Design of Hapten Synthesis and Antibody Characterization of G-group Aflatoxins

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Food and feed contamination with Aflatoxins pose a serious threat to human health and animal husbandry development and has caused widespread concern, among them, G-group Aflatoxins as the main pollutant has attracted more and more attention. In order to establish a rapid, sensitive, specific and efficient immunoassay method for G-group aflatoxins, this study aimed to designed to synthesize 3 immunogens and coating antigens and identified by UV and SDS-PAGE. Then used to immunize Balb/c mice with prepared of three immunogen the titers were determined by indirect ELISA and the sensitivity was determined by competitive indirect ELISA (cELISA), the specificity was assessed by the cross-reaction test (CR). The results of UV and SDS-PAGE showed that the three immunogens and the corresponding coated antigens were successfully synthesized and the best one was SA method among three synthesis methods of G-group AF artificial antigen and its conjugation ratio of AFG1 to BSA was about 5.64. The results of ELISA showed that the antibody titers obtained by each method were higher than 1:6.4×10^5, and the specificity of the antibodies was consistent with the results of cELISA. The experimental results not only obtained the ideal G group aflatoxin antibody, but also established a substance and technology foundation for G group aflatoxin immunization methods, and can be referenced in the similar tests.

Key words: G-group Aflatoxins, Hapten, Artificial Antigen, Polyclonal Antibody, Specificity Analysis.

Abstract

Aflatoxins (AFs) has acute, chronic, carcinogenic and immunosuppressive toxic effects on human health. Among them, AFB1 has the strongest toxicity and wide pollution, accounting for more than 50 %. Most countries have made clear provisions on the maximum residue limit (MRL) of AFB1 in food (Wu et al., 2013; Rushing & Selim, 2019). AFB1 produced under natural conditions is in the form of a fat-soluble substance, which is difficult to purify and extract. Therefore, the effective immunoassay method is to prepare single antibodies with high 

Introduction

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sensitivity and specificity for B group AF and G group AF respectively. After that, form mixed universal antibodies, but the high quality antibody is prepared from semi-antigen molecular design and antigen synthesis (Gefen et al., 2015). The synthesis methods of G AF antigen have been reported, but the research on comparative analysis of different semi-antigen synthesis, antigen synthesis and antibody characteristics has not been reported. In this study, AFG1 was reactive starting raw materials, which was designed to synthesize artificial antigen by different AFG1 semi-antigen molecular design and antigen synthesis methods, and prepared polyclonal antibody (pAb), analyze its characteristics, and select the best hapten molecular design and antigen synthesis method. In combination with antigen synthesis, the preparation of G-group AF high-quality antibodies with strong specificity, high affinity, and it laid a foundation for the establishment of TAFS immunoassay.

Materials and methods

Reagents. AFB1, AFB2, AFG1 and AFG2 standard products, products of german prio lab company, purchased from Zhengzhou Yingke reagent consumables business department; Bovine serum albumin (BSA), ovalbumin (OVA), Freund’s complete adjuvant (FCA), Freund’s incomplete adjuvant (FIA), products of Pierce Company Purchased from Yunke reagent consumables station, Zhongyuan District, Zhengzhou; and goat anti-mouse IgG conjugated with horseradish peroxidase (GaMIgG-HRP) were obtained from Huamei Biological Engineering Co., Ltd. Other reagents were commercially available, and were analytical or chromatographic pure, the test water is triple distilled water.

Solutions. The diluent used in the enzyme-linked immunosorbent assay (ELISA) is 0.01 mol·L⁻¹ pH 7.4 phosphate buffer solution (PBS); the washing solution is PBS containing 0.5 g/L Tween-20 (PBST); Blocking buffer containing swine serum (5 %, v/v) in PBST; the coating solution is 0.1 mol/L carbonate buffer solution (CBS) with pH 9.6.

Experimental instrument. Pharmacia protein nucleic acid analyzer, Amersham company; Wd-9403d ultraviolet instrument, Beijing Liuyi Instrument Factory; RF-5301 PC fluorescence spectrometer, Shimadzu company, Japan; Jy-3000 electrophoresis instrument, Beijing Junyi Dongfeng electrophoresis equipment Co., Ltd; F-4500 fluorescence spectrophotometer, Hitachi, Japan; Multifunctional automatic microplate reader (MK3), was purchased from Thermo in USA.

Experimental animals. SPF 4-week-old female Balb/C mice, Provided by Experimental Animal Center of Xinxiang Medical College, Animal License Number: SCXK (Yu) 2010-0002.

