Paper-based ELISA Diagnosis Technology for Human Brucellosis Based on a Multiepitope Fusion Protein

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Research Article

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Abstract

**Background:** At present, as a serious zoonotic infectious disease, the incidence of brucellosis is increasing each year worldwide, exhibiting signs of resurgence. Brucellosis seriously threatens the health of humans, and it is necessary to strengthen the methods utilized for its rapid and accurate diagnosis.

**Methods:** Bioinformatic technology was used to predict B-cell epitopes of the main outer membrane proteins of Brucella and subsequently verified the antigenicity of these epitopes. Prepared a Brucella multiepitope fusion protein and verified the antigenicity of the protein by indirect ELISA. Whatman filter paper was then modified with nano-zinc oxide to construct a paper-based ELISA (p-ELISA) technology for the diagnosis of brucellosis.

**Results:** A total of 22 linear B cell epitopes were predicted. Each epitope could recognize some brucellosis sera. The constructed multiepitope fusion protein had good antigenicity and significantly reduced cross-reaction compared with LPS. The sensitivity and specificity of the method were 92.38% and 98.35%, the positive predictive value was 98.26%, and the negative predictive value was 91.67%.

**Conclusions:** A multiepitope fusion protein of Brucella was successfully prepared, and a rapid diagnostic technique for brucellosis was established. This technology has potential application value and can be used for the rapid diagnosis of brucellosis.

Background

Brucellosis is a reemerging zoonotic infectious disease. In recent years, the incidence rate has been increasing yearly. It not only seriously threatens the health of the people but also causes huge economic losses to the animal husbandry industry. Brucellosis has a complex condition and a long course of disease, causing a huge economic burden and a waste of medical resources in countries all over the world, especially after misdiagnosis, which will increase the treatment cost[1]. Therefore, the rapid and accurate diagnosis of brucellosis is necessary. At present, the diagnostic methods for brucellosis include traditional pathogenic detection, serological diagnostic techniques and molecular biology methods[2, 3].

The traditional isolation culture method requires complicated laboratory and medium conditions, which need to be carried out in a P3 laboratory, and the culture time is long, usually several weeks[4]. Molecular biology methods such as PCR technology have specific and rapid characteristics, but nucleic acid contamination will cause false positive results, the sensitivity and specificity are not high, and it requires relatively expensive instruments and professional operation, which is not conducive to popularization in grassroots units[5]. Serological diagnostic techniques mainly include the agglutination test, complement fixation test (CFT), enzyme-linked immunosorbent assay (ELISA), immunochromatographic diagnostic test (ICDT), and fluorescence polarization assay (FPA), which are currently the commonly used screening methods for brucellosis[6]. This method has the advantages of high sensitivity and short operation time, but the diagnosis of serological methods is greatly affected by antigens. The key technology to improve the sensitivity and specificity of serological diagnostic methods is to find suitable diagnostic antigens.
Paper-based enzyme-linked immunosorbent assay (p-ELISA) is an emerging detection technology. It has the same principle as the traditional 96-well plate ELISA except that it uses paper as a solid phase carrier. Due to its small reagent dosage, rapid detection, low cost, and lack of need for special equipment, it has attracted increasing clinical attention\[7, 8\]. The main advantages of paper as a carrier are as follows: wide range of paper sources and low cost; through capillary action, no external force is needed to make the liquid flow; good biocompatibility; small reagent loading volume; and easy to use and carry\[9\].

In this study, we used immunoinformatic technology to predict the B cell epitopes in the major outer membrane proteins of Brucella, synthesized polypeptides and coupled them with KLH, screened these polypeptides by traditional ELISA methods, selected effective polypeptides as diagnostic antigens, and established a nano-ZnO-modified p-ELISA for brucellosis diagnosis based on a multiepitope fusion protein.

**Methods**

**Human serum samples**

There were 121 human brucellosis sera (gifted by the School of Public Health of Jilin University), 90 control sera, including 50 healthy sera and 40 patient sera (confirmed by blood culture to be infected with other pathogens, collected by the infection department of the First Clinical Hospital of Jilin University; information on the patients is shown in Table S1).

