Selection, Identification, and Application of Aptamers against Agaricus bisporus Lectin to Establish an Aptamer-AuNPs Colorimetric Method for Detection of ABL

Hao Gao, Yiling Tian, Min Zhang, Jianhui Liu, Yaowu Yuan, Jianxin Tan, and Aijin Ma

1 College of Food Science and Technology, Hebei Agricultural University, No. 289 Lingyusi Street, Baoding 071001, Hebei, China
2 Shijiazhuang Junlebao Dairy Co., Ltd., No. 36 Shitong Road, Shijiazhuang 050221, Hebei, China
3 Beijing Technology and Business University, Beijing, No. 11 Fucheng Road, Beijing 100048, China

Correspondence should be addressed to Jianxin Tan; jianxintan@sina.com and Aijin Ma; maaj@btbu.edu.cn

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Agaricus bisporus lectin (ABL), which is one of the antinutritional factors in A. bisporus, is an important allergen and harmful to human health. Due to the shortcomings of the current detection methods, it is extremely urgent to establish a rapid and sensitive detection method for ABL in foods. To isolate the ssDNA aptamer of ABL, 13 rounds of subtractive systematic evolution of ligands by exponential enrichment (SELEX) selection were carried out. As a result, six candidate aptamers were selected and further examined for their binding affinity and specificity by enzyme-linked aptamer method. One aptamer (seq-41) against ABL with a high affinity and specificity was isolated and demonstrated to be the optimal aptamer whose dissociation constant reaches the nanomolar level, $K_d = 31.17 \pm 0.1070 \text{ nM}$. Based on seq-41, an aptamer-AuNPs colorimetric method was established to detect ABL with a linear range of $0.08 \sim 1.70 \mu g/mL$ and the detection limit is $0.062 \mu g/mL$. This study provides a novel aptamer-AuNPs colorimetric method with high sensitivity and specificity for detection of ABL and a novel strategy for development of detection method of fungal or plant allergens.

1. Introduction

Agaricus bisporus, also known as button mushroom, white mushroom, or champignon, is one of the most widely cultivated and consumed edible mushrooms in the world [1]. As one of the most studied bioactive proteins [2–4], A. bisporus lectin (abbreviated ABL) is a 64 KDa glycoprotein composed of four subunits [5] and has binding specificity to Gal ($\beta$-1, 3) GalNAc [6]. On the one hand, lectins display antifungal and immunomodulatory activities [7]. On the other hand, lectins have blood coagulation activity and the antinutritional effects on animals [8] because they are one of the components formed in the long-term evolution of plants to resist diseases, insect pests and animal digestion. These lectins may contribute to the development of multiple food sensitivities, food allergies, and autoimmune conditions [9]. Therefore, the establishment and development of methods for rapid detection and identification of allergens in food (e.g., lectins) can effectively prevent people from allergies, which is very important for health.

At present, the common qualitative and quantitative assays of the lectin from plant and fungi are generally based on the agglutination assay and immunosassay. For example, the hemagglutination activity test can be served to monitor and quantify lectin activity at each step during purification [10]. In addition, based on the protein properties of lectins and immunological techniques, a variety of assay methods for quantifying lectin activity have been developed, such as electrochemical immunoassay (ECIA) [11] and enzyme-linked immunosorbent assay (ELISA) [12, 13]. Comparatively, the procedure of hemagglutination test is relatively simple and easy to be operated. However, the same lectin
may show different agglutination activity to red blood cells from the different kinds of animals, and its accuracy and sensitivity are relatively low. Accordingly, this method may be more suitable for semiquantitative or qualitative analysis of lectin content. Except for the hemagglutination test, other detection methods have the disadvantages of cumbersome steps and complex operation. Hence, it is particularly important to develop a new convenient and sensitive method for the detection of lectin in order to provide technical support for ensuring food safety and human health.

