The Expression of apxIVA Gene of Actinobacillus pleuropneumoniae Serotype 5 and Establishment of an Indirect ELISA for Distinguishing Infection and Immunization

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Abstract: Actinobacillus pleuropneumoniae is the etiological agent of Porcine pleuropneumonia, which causes severe losses in pig farming. To establish an serological method to detect A. pleuropneumoniae infection, bioinformatics method was utilized to analyzed the sequence of apxIVA gene of A. pleuropneumoniae serotype 5 completed by our lab, a 1152 bp fragment of the N’-terminal of apxIVA gene was amplified and cloned into the prokaryotic expression vector pET-32a(+) by Ni-NTA, this protein was used to establish an indirect ELISA to detect A. pleuropneumoniae infection. This method showed high specificity and could react positively with antibodies of live A. pleuropneumoniae infection while negatively with those of inactivated A. pleuropneumoniae immunization. In conclusion, this indirect ELISA could be used to detect A. pleuropneumoniae infection in mice model.

Keywords: Actinobacillus pleuropneumoniae, ApxIVA, serotype 5, indirect ELISA, infection, immunization.

Porcine pleuropneumonia is an acute or chronic respiratory disease characterized by hemorrhagic, fibrinous, and necrotic lung lesions, and A. pleuropneumoniae is the etiological agent [1]. The disease is highly contagious and associated with all ages of growing pigs, often co-infecting with other pathogens such as Hog cholera virus, Porcine reproductive and respiratory syndrome virus and Pasteurella multocida etc, resulting in severe economic losses in swine industry. The morbidity ranges from 8.5% to 100% while the fatality ranges from 0.4% to 100%. A. pleuropneumoniae infection is more prevalent under intensive breeding conditions. To date, 15 serotypes have been reported with all serotypes can secrete apxIV which can be produced only during infection, but not under in vitro conditions [4]. ApxIV can also induce specific antibodies of high titres [5]. Based on the above character, bioinformatics method was used to analyzed the apxIVA gene sequence of serotype 5 completed by our lab, and a 1152 bp long fragment of the N’-terminal of apxIVA gene of serotype 5 was cloned and expressed to establish an indirect ELISA for detecting A. pleuropneumoniae infection in mice model.

MATERIALS AND METHODOLOGY

Bacterial Strain and Grown Conditions

E. coli JM 109, DH5a and BL21 (DE3) were purchased from Invitrogen corporation (California, USA) and cultured in LB, A. pleuropneumoniae serotype 1 strain CVCC 259 and serotype 5 strain CVCC 263 and serotype 7 strain CVCC 265 were purchased from China institute of Veterinary Drug Control (Beijing, China) and proliferated in Tryptic Soy Broth (TSB) with 0.001% nicotinamide adenine dinucleotide (NAD).

Experiment Mice

Female Kun-Ming mice (six weeks old, 17–22 g), purchased from the experimental animal center of Sichuan University (Chengdu, China), were maintained in animal holding laboratory under controlled condition with temperature of 25±10°C, humidity of 40±10% and had free access to standard mouse diet and water.

Sera

For the production of mice antisera against live or inactivated A. pleuropneumoniae serotype 5, the 50% of lethal dose (LD50) were determined firstly. Six-week-old Kun-Ming mice were randomly divided into five groups, each of them consisting of eight mice. and injected intraperitoneally (i.p.) with 200 µl TSB containing A. pleuropneumoniae (1.0×10⁸ to 1.0 ×10⁹ CFU/mouse). The number of surviving mice was recorded 5 days after
infection. LD50 was calculated by the method of Reed and Muench and determined to be 6.3 × 10^6 CFU. Then Six-week-old Kun-Ming mice were randomly divided into two groups, each of which consisting of twelve mice. Mice in group A were injected i.p. with 200 μl of A. pleuropneumoniae preparation containing 6×10^5 CFU per mouse on days 0, 14 and 28 respectively; while mice in group B were inoculated i.p. with the same amount of inactivated A. pleuropneumoniae as group A per mouse on days 0, 14 and 28 respectively. Mice were bled from eyes 10 days following the final boost and sera were collected. For the production of mice antisera against live A. pleuropneumoniae serotype 1 or 7, twelve mice were inoculated with live A. pleuropneumoniae preparation containing 1/10 LD50 per mouse following the method above.

Construction of Recombinant Prokaryotic Expression Vector

* A. pleuropneumoniae* present in 1 ml of an overnight culture were collected by centrifugation and DNA was extracted using a sodium dodecyl sulfate (SDS) - proteinase K and phenol/chloroform protocol as described in previous research [6]. Then, bioinformatics method was used to analyze the apxIVA gene sequence of A. pleuropneumoniae serotype 5 completed by our lab (Genbank NO: GQ332268), a pair of specific primers was used to amplify the 1152 bp fragment of the 5’-terminal of apxIVA A gene. The two primers were 5’CCA GAATTC AGC TAC GAA ACA AGG- 3’ with EcoRI site (underlined) comprising position 665–672 of apxIVA coding sequence and 5’CTC CTGCAG TTA ATT CAG ATC GGA GGC A -3’ with XholI site (underlined) comprising position 1791–1806 of apxIVA coding sequence. The oligonucleotides were synthesized by Takara biotechnology (Dalian, China) co., ltd. Amplification of the apxIVA gene was performed with polymerase chain reaction (PCR) and the PCR products were cloned into the pMD18-T plasmid (Takara) to yield recombinant plasmid. Confirmation of clones containing recombinant plasmid was achieved by PCR and restriction enzyme (RE) digestion, and the correct clones were sequenced by Takara biotechnology (Dalian). The recombinant plasmid were then digested with EcoRI and XholI and cloned into pET-32a(+) (Novagen, Madison, USA) digested with EcoRI and XholI to generate recombinant prokaryotic expression plasmid.

Expression and Purification of apxIVA

The BL21 strains containing the recombinant prokaryotic plasmid were grown overnight with shaking at 37°C. The overnight cultures were diluted in the proportion of 1:100 and incubated at 37°C with vigorous shaking for 3-4 hours (h). The production of poly-His-tailed ApxIV a proteins were induced by addition of 1 mM IPTG and incubation for a further 3-4 h. The cells were harvested and analyzed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). Optimization of expression conditions at different IPTG concentration ranging from 0, 0.2, 0.4, 0.6, 0.8, 1.0 to 1.2 mM and different induction time ranging from 0, 2, 3, 4, 5 to 6 h were also conducted.

For purification of recombin apxIVA, the cells were harvested from liquid cultures by centrifugation and suspended in phosphate-buffered saline (PBS) (pH 7.2) containing lysozyme (1mg/ml) by 1/10 (v/v), after ice bathing for 30 min, the suspension were sonicated and centrifuged at 12000g for 10 min. The pellets were homogenated and washed with washing buffer [50 mM Tris-Hcl (PH8.0), 1 