**Selection of the outer membrane proteins of Brucella**

The antigenicity of the outer membrane proteins of Brucella was investigated by consulting the literature (https://www.ncbi.nlm.nih.gov/protein/). The amino acid sequence was obtained, and the conservation of the amino acid sequence was analyzed by BLAST.

**B cell epitope prediction and peptide synthesis**

The conserved amino acid sequence of the Brucella outer membrane protein was used to predict B cell epitopes by using the B cell epitope prediction tool bepipred linear epitope prediction 2.0 in IEDB (http://tools.iedb.org/bcell/). The predicted B cell epitopes were delivered to SGS and coupled with keyhole limpet hemocyanin (KLH). The predicted B cell epitope was handed over to Sangon Biotech (Shanghai, China) for synthesis and coupled with keyhole limpet hemocyanin (KLH).

**Screening of peptide epitopes**

To screen effective peptide epitopes, we verified the antigenicity of predicted epitopes by indirect ELISA. The experimental procedure was as follows: peptides coupled to KLH were diluted with carbonate buffer (pH = 9.6) to final concentrations of 30 μg/mL and 100 μL/well in a 96-well plate (Corning, USA) and incubated overnight at 4°C. Next, 300 μL/well of blocking solution (PBS containing 5% skimmed milk powder) was incubated at 37°C for 1 h, and the cells were washed 3 times with PBST (PBS containing
0.05% Tween 20). Afterwards, 100 µL/well of 1:400 serum was added and incubated at 37°C for 1 h followed by washing 3 times with PBST. Next, 1:5000 diluted HRP-labeled protein G (Thermo, USA) was added; it was reacted at room temperature for 15 min and washed again with PBST 3 times. Next, 100 µL of TMB substrate solution was added to each well and reacted for 15 min at room temperature followed by the addition of 50 µL of stop solution (2 M H₂SO₄). The optical density was measured at 450 nm (OD450) using an ELISA plate reader (BioTek, USA). At the same time, KLH (30 µg/mL) and lipopolysaccharide (LPS, 1 µg/mL, provided by China Animal Health and Epidemiology Center (Qingdao, China)) were used as blank carriers and positive antigen controls to detect serum.

**Fusion protein preparation**

The selected effective peptides are connected in series, and the adjacent two peptide chains are connected with the linker 'GGGS'. The plasmid was constructed by full gene synthesis, subcloned into the expression vector pET-21a (Sangon Biotech, Shanghai, China) and further transformed into E. coli competent BL21(DE3) cells (Sangon Biotech, Shanghai, China). The cells were cultured, IPTG was used to induce expression, bacteria were collected, the protein was purified, and the target protein was verified by SDS-PAGE and Western blotting. The specific steps are described below.

After transferring the recombinant plasmid into BL21(DE3), 800 µL of nonresistant LB medium was added, followed by incubation at 37°C for 45 min and centrifugation at 5000 rpm for 3 min. Most of the supernatant was discarded (leave approximately 100-150 µL), the bacteria were resuspended, the LB plate with corresponding resistance was selected, and it was coated. After air-drying, it was inverted and cultured overnight in a 37°C incubator. The monoclonal colonies on the plate were chosen, placed into 10 mL of LB liquid medium and incubated at 37°C and 200 rpm. The cultured bacterial solution was transferred to 750 mL of LB liquid medium at 37°C and 200 rpm, cultured to OD₆₀₀=0.6-0.8 with IPTG (0.5 mM) at 16°C and induced overnight. Then, the cells were centrifuged at 6000 rpm for 5 min, the supernatant was discarded, and the bacteria were collected. Bacteria were blown away with 20-30 mL 10 mM Tris-HCl (pH = 8.0) solution and ultrasonically broken (500 W, 60 times, 10 s each time, 15 s interval). After sonication, 100 µL of the bacterial suspension was centrifuged at 12000 rpm for 10 min, and 50 µL of supernatant was transferred to another EP tube. After the supernatant was removed, the precipitate was blown away with 50 µL of 10 mM Tris-HCl (pH = 8.0) solution. SDS-PAGE and Western blotting were used to detect protein expression. A nickel column (Ni Sepharose 6 Fast Flow, GE Healthcare) for affinity chromatography was used for protein purification. Taking 5 mL of Ni-NTA, the equilibrium column was washed with 5 times the column bed volume of binding buffer at a flow rate of 5 mL/min. The crude protein was incubated with the equilibrated column packing for 1 h; the incubated product was loaded onto the column and the effluent liquid was collected; the equilibrium column was washed with binding buffer; the column was washed with washing buffer, and the effluent liquid was collected; with the column was eluted with elution buffer, and the effluent liquid was collected; and the crude protein was treated, washed with effluent and eluted with effluent separately, followed by sample preparation and, SDS-PAGE and WB detection. The concentrated protein was divided into 1 mL/tube and stored at -80°C.
**Antigenicity identification of fusion protein**