The hapten molecular design and artificial antigen synthesis of G-group AF. AFG1 was selected as the starting material, according to the active groups and sites on its molecular structure (see Figure 1), and on the basis of reference to relevant literature, the molecular design of AFG1 hapten and the synthetic methods of artificial immunogen were mainly semi acetal (SA) and epoxide (EP), enol ether derivative (EED) and other methods to synthesize AFG1-BSA.

**Fig. 1. Molecular structure of AFG1**

**SA method.** AFG1 is converted to AFG2a under the action of H2SO4, the aldehyde group of afg2a and the amino group of BSA form unstable Schiff base. Under the reduction action of NaBH4, which formed a stable AFG2A-BSA (Chu et al., 1985; Li et al., 2015). The synthesis route is shown in Figure 2.

**Fig. 2. The synthetic route of AFG1-BSA by SA method**

**EP method.** Using the double furan ring double bond of AFG1, under the action of oxidation, the 3 and 4 positions of AFG1 form epoxides, react with – NH2 of BSA to form secondary amines, and couple with BSA in the form of – CONH – to synthesize AFG1-BSA. The synthesis route of (Martin & Garner, 1977; Zhang et al., 2009) is shown in the figure 3.

**Fig. 3. The synthetic route of AFG1-BSA by EX method**

**EED method.** The AFG1 4-bit active site is obtained by adding an ethanolic acid attachment arm to obtain an AFG1-Ga, and the carboxyl group of AFG1-Ga was conjugated with BSA (Iyer & Harris, 1993). The synthesis route of is shown in the figure 4.
Identification of G-group AF artificial antigen. UV identification. A certain amount of AFG1, BSA and AFG1-BSA, and AFG1 were weighed and dissolved with methanol the methanol as a blank control; BSA and AFG1-BSA were dissolved in methanol PBS solution \[V(\text{methanol}):V(\text{PBS})=4:6\] , which was used as blank control. UV scanning was performed at 220 ~ 450 nm and analyze the scanning atlas, refer to the method of Wang Yanan et al. calculate the molecular binding ratio of AFG1 and BSA (Wang et al., 2014; 2016).

SDS-PAGE identification. The concentration of concentrated glue is 50 g/L, voltage 90 V; separation glue mass concentration is 120 g/L, voltage 45 V; 10 μL per hole per hole, 10 μg of protein, Coomassie blue staining, the molecular binding ratio of AFB1 and BSA was calculated by UV gel imaging system analysis software.

Fluorescence intensity identification. A certain amount of BSA and AFG1-BSA were weighed and dissolved in methanol PBS solution \[v(\text{methanol}):v(\text{PBS})=4:6\] , the mass concentration was 1.0 mgmL^{-1}, and the fluorescence intensity was measured at the excitation wavelength of 365 nm and emission wavelength of 440 nm.

Analysis of Preparation and Immunological Characteristics of AFG1 pAb. Balb/C mice were immunized with the antigen AFG1-BSA synthesized by three different methods, and each antigen was immunized with 1 group. There were 3 groups with 5 mice in each group. The immunization dose was calculated according to the amount of BSA in AFG1-BSA, 50 μg for each, with a volume of 0.2ml. Subcutaneous injection was performed at 4 ~ 6 points on the back for 5 times, with a time interval of 4 w. The suborbital sinus of the eyeball was removed 21 days after the last immunization, and blood samples were collected, and serum was separated to obtain AFG1 pAb.

Analysis of immunological characteristics of AFB1 pAb. The titer was determined by indirect ELISA (Zhang et al., 2008). Sensitivity was determined by indirect competitive ELISA (icELISA) to determine the 50 % inhibition concentration (IC_{50}) of AFG1 pAb to AFG1, and sensitivity was measured by IC_{50} (Li et al., 2015). Specificity identification, AFG1, G2, B1 and B2 were selected as inhibitors by cross reaction test. The IC_{50} of each inhibitor was determined by icELISA. The cross-reaction rate (CR%) was the percentage of IC_{50} of AFG1 PAB to AFG1 and IC_{50} of AFG2, AFB1 and AFB2.

Results

UV identification. It can be seen from Figure 5 that the maximum absorption peak of BSA is at 278 nm, AFG1 has four absorption peaks at 243, 257, 264 and 362 nm between 200 and 500 nm, and the maximum absorption peak is at 362 nm. AFG1-BSA (SA) and AFG1-BSA (EP) have characteristic absorption peaks at 346 nm and 413 nm, and AFG1-BSA (EED) has characteristic absorption peaks at 423 nm. The three methods all have different UV absorption characteristics from BSA and AFG1, indicating that the artificial antigen AFG1-BSA was successfully synthesized by the three methods. The calculated molecular binding ratio of AFG1 and BSA was 5.64:1 (AFG1-BSA(SA)), 7.17:1 (AFG1-BSA(EP)) and 3.43:1 (AFG1-BSA(EED)).