Aptamers are short single-stranded DNA or RNA with high affinity and specificity to the target molecule [14] and are screened by systematic evolution of ligands by exponential enrichment (SELEX) technology [14, 15]. The concept of aptamer was first proposed by Ellington [16] and Tuerk [17]. The SELEX technology has been widely explored and used in many fields during the past 30 years. Aptamers are also called “chemical antibodies” because their characters and applications are similar to antibody [18]. However, compared with antibodies, aptamers have many advantages, such as a wide range of target molecules, short preparation cycle, strong specificity, high affinity, lower immunogenicity, and lower cost of synthesis [19]. As a new type of specific recognition probe, aptamer provides a sensitive and specific optional mode for the development of the new detection methods in the field of food safety, for example, detection of biological toxins [20, 21], pathogenic bacteria [20], antibiotics [22], and heavy metals [23]. Colloidal gold, as a new type of nanomaterials, is popularly used as biological and chemical sensors owing to its merits of easy labeling, surface plasma resonance effect, and other chemical and optical properties [24]. Particularly, the gold nanoparticle-based colorimetric assay is widely used in environmental monitoring, public health, and disease diagnosis because of their simple, cost-effective fabrication and ease of use [24]. Combined usage of aptamer and gold nanoparticle-based colorimetric assay displays both advantages and provides a much more powerful and useful strategy for developing usable methods and their applications [25].

The study aims to explore a simple, easy, and sensitive method for the determination of *Agaricus bisporus* lectin. We use subtractive-SELEX technology to obtain high-affinity aptamers after 13 selection rounds. Using the selected aptamer as a specific recognition probe and AuNPs as an indicator, a visual detection method of ABL was established to effectively monitor the lectin content of *A. bisporus* products. The results of this study will provide a new reference for developing detection methods of other allergens from fungi or plants.

2. Materials and Methods

2.1. Materials. A single-strand DNA (ssDNA) library (5′-ATAGGAAGTCACGACAGCAGCAG-N40-TATGGTGCGTC-TACCTCTTGA-3′) and primers (forward primer1/FP1: 5′-ATAGGAAGTCACGACAGCAG-3′, reverse primer1/RP1: 5′-TCAAGAGGGTAGAGGACACA-3′, and reverse primer2/RP2: 5′-biotin-TCAAGAGGGTAGAGGACACATA-3′) for the amplifying and screening of ABL aptamer were designed and synthesized by Sangon Bioengineering (Shanghai) Co. Ltd. China. 2×Es Taq Master Mix (Dye) was purchased from Beijing ComWin Biotech Co., Ltd. China. *Agaricus bisporus* lectin FL-1420 was purchased from Vector Laboratories Inc. Bovine serum albumin, acrylamide, TEMED, streptavidin-horseradish peroxidase (HRP-SA), and 3, 3′, 5, 5′-tetramethylbenzidine (TMB) were purchased from Beijing Solarbio Science & Technology Co., Ltd. China. Streptavidin magnetic beads were ordered from Suzhou Beaver Biomedical Engineering Co., Ltd. China. Nucleic acid coprecipitation reagent kits were bought from Takara Bio Medical Technology Co., Ltd. All other reagents were from Sinopharm Chemical Reagent Company, China.