Indirect ELISA was used to verify the antigenicity of the fusion protein. The procedure was as follows: 96-well microtiter plates were coated with 1 μg/well fusion protein at 4°C overnight, and then, 5% skimmed milk powder was blocked at 37°C for 2 h. Serum was diluted 1:400 and added at 100 μL per well, followed by incubation at 37°C for 1 h. HRP-labeled protein G was diluted with 1:8000, added at 100 μL/well, incubated at 37°C for 1 h, and finally developed with TMB substrate solution for 15 min. For termination, 2 M H₂SO₄ was used, and the OD450 was measured. After each step, the cells were washed with PBST (0.05% Tween-20 in PBS) 3 times.

**Synthesis of nano-ZnO and paper modification**

ZnO nanorods were synthesized on Whatman No. 1 filter paper by a hydrothermal method[10]. The steps were as follows: Whatman filter paper was soaked in 100 mm zinc acetate solution for 60 s and then annealed at 100 °C for 1 h to form a seed layer (seed layer). Then, the filter paper with a seed layer was transferred to a hydrothermal reaction vessel containing an equimolar solution (100 mm, pH = 6.5) of hexamethylenetetramine (HMTA, Sigma) and zinc nitrate (Zn(NO₃)₂.6H₂O, Sigma). ZnO nanorods were synthesized at 90 °C for 5 h. Next, Whatman filter paper with ZnO nanorods was immersed in anhydrous toluene solution (Sigma) with 1% APTES (Sigma) for 5 min, heated and dried at 100 °C for 15 min, and silanized. Scanning electron microscopy (SEM, JSM-7500F), X-ray diffraction (XRD, Bruker D8) and X-ray photoelectron spectroscopy (XPS, Escalable250Xi) were used to characterize the structure and surface of the paper. Whatman filter paper modified with ZnO nanorods was punched into circular paper pieces with a diameter of 10 mm by a punch, and A4 plastic packaging paper was punched into small holes with a diameter of 6 mm by a punch. The 10 mm filter paper pieces were placed in the center of the 6 mm holes of the plastic packaging paper, fixed by a plastic packaging machine, and cut into small strips with 3 holes for standby.

**Establishment of p-ELISA**

Five microliters of fusion protein solution was placed in each well (30 μg/mL in PBS), incubated at room temperature for 30 min, washed with 20 μL of deionized water 3 times, and blocked with 20 μL of 5% skimmed milk powder at room temperature for 15 min; PBST was used for washing 3 times, and 5 μL of serum was added (diluted with 1:400); PBST was used for washing 3 times, and 5 μL of HRP labeled protein G was added (diluted with 1:8000), followed by incubation at room temperature for 210 s; and PBST was used for washing 3 times, 5 μL of TMB substrate solution was developed for 10 min, and an HP Laser Jet Pro MFP M227 was used for scanning to obtain the image. ImageJ software carries out gray intensity analysis for quantitative analysis.

**Traditional p-ELISA**

To compare with the nanomodified p-ELISA (nano-p-ELISA) method, we also performed the traditional p-ELISA (tra-p-ELISA) method. The specific steps have been described in the literature[11]. Five microliters
of chitosan was added to deionized water (0.25 mg/mL) and placed onto Whatman No.1 filter paper followed by drying at room temperature; then 5 μL of 2.5% glutaraldehyde solution (PBS) was added, followed by resting at room temperature for 2 h and washing with 20 μL of deionized water twice. The remaining steps are same as described in section 2.8.