SDS-PAGE identification. It can be seen from Figure 6 that the migration rate of the immunogen AFG1-BSA synthesized by the three methods on the gel plate is less than that of BSA, indicating that the MW of AFG1-BSA is greater than that of BSA, this indicates that the synthesis of AFG1-BSA is successful.
Fluorescence intensity identification. It can be seen from Figure 7 that under the same excitation light source and the same protein mass concentration (1.0 mg/mL) conditions, compared with BSA, the antigens AFG1-BSA (SA), AFG1-BSA (EP), and AFG1-BSA (EED) synthesized by the three methods have enhanced fluorescence intensity to different degrees, indicating that the complete antigen synthesis was successful.

![Figure 7. The measurement of fluorescence intensity of artificial antigen AFG1-BSA](image)

Titer determination. It can be seen from Figure 8 that after 5 immunizations, 1 mouse with the highest titer was selected from each group, and a total of 3 mice were selected. The titer was measured by indirect ELISA and compared. The AFG1 pAb titer of 3 mice all reached 1:(1.6×10³) or more, it shows that the three synthetic complete antigens AFG1-BSA all have good immunogenicity. The immune effect of the three groups is evaluated according to the titer, and the order is AFG1-BSA (SA), AFG1-BSA (EP) and AFG1-BSA (EED).

![Figure 8. The titer curves of AFG1 pAb](image)

Sensitivity analysis. It can be seen from Figure 9 that the icELISA curve of the 3 mice with the highest titer selected after immunization showed a good linear relationship. The $A_{50}$ value was converted to $B/B_0\%$, and then the regression analysis was performed on $Lg [AFG1/100]$, all meet the criteria of linear relationship. The regression equation, $R^2$ value and $IC_{50}$ value of the three mice's serum AFG1 pAb inhibition curve are shown in Table 2. Among them, the serum of the AFG1-BSA (SA) group mouse has the best inhibition effect, with an $IC_{50}$ of 17.43 μg/kg⁻¹. Followed by the $IC_{50}$ of the AFG1-BSA (EP) group, which was 25.11 μg/kg⁻¹, and the AFG1-BSA (EED) group had a poorer sensitivity, which was 70.41 μg/kg⁻¹.

![Figure 9. The inhibition curves of AFB1 pAb against AFB1](image)

Specificity analysis. It can be seen from Table 1 that the AFG1 pAbs obtained after the three coupling methods synthesize antigens to immunize animals can recognize AFG1 100 %, and AFG1 pAb (SA) has the best sensitivity, with $IC_{50}$ of 17.43 μg/kg⁻¹, the broad spectrum is also the best, the CR with AFG2 is 94.47 %, and the CR of AFB1 and AFB2 are both less than 10 %. AFG1 pAb (EP) has good sensitivity, with $IC_{50}$ of 25.11 μg/kg⁻¹, but it has poor broad-spectrum, and its CR with other AFs is less than 10 %. AFG1 pAb (EED) has poor sensitivity to AFG1, with $IC_{50}$ of 70.41 μg/kg⁻¹. The results show that AFG1-BSA prepared by hemiacetal method has high sensitivity, strong specificity and best broad spectrum.

| AF | AFG1 pAb (SA) | AFG1 pAb (EP) | AFG1 pAb (EED) |
|----|---------------|---------------|---------------|
| IC₅₀ / (μg·kg⁻¹) | Cross-reactivity | IC₅₀ / (μg·kg⁻¹) | Cross-reactivity | IC₅₀ / (μg·kg⁻¹) | Cross-reactivity |
| B1 | >1×10⁹ | <10 | >1×10⁹ | <10 | >1×10⁹ | <10 |
| B2 | >1×10⁹ | <10 | >1×10⁹ | <10 | >1×10⁹ | <10 |
| G1 | 17.43 | 100 | 25.11 | 100 | 70.41 | 100 |
| G2 | 18.45 | 94.47 | >1×10⁹ | <10 | >1×10⁹ | <10 |

![Table 1](image)
**Discussion**

**About AF artificial antigen synthesis pathway.** High-quality antibodies are the core reagents for establishing immunoassay methods. In terms of current research progress, there are two ways to realize immunoassay of total TAFs. One is to prepare single antibodies with high sensitivity and strong specificity for group B AFs and group G AFs, and then mix and use mixed universal antibodies. This method is currently more commonly used and has achieved good application effects. The second is to prepare a single universal antibody that can simultaneously recognize AFB1, AFB2, AFG1, and AFG2 with high sensitivity and broad recognition spectrum. This is the most ideal technical method, but it is difficult to prepare such high-quality monoclonal antibodies. Since AF is a small molecule hapten, it is necessary to synthesize an artificial immunogen to assist the proliferation and differentiation of B cells with the help of activated T cells to obtain qualified antibodies.