2.2. Aptamer Selection. The aptamer selection procedures followed the protocol of Du et al. [26] with some minor changes. First, ABL or BSA was adsorbed onto the wells of 96-well plates. In detail, at the first round, the plates were coated with 200 μL/well of 50 μg/mL of ABL or BSA dissolved in binding buffer (20 mM HEPES, 120 mM NaCl, 5 mM KCl, 1 mM CaCl₂, and 1 mM MgCl₂, pH 7.4) and incubated overnight at 4°C. After washing three times with the binding buffer, the plates were blocked with 3% (w/v) BSA solution for 2 h at 37°C and then the unbound BSA was removed. Next, 2 mM of the random ssDNA pool was prepared in 100 μL of binding buffer and denatured for 5 min at 95°C, cooled for 5 min at 4°C, and then incubated for 10 min at room temperature. The denatured ssDNA pool was added into the BSA coated wells and incubated for 40 min at 37°C to eliminate the ssDNAs that were able to bind BSA. The unbound ssDNAs were collected, added into the ABL-coated wells, and incubated for 60 min at 37°C. After incubation, the wells were washed three times with the binding buffer containing 0.05% Tween-20 to remove unbound ssDNAs. The remaining ssDNAs that bound to ABL were incubated in elution buffer (20 mM Tris-HCl, 4 mM guanidine isothiocyanate, and 1 mM DTT, pH 8.3) for 10 min at 90°C and then collected and recovered by alcohol precipitation. The enrichment of ABL binding ssDNAs was monitored by the recovery of ssDNA. The purified ssDNA were amplified by PCR (2×Es Taq Master Mix (Dye) 5 L, FP1 (20 μM) 0.5 L, RP2 (20 μM) 0.5 μL, ssDNA Template 2.0 μL, ddH₂O 9.5 μL in a total volume of 25 μL). PCR was performed following the program: pre-denaturation at 95°C for 5 min, followed by 20 cycles of 95°C for 30 s, 58°C for 30 s, and 72°C for 30 s with a final extension at 72°C for 5 min. After amplification, PCR products were denatured using 150 mM NaOH solution, and the unlabeled ssDNA strands were separated using streptavidin-coated magnetic beads and concentrated by ethanol precipitation. The isolated ssDNA strands were subjected to the next round of SELEX until the thirteenth round of selections were completed.

2.3. Aptamer Sequencing and Analysis. The entire enriched ssDNA pool of the last SELEX round was amplified by PCR and ligated into pMD18-T vector. The resultant ligation products were transformed into *E. coli* DH5α competent
cells. The plasmids containing the selected candidate aptamers harbored in the positive colonies were extracted and sequenced by Beijing Genomics Institute (BGI), China. Multisequence alignment and phylogenetic tree of the enriched sequences were performed with Clustal X 2.1 and MEGAX 10.0.5 software. Secondary structure and the free energy prediction of the enriched sequences were conducted by using RNA Structure Network Service (http://rna.urmc.rochester.edu/RNAstructureWeb/). Based on the primary sequences, secondary structure of the candidate aptamers, and the phylogenetic tree, six representative candidate aptamers were selected for subsequent affinity and specificity analysis.

2.4. Evaluation of the Aptamer Binding Affinity and Specificity. For determination of affinity, firstly, 100 μL of the biotin-labeled candidate aptamers in different concentrations (10, 50, 100, 200, 300, and 400 nM) were denatured at 95°C for 5 min and then immediately cooled at 4°C for 5 min, followed by incubation for 10 min at room temperature. Next, the solution was transferred into the ABL precoated wells and incubated at 37°C for 1 h. Each well was washed three times with 200 μL washing buffer (binging buffer plus 0.05% Tween-20) and then incubated with 100 μL streptavidin-horseradish peroxidase (dilution 1:5000) at 37°C for 30 min. After washing three times with washing buffer, 100 μL substrate (TMB) was added into each well and incubated for 10 min at room temperature, and then, 50 μL of 2M H₂SO₄ was added to stop the reaction. The optical density (OD) at 450 nm was measured using BSA precoated well as blank. The dissociation constant (Kₐ) value of aptamer, which reflects the aptamer affinity, was analyzed with GraphPad Prism software (version 6.01, Windows) and calculated according to the equation Y = B_max X / (Kₐ + X), where B_max is the number of maximum binding sites, X is the aptamer concentration, and Y is the absorbance value.

To evaluate the specificity of the selected aptamer candidates, 1 μg ABL lectin extract from fresh A. bisporus fruit, and BSA were precoated in a 96-well microplate and incubated with 200 nM biotinylated aptamer for binding with ABL. The following steps were the same as that of affinity test. Finally, the amount of the binding aptamers is determined by absorbance value at 450 nm.