Statistical analysis

Dot plot and receiver operating characteristic (ROC) curve analyses were performed using GraphPad Prism version 6.05 for Windows. The OD450 and gray intensity were determined by Student's t-test (unpaired t-test). *P*-values < 0.05 were considered to indicate significant differences.

Results

Brucella outer membrane protein epitope prediction and peptide synthesis

Five highly conserved proteins, omp16, omp25, omp31, omp2b and BP26, were selected. All information on Brucella species and protein accession numbers are shown in Table S2. A total of 22 epitopes were predicted for the selected proteins, and the detailed epitope information is shown in Table 1. The 22 polypeptide epitopes were synthesized and coupled to KLH. KLH was uniformly coupled to the tail end of the polypeptide (right side). Each polypeptide (10 mg) was coupled to 10 mg of KLH, and the purity was > 90%.
Table 1
Detailed information of 22 predicted B cell epitopes

| Protein | Epitope (amino acid sequence) | Start-end position | Peptide ID |
|---------|-------------------------------|--------------------|------------|
| BP26    | AFAQENQMTTQPARIAV             | 26–42              | P19266-1   |
|         | KAGIEDRDLQTGGIN               | 88–100             | P19266-2   |
|         | QPIYVYPDKNLKEPTITGY          | 104–124            | P19266-3   |
|         | GVNOQGDLNLVNDNPSAVIN          | 151–170            | P19266-4   |
|         | LSRPPMPMPMP                  | 204–212            | P19266-5   |
|         | AAAPDNSVPIAAGENSYNVSVNVVFE    | 223–248            | P19266-6   |
| Omp2b   | SGAQAADAIVAPEPEAVEY          | 31–49              | P19266-7   |
|         | DVKGGDDVYSGTDRNGWDK           | 79–97              | P19266-8   |
|         | NNSGVGKYGNETSSGTV             | 129–146            | P19266-9   |
|         | TVTPEVSYTKFGEWKNTVAEDNAWGGI  | 341–368            | P19266-10  |
| Omp16   | AAAPGSSQDFTV                 | 44–55              | P19266-11  |
|         | SRGVPTNRMRTISYGNERPVAVCD     | 125–148            | P19266-12  |
| Omp25   | GRAKLENRTNGGTS               | 56–69              | P19266-13  |
|         | GNPVQTTGETQ                  | 115–125            | P19266-14  |
|         | GGIKNSLRBIGGEGSSKSKTQT       | 154–174            | P19266-15  |
|         | GWTVGAGIEYAA                 | 175–186            | P19266-16  |
|         | TDYGKKNNFGLNDLDTRGSFKTNDIR   | 199–223            | P19266-17  |
| Omp31   | VSEPSAPTAAPVDTFSWTGGYIGINA   | 24–49              | P19266-18  |
|         | GKFHPFSSFDFKEDNEQVSGSL       | 53–75              | P19266-19  |
|         | TGSISAGASGLEGKAE             | 112–127            | P19266-20  |
|         | GDDASALHTWSDKTKAGWTLGAGAEYA  | 168–194            | P19266-21  |
|         | DLGKRNLDV                   | 109–217            | P19266-22  |

**Peptide screening**

The KLH-conjugated polypeptide was used as the antigen, and the indirect ELISA method was used to verify the antigenicity of the polypeptide. Positive serum was used to screen the polypeptide. The results
of iELISA show that each peptide can recognize some sera, and the recognition ability is quite different (see Fig. 1).

**Fusion protein preparation**

SDS-PAGE identification results showed that after 0.5 mM IPTG-induced expression overnight, the expression of the fusion protein reached an ideal amount, and the expression molecular weight was approximately 66 kDa. The results are shown in Figures 2A and 2B. After Western blot identification and analysis, the molecular weight of the histidine tag was consistent with the SDS-PAGE results, and after mass spectrometry verification, it was confirmed that the expressed protein was the target protein. The detailed results are shown in Figure 2C.