**About G group AF artificial immunogen synthesis method and immune effect analysis.** Due to the late start of the study of G-group AF hapten molecular design, antigen synthesis methods and the characteristics of the antibodies produced, comprehensive analysis of the above three methods, the author believes that the hemiacetal method is the most effective. AFG1 is acidified to form AFG2a, using the 3-hydroxyl active group of AFG2a to synthesize antigen with Schiff's base as the spacer arm, the produced antibody has the characteristics of high titer, good sensitivity, strong specificity and broad recognition spectrum. The epoxide method can induce the body to rapidly produce antibodies, but its sensitivity is not as good as the hemiacetal method. The enol ether derivative method has obvious defects, the ability to induce the body to produce antibodies is poor, and the antibodies produced are less sensitive to AFG1. Therefore, this method has academic value for scientific research, but it is rarely applied in actual production.

**About the development trend of AF artificial immunogen synthesis method.** Due to the AFB1 is closely related to other AF pollution, AFB2 pollution is accompanied by AFB1, and high concentrations of AFB1 have an inhibitory effect on AFG1 and AFG2. Therefore, there are two immunological detection and evaluation methods for AF contamination in food. Some countries, including China, adopt AFB1 limit standards. However, in order to solve the simultaneous existence of multiple toxin pollution and the toxic additive effect and the corresponding lack of detection standards, some countries adopt the total amount of AF (B1+B2+G1+G2) limit standard. In view of this, the focus of the research of AF hapten molecular design, antigen synthesis method and immune effect analysis is to improve the sensitivity and standard, in the meanwhile, improve the sensitivity and broad recognition spectrum of antibodies to AFB1, AFB2, AFG1, and AFG2, and meet the requirements for the total amount of AF. At present, the molecular design and antigen synthesis of AF hapten still remain at the level of empirical design and predictive design, and trial-and-error assays are mostly used. Although a variety of identification methods such as infrared (IR), ultraviolet (UV), mass spectrometry (MS), gel electrophoresis (SDS-PAGE) and nuclear magnetic resonance (NMR) have been established for synthetic artificial antigens, but in the end, it is necessary to verify its scientific rationality through animal experiments, there will be blindness and contingency to a certain extent (Guan et al., 2013). With the development of new disciplines such as molecular immunology, quantum chemistry, and molecular mechanics, as well as the popularization and application of molecular information technology, molecular simulation technology, and computer-aided technology, these provide theories and methods for reference to improve the rationality, timeliness and predictability of hapten molecular design and antigen synthesis (Xu et al., 2009; Kortkhojja et al., 2013; Sevy & Meiler, 2014). Kim et al. (2011) used computer-aided technology to prepare TAFs mAbs with strong specificity and broad recognition spectrum. The IC50 for AFB1, AFB2, AFG1, and AFG2 were 4.36, 7.22, 6.61, 29.41 μg/kg\(^{-1}\), respectively, ELISA detection technique to achieve TAFs mode. Zhou Qian et al. (2010) successfully prepared G-group AF mAb with the help of molecular simulation technology, with IC50 of 17.18 and 19.75 μg/kg\(^{-1}\) for AFG1 and AFG2, respectively, and established an AF G-group ELISA detection method.

**Conclusion**

This study refers to a large number of literatures, according to the molecular structure and active site of AFG1, the SA method, EP method and EED method are used to prepare the G group AF artificial immunogen AFG1-BSA. The synthetic artificial antigens were identified by UV, SDS-PAGE and fluorescence intensity measurement. The results showed that among the three methods for the synthesis of G-group AF artificial immunogens, the SA method had the best effect, and the molecular binding ratio of AFG1 to BSA was 5.64:1, the AFG1 pAb produced by animal immunization has the characteristics of high titer, sensitivity, specificity, and broad spectrum. The research results establish a basis for the AF Group G immunoassays and the total amount of AF.

**Author Contributions**

Yanan Wang participated, original draft preparation, writing, and software. The author has read and agreed to the published version of the manuscript.

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Conflict of interest
The author declare no conflict of interest.

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