based on the target-induced disassembly process.

Not only the heterogeneous core–satellite assemblies can produce sensitive SERS signals, gold–silver core–shell structures could also serve as excellent SERS substrates for target detection with high sensitivity. Based on this principle, various aptasensors have been developed. Yang et al. designed gold nanotriangle (AuNT)–silver core–shell nanomaterials (AuNT@Ag) for the sensitive and selective SERS detection of AFB1 [46]. To achieve AFB1 detection, aptamer-MNP and aptamer-AuNT@Ag were used as capture probe and reporter probe, respectively. Under optimized conditions, a detection limit of 0.54 pg mL$^{-1}$ was obtained. Another aptasensor based on core–shell nanomaterials composed of Au(core)@Au–Ag(shell) was also constructed by Shao et al. for the detection of OTA [47]. In this strategy, cDNA-MNP and aptamer-Au(core)@Au–Ag(shell) were prepared. Due to the high binding affinity between OTA and its aptamer, the presence of OTA induced cDNA-MNP to break away from aptamer-Au(core)@Au–Ag(shell). And a significant change in Raman intensity was observed. An ultralow detection limit of 0.004 ng mL$^{-1}$ was achieved successfully in red wine samples. Collectively, in comparison to the mono-metallic counterparts, these heterogeneous core–shell nanostructures possessing built-in nanogap hotspots have shown superior SERS signal amplification, holding great potential for ultrasensitive detection.

2.6.2. Bimetallic nanomaterial-based electrochemical aptasensors

In addition to the LSPR, other physicochemical characteristics such as catalytic and electrocatalytic properties of bimetallic nanomaterials may also be tremendously different from those of the parent metals. Alloyed structures, in particular, often result in a tunable electrochemical activity, attributing to the synergy effect of different electronic structure, achieved by gradually varying the composition [48]. Ji et al. developed a palladium–platinum nanoparticle (Pd–PtNP)-based aptasensor for ZEN detection [49]. In their design, a glassy carbon electrode (GCE) was modified with spherical Au–PANI–Au (PANI: polyaniline) nanocomposites to increase the conductivity of the electrode and to provide more aptamer-binding sites. The cDNA-modified sakura-shaped Cu@L-Glu/Pd-PtNPs was employed as a signal tag. The hybridization reaction between the aptamer and cDNA brought the Cu@L-Glu/Pd-PtNPs close to the GCE surface, which produced obvious electrochemical signals. After the specific ZEN-aptamer binding, cDNA-Cu@L-Glu/Pd-PtNPs was released from the GCE, resulting in a significant signal decrease. Through the effective signal amplification mediated by Cu@L–Glu/Pd–PtNPs, an ultralow detection limit of 0.45 fg mL$^{-1}$ was achieved. By utilizing rGO–TEPA–Au@PtNRs as the signal tag, Zhong et al. developed another novel aptasensor for T-2 toxin detection [50]. Similarly, the high catalytic ability of Au@PtNRs enabled an efficient signal amplification, leading to a sensitive detection of T-2 toxin as low as 1.79 fg mL$^{-1}$. Collectively, bimetallic nanomaterials which demonstrated superior catalytic and electrocatalytic activities are promising signal enhancers in electrochemical aptasensors. Such devices provide high sensitivity, representing a promising strategy for the detection of small trace toxins in food samples.
3. Conclusion and outlook

Aptamers, also known as chemical antibodies, play an increasingly important role in the molecular recognition of microbial toxins because of their multiple advantages including high binding affinity, high specificity, high stability, repeatable synthesis and modification, predictable structure, and flexible amplification strategies that are easy to develop. At the same time, the outstanding physical and chemical properties of metallic nanomaterials make them excellent candidates for signal transduction. Therefore, the combination of aptamers and nanomaterials holds great potential for microbial toxin detection.

Despite a numerous aptasensors have been developed by incorporating noble metal nanomaterials for selective and sensitive detection of food toxin, most of the noble metal nanomaterial-based aptasensors developed up to date are still at the experimental stage. There are abundant challenges that remain to be overcome before it can be employed for practical detection applications. For instance, due to the complex food matrices, appropriate attention needs to be paid to the non-specific adsorption of non-target molecules on nanomaterials, which might lead to false positive signals. Moreover, some noble metal nanomaterial-based aptasensors still rely on advanced precision instruments and professional operators, which limit their widespread use in resource-limited settings.

Nevertheless, noble metal nanomaterial-based aptasensors are a promising tool for microbial toxin monitoring. As further research progresses, we will witness a thriving and prosperous development of noble metal nanomaterial-based aptasensors for portable and on-site detection of food microbial toxin in the future.

Declaration of competing interest

None.

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