**Antigenicity identification of fusion protein**

The antigenicity of the purified fusion protein was verified by iELISA. Compared with the LPS antigen, the area under the diagnostic curve of the fusion protein was 0.9877 (95% CI: 0.9758 to 0.9996), while the area under the LPS curve was 0.9174 (95% CI: 0.8796 to 0.9552) (the results are shown in Figure 3). The optimal cutoff value was calculated by the Youden index, the positive predictive value and negative predictive value of the fusion protein were higher than those of LPS, and the diagnostic predictive value of the two antigens under this cutoff value was analyzed (Table 2).

| Cutoff value            | Positive | Negative | PPV (%) | NPV (%) |
|-------------------------|----------|----------|---------|---------|
| ≥0.470 (fusion protein) | 117      | 4        | 95.90   | 95.51   |
| ≥0.4095 (LPS)           | 115      | 6        | 85.19   | 92.10   |
| ≥50.98 (nano-p-ELISA)   | 113      | 8        | 98.26   | 91.67   |
| ≥45.66 (tra-p-ELISA)    | 113      | 8        | 97.41   | 91.58   |

TP, true positives; TN, true negatives; FP, false positives; FN, false negatives; PPV, positive predictive value (TP/TP+FP)×100; NPV, negative predictive value (TN/TN+FN)×100.

**Synthesis and characterization of nano-ZnO**

Scanning electron microscopy showed that we synthesized nano-zinc oxide on the surface of Whatman filter paper (Fig. 4B). At the same time, XRD results showed that the functionalization of nano-zinc oxide was successfully performed (Fig. 4A). XPS shows that the concentration of Zn atoms is 40.79% and the concentration of oxygen atoms is 59.21%, which further indicates that the concentration of zinc and oxygen in the sample is ZnO.
**Evaluation of the diagnostic effect of p-ELISA**

A total of 211 sera were detected by the p-ELISA method modified by nano-ZnO (nano-p-ELISA) and the p-ELISA method modified by chitosan and glutaraldehyde (tra-p-ELISA) to evaluate the diagnostic effects of the two methods. The gray intensity of 121 brucellosis-positive sera and 90 brucellosis-negative sera was analyzed by ROC curve analysis, and the results of the nano-p-ELISA showed that the area under the curve was 0.9900 (95% CI, 0.9816 to 0.9984), indicating that the diagnosis of this method has a very high accuracy. The optimal cutoff value was 50.98. Under this cutoff value, the sensitivity of this method was 92.38% (95% CI, 0.8554 to 0.9665) and the specificity was 98.35% (95% CI, 0.9416 to 0.9980) (Fig. 5A). Using this optimal cutoff value to further analyze the diagnostic effect, 113 of the 121 positive samples were accurately diagnosed, and 88 of the 90 negative samples were correct. The positive predictive value of the nano-p-ELISA was 98.26%, and the negative predictive value was 91.67% (Table 2). The gray intensities of the positive and negative samples were significantly different \( P < 0.001 \) (Fig. 5B).

The results of the tra-p-ELISA showed that the area under the curve was 0.9894 (95% CI, 0.9817 to 0.9970), indicating that the diagnosis of this method also has high accuracy. The optimal cutoff value is 45.66. Under this cutoff value, the diagnostic sensitivity of this method was 94.24% (95% CI, 0.8897 to 0.9748), and the specificity was 98.26% (95% CI, 0.9386 to 0.9979) (Fig. 5C). With this optimal cutoff value, 113 of the 121 positive samples were accurately diagnosed, and 87 of the 90 negative samples were correctly judged as negative. The positive predictive value of the tra-p-ELISA was 97.41%, and the negative predictive value was 91.58% (Table 2). The gray intensities of the positive and negative samples were significantly different \( P < 0.001 \) (Fig. 5D).