2.5. Establishment of an Aptamer-AuNPs Colorimetric Method for Detection of ABL

2.5.1. Preparation of AuNPs. AuNPs were prepared by sodium citrate reduction method created by Frens [27]. 95.8 mL ddH₂O was added to the beaker, and then 4.2 mL 1% chloroauric acid solution was added and heated to boiling, 10 mL 1% sodium citrate solution was quickly added and heated to boiling for 20 min, and then the solution was continually stirred. The AuNPs solution was finally obtained after cooling to room temperature and the solution color turned to wine-red.

2.5.2. Optimization of NaCl Concentration. 20 μL of NaCl solution in different concentrations (0, 50, 100, 150, 200, 250, 300, 350, and 400 mM) was mixed with 100 μL of colloidal gold solution, respectively, and incubated at 37°C for 10 min. The color change was observed and measured over the wavelength ranging from 400 nm to 700 nm.

2.5.3. Optimization of Aptamer Concentration. 100 μL of different concentrations of the aptamer solution (0, 0.5, 1.0, 2.0, 4.0, 6.0, 8.0, and 10.0 μM) was added to 100 μL of colloidal gold solution, mixed, and incubated at 37°C for 3 h, and finally 20 μL 250 mM NaCl was added. The color change was observed and the absorbance value was measured at 520 nm.

2.5.4. Quantitative Detection of ABL and the Establishment of the Standard Curve. 100 μL of 2 μM aptamer solution was added to 100 μL colloidal gold solution and incubated at 37°C for 3 h in a shaker. Then, different concentrations of ABL standard samples (0, 0.043, 0.087, 0.17, 0.30, 0.35, 0.39, 0.87, 1.28, 1.7, and 2.0 μg/mL) were added and incubated at 37°C for 1 h. Finally, after adding 20 μL of 250 mM NaCl, the color change was observed and the ratio of the absorbance value at 620 nm to 520 nm was measured. A standard curve was generated using GraphPad Prism 6.01 software.

2.5.5. Specificity Tests. To evaluate the specificity of the aptamer, the same procedures were performed as mentioned in Section 2.5.4 with the samples of standard ABL, *Agaricus bisporus* extract, BSA, *Flammulina velutipes* extract, and *Lentinula edodes* extract in the reaction mix, and the specificity was determined by the ratio of the absorbance value at 620 nm to 520 nm. Fresh lectin extracts were prepared as follows: fresh fruiting bodies of *A. bisporus*, *F. velutipes*, and *L. edodes* were mixed with 0.05 M PBS (pH 7.2), ground in a blender, and soaked overnight at 4°C. The homogenate was filtered through cheesecloth, and the filtrate was centrifuged at 14000×g for 15 min at 4°C. The supernatants were collected as fresh *A. bisporus*, *F. velutipes*, and *L. edodes* extract samples.

3. Results and Discussion

3.1. Selection of the Specific Aptamers against ABL. Proteins can be physically adsorbed by the interaction between their hydrophobic groups and the hydrophobic groups on the surface of polystyrene microplate. As glycoprotein, ABL can fix itself on the microplate by hydrophobic interaction. Taking advantage of this feature, we can separate and isolate the ssDNAs which stably bind to the ABL from the ssDNAs that do not bind by repeatedly washing plates. The application of subtractive-SELEX allows removing nonspecific binding ssDNAs before each binding to the target [28]. This negative screening step greatly improves screening efficiency [29]. By using these two strategies, the selection of ABL aptamer was performed.
To select the specific aptamers against ABL, we first designed a ssDNA library containing a central randomized sequence of 40 nucleotides flanked by defined primer-binding sites of 20 nucleotides for PCR amplification. In the first round of SELEX selection, 2 nmol ssDNAs were fixed in the wells of the microplate to ensure that there was the sufficient diversity of oligonucleotide sequences in the initial selection system. In subsequent steps, gradually reducing the amount of ssDNA pool and ABL is conducive to improve the screening efficiency of ABL aptamers with high affinity. Table 1 listed the amount of ssDNA and ABL input during each round of aptamer-selection procedures.

As shown in Figure 1(a), at the first seven rounds of selection, the ssDNA recovery rate of each round obviously increased. After that, the recovery rates showed gently increase trend and reached up to the highest one at the end round. These results indicated that the specific ssDNAs binding to ABL were effectively enriched and the selection strategy was practicable. The bands around 80bp (Figure 1(b)), which were recovered from each selection round, indicated that the products of each selection round were correct and sufficient.

### 3.2. Sequence Analysis and Characterization of the Selected Aptamers against ABL

The products of the final SELEX round were cloned and sequenced. A total of 43 sequences were obtained. Among them, seq-1 appeared the highest frequency with nine repeats. For seq-2 to seq-6, each of them had two repeats. The rest only appeared once. The existence of high-frequency sequences provided evidence that the selection method served in this study was able to enrich aptamers of ABL with high efficiency. The phylogenetic tree was constructed according to the sequence homology comparison (Figure 2). As shown in Figure 2, the seq-2, seq-26, seq-32, seq-36, seq-38, and seq-39 were in a group and formed a large branch since they have relatively higher polymorphism. The rest 37 sequences clustered into another large group with much more homology and conserved sequences of the primary sequence motifs.

According to the comparison of the secondary structure and the minimum free energy, the isolated sequences were classified into six categories. A representative sequence from each group was picked out to generate diagram of their secondary structure (Figure 3). All the secondary structures contain closed or open loop, hairpin, and cruciform structure, which may form the complex three-dimensional structures and bind to the target via noncovalent bonds interaction with the target. The different secondary structure and the conformation changes affected by different environments may influence the affinity of aptamer to the target due to the variations of hydrophobic interactions, hydrogen bonding, and salt linkages formed between them [30]. The sequences of the representative candidate aptamers, their copy number out of the 43 cloned sequences, and the free energy were listed in Table 2.

### 3.3. Determination of the Affinity and Specificity of the ABL Candidate Aptamers against ABL

The enzyme-linked aptamer method was used to determine the affinity of the six candidate aptamers picked out from the 43 sequences. The dissociation constant (Kd) of the candidate aptamers was calculated by nonlinear regression analysis (Table 3). The binding affinity of aptamer has a high correlation with Kd value, which means Kd value reflects the aptamer affinity. The Kd values of the six aptamers were between 31 and 135 nM, indicating that all the selected candidate aptamers exhibit a high affinity towards ABL although their Kd values are more than four times different.

The nonlinear saturation curves of the six candidate aptamers were fit and generated by Graphpad Prism 6.01 software (Figure 4). As shown in Figure 4, when the aptamer concentration increased to 200 nM, the absorbance value tended to be stable, implying that the binding reaction reaches saturation. Among the six candidates, since the Kd values of seq-1 and seq-41 were relatively smaller, these two candidate aptamers were selected to further test the specificity of them.

Figure 5 showed the specificity of seq-1 and seq-41 to the standard ABL, the *Agaricus bisporus* extract, and BSA (negative control). The results revealed that the absorbance values of seq-1 and seq-41 were significantly higher than that of BSA, which demonstrated that both candidate aptamers had high specificity to both standard ABL and *Agaricus bisporus* extract. Comparatively, seq-41 was the best one of ABL aptamer due to the Ab450nm value of seq-41 was higher than that of seq-1. Thus, considering the affinity and specificity results, seq-41 was verified to be the best aptamer against ABL.

### 3.4. Establishment of an Aptamer-AuNPs Colorimetric Method for Detection of ABL

#### 3.4.1. Optimization of NaCl Concentration

Negatively charged AuNPs usually aggregates under the action of electrostatic force when the salt solution, for example, NaCl, is added [31]. The concentration of sodium chloride added directly affects the degree of AuNPs aggregation and will further interfere with the interaction between aptamer and colloidal gold. Therefore, in order to obtain the detection
results with high sensitivity and wide detection range, the optimization of NaCl concentration is particularly important. As shown in Figure 6, along with the salt concentration increasing in the range from 0 to 400 mM, the color of colloidal gold solution changed from red to purple and finally to gray. Meanwhile, the Ab520 value continued to decline and finally stabilized, indicating that the colloidal gold gradually accumulated under the action of salt. When the salt concentration is between 50 and 100 mM, the red color was gradually getting lighter and the precipitation was not observed. If the salt concentration increased from 150 mM to 200 mM, the colloidal gold solution turned purple and most of the AuNPs deposited. When the salt concentrations were in the range of 250–400 mM, the color of the colloidal gold solution became gray and the AuNPs were completely precipitated. Combined with the above observed results, the salt concentration at 250 mM was selected and served for subsequent test.

3.4.2. Optimization of Aptamer Concentration. Oligonucleotides, like aptamer, can bind to colloidal gold [32], which can protect gold nanoparticles from aggregation under high salt conditions. However, when the amount of aptamers is too high, the gold nanoparticles will not agglomerate in high concentration salt solution and the color will not change obviously, which will eventually lead to elevating the detection limit. As shown
in Figure 7, when the aptamer concentration was in the range of 0-1.0 μM, the absorbance value increases obviously and the color changes from gray to purple-gray, which indicated that aptamer was not enough to protect the aggregation and precipitation of gold nanoparticles under high salt concentration. When the aptamer...
### Table 2: The sequences and the features of the selected representative candidate aptamers.

| Sequence name | Sequence 5’-3’ | Length | Free energy (kcal/mol) | Copy |
|---------------|----------------|--------|------------------------|------|
| seq-1         | ATAGGAGTCAGACGACACCAGCATTCCGTTGATCCCTGAGAATAGAAGTAAGTTCTTTGTTGATGTGCGTCTACCTCTGA | 79   | -4.8                   | 9    |
| seq-2         | ATTAGGAGTCAGACGCACCCAGAAGATCATGGTGACCGGATCCAATTCTATTTATATGTCATTTATGTCGCTACCTCTTTGA | 81   | -4.5                   | 2    |
| seq-3         | ATAGGAGTCAGACGACACCAGCATTCCGTTGACTGTGTAGTGTATGTTTGGAGTTGTATGTGCTACCTCTTG | 79   | -4.4                   | 2    |
| seq-4         | ATAGGAGTCAGACGACACCAGCATTCTAGTGTACAGTGTACAGTCAGTGCTACCTCTTTGA | 80   | -6.3                   | 2    |
| seq-17        | ATAGGAGTCAGACGACCCAGCAGCTGACCTTTTCTGTGTAGTGTACAGTGTACAGTCAGTGCTACCTCTTTGA | 80   | -10.4                  | 1    |
| seq-41        | ATAGGAGTCAGACGACCCAGCAGCTGACCTTTTCTGTGTACAGTGTACAGTCAGTGCTACCTCTTTGA | 80   | -11.6                  | 1    |
concentration was equal to or higher than 2 μM, the color of colloidal gold solution changed from purple-gray to red, and the absorption value tended to be stable. These

data are presented in Table 3.

**Table 3: Equilibrium dissociation constant (K_d) of the six selected candidate aptamers.**

| Sequence | K_d (nM) | R^2  |
|----------|----------|------|
| seq-1    | 37.94 ± 0.1445 | 0.9841 |
| seq-2    | 130.1 ± 0.2984  | 0.9712 |
| seq-3    | 135.1 ± 0.4183  | 0.9796 |
| seq-4    | 57.16 ± 0.2055  | 0.9622 |
| seq-17   | 98.03 ± 0.2510  | 0.9783 |
| seq-41   | 31.17 ± 0.1070  | 0.9857 |

Figure 4: Binding affinity of the six selected candidate aptamers to ABL. Nonlinear regression analysis was performed using GraphPad Prism 6.01.

Figure 5: Specificity of the selected candidate aptamers seq-1 and seq-41 towards ABL. The absorbance values of seq-41 and seq-1 were measured at 450 nm. ABL, ABE, and BSA are the abbreviations of *Agaricus bisporus* lectin, *Agaricus bisporus* extract, and bovine serum albumin, respectively.

Figure 6: Optimization of the NaCl concentration. The embedded image showed the color changes (from left to right) with the increase of the NaCl concentration.
evidences suggested that 2 μM aptamer was appropriate for further experiments.

3.4.3. Setting Up a Standard Curve of the Aptamer-AuNPs Colorimetric Method for Detection of ABL. In the aptamer-AuNPs colorimetric system, ABL and AuNPs competitively bind to aptamers. Under the condition of a high concentration salt solution, lectin binds with aptamer, thus releasing the free AuNPs and exposing them to the high salt solution to aggregate and precipitate. Meanwhile, the color change of AuNPs can be observed by naked eyes and monitored by UV-vis spectroscopy. As shown in Figure 8, with the increase of ABL concentration, the ratio of absorbance A620/A520 gradually increased, revealing that there was a good linear relationship between them. Based on these results, a standard curve was generated (Figure 8). The linear regression equation was $y = 0.4146x + 0.4307$ ($R^2 = 0.9964$) within a linear range of 0.08–1.70 μg/mL and the limit of detection of this method can reach 0.062 μg/mL. The limit of detection (LOD) was calculated according to the definition of LOD (LOD = 3Sb/m, n = 9; Sb: the blank standard deviation; m: the slope of the linear regression equation) in a standard solution. The results demonstrated that the colorimetric method could be used to quantitatively detect ABL. Currently the main method for determination of ABL activity (or ABL concentration) is the hemagglutination activity test. According to the ABL information provided by Sigma-Aldrich, the activity of ABL is defined as the lowest concentration to agglutinate a 2% suspension of human blood group O erythrocytes after 1 h incubation at 25°C. Agglutination activity is usually expressed in μg/mL of lectin and is determined from serial dilutions in phosphate buffered saline, pH 6.8, of a 1 mg/mL solution (https://www.sigma-aldrich.com/catalog/product/sigma/l5640?lang=zh &region=CN). That is to say, the limit of detection for ABL is usually at the level of μg/mL. EY Laboratories, Inc., USA, states on its web page (http://eylabs.com/agaricus-bisporus/) that 5–10 μg/mL ABL causes agglutination of type A or B erythrocytes and 2 μg/mL will agglutinate type O erythrocytes. Alborés et al. reported that the minimum concentration of Agaricus bisporus lectin detected by mouse erythrocytes with hemagglutination test was 7.8 μg/mL [33]. Comparing to these facts, the detection limit of aptamer-AuNPs colorimetric method constructed in this study is about one hundred times as high as that of the hemagglutination method, which indicated that the method had much high sensitivity. Moreover, no one has reported the isolation of ABL aptamer as well as the ABL assay method based on the aptamer of ABL. Consequently, as we know, this is the first report about the establishment of the aptamer-AuNPs colorimetric method for detection of ABL.

3.4.4. Specificity Verification of the Aptamer-AuNPs Colorimetric Method for Detection of ABL. Some foods consumed daily may contain more than one kind of mushrooms. To explore the selectivity of the aptamer-AuNPs colorimetric method for detection of ABL, the standard ABL, lectin extracts from Agaricus bisporus, Flammulina velutipes, and Lentinus edodes were tested with the method. The results...
showed that the ratio of A620/A520 of standard ABL and fresh ABL extract (ABE) was dramatically higher than that of other samples, highly indicating that this method can be applied to the detection of ABL with high specificity and sensitivity (Figure 9). This may provide a potential assay method for detection of ABL from foods and agricultural products.

4. Conclusions

In conclusion, in this study, aptamer seq-41 was selected from the ssDNA pool after 13 SELEX selection rounds and it was proved that seq-41 had high affinity and specificity to ABL. Based on seq-41, an aptamer-AuNPs colorimetric method for the qualitative and quantitative detection of ABL was established. Compared to the hemagglutination assay method, this method has high sensitivity and specificity, which provides a reference for the future research and development of the ABL detection method.

Data Availability

All data generated or analyzed during this study are included in this article.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

Authors’ Contributions

Hao Gao and Yiling Tian contributed equally to this manuscript.

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