**Discussion**

The existing diagnostic methods for brucellosis have disadvantages such as complicated operation, long time consumption, low sensitivity, and proneness to cross-reaction\[12\]. Therefore, a simple, fast and sensitive diagnostic method is sought. Early diagnosis and early treatment of the disease are of great significance to reduce economic loss and medical burden\[13\]. The development of new diagnostic techniques for brucellosis is of great significance for the prevention and control of brucellosis.

At present, many vaccine studies have shown that many outer membrane proteins in Brucella have strong antigenicity\[14–16\]. Animal experiments have also shown that these ingredients have a certain immunoprotective effect on Brucella and are good vaccine candidates\[17, 18\]. Therefore, these components also provide a direction for researchers to develop new diagnostic antigens for brucellosis, and the development of immunoinformatic technology provides tools for the development of new diagnostic antigens. Immunoinformatic is based on bioinformatic tools, an emerging science that integrates life sciences, computer science, and mathematics\[19, 20\]. Immunoinformatic technology uses bioinformatic tools to treat pathogens without cultivating them. Processing and analysis of, for instance, the genome and proteome can be used to complete the gene prediction and the prediction of cell epitopes in the protein\[21\]. Immunoinformatic technology has the advantages of speed and economy and has
been widely used in vaccine design, disease prevention, diagnosis and treatment. In this study, we selected five Brucella antigen proteins, omp16, omp25, omp31, omp2b and BP26, and predicted 22 epitopes. The results of iELISA confirmed that each peptide can recognize a portion of brucellosis sera, but the recognition ability of each polypeptide is limited. Therefore, we concatenated 22 epitopes to synthesize a fusion protein. Using the fusion protein as an antigen, the ability to recognize serum is greatly improved. Compared with LPS, the use of fusion proteins can significantly reduce serological cross-reactions.

P-ELISA has attracted the attention of many researchers due to its advantages, such as strong specificity, simplicity, rapidity, portability, and low cost, especially in the fields of medical testing, environmental testing and food safety analysis[22, 23]. It combines the advantages of traditional ELISA and paper and provides a new method and new idea for resource-poor areas and point-of-care testing. At present, the commonly used p-ELISA is a method of modifying the surface of paper with glutaraldehyde and chitosan. Currently, nanomaterials have the characteristics of a large specific surface area and quantum size effect and have been widely used in the fields of biology and medicine[24, 25]. The modification of nanomaterials on the surface of paper can increase the surface area of the paper and establish a p-ELISA diagnostic method for nanomaterial modification[10, 26]. In this study, we modified the surface of paper with nano-ZnO and used a synthetic fusion protein as the antigen to establish a new diagnostic technique for brucellosis. The diagnostic effect is good, and it can be modified with glutaraldehyde and chitosan. The p-ELISA diagnostic method is comparable. In addition, p-ELISA paper modified by nanomaterials can be stored at room temperature for a long time, and we also found in experiments that the color development time of p-ELISA modified by nanomaterials can be significantly extended, which may be due to the antioxidant effect of nanomaterials, thus extending the effective time of the color reagent. Compared with the traditional ELISA method, the p-ELISA diagnostic method uses very few reagents and shortens the time for the entire process. It is a relatively promising on-site rapid detection technology.

Conclusions

In summary, using bioinformatic technology combined with nanomaterials, this performance has established a new type of brucellosis diagnostic technology, which has good potential application value. However, the brucellosis sera selected in this study were all clinically screened positive sera, and the number was limited. The diagnostic validity of this method requires a large number of clinical random samples for verification.

List Of Abbreviations

paper-based ELISA, p-ELISA; complement fixation test, CFT; enzyme-linked immunosorbent assay, ELISA; immunochromatographic diagnostic test, ICDT; fluorescence polarization assay, FPA; keyhole limpet hemocyanin, KLH; receiver operating characteristic, ROC
Declarations

Ethics approval and consent to participate

Not applicable

Consent for publication

Not applicable

Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests

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Authors' contributions

D-HY and Q-QB conceived and designed the study. D-HY, Q-QB, X-LW and HL performed the ELISA assays. D-HY, and S-JH analyzed the data. D-HY drafted the manuscript. M-JS, HJ and J-PZ reviewed and made improvements in the manuscript. All authors read and approved the final manuscript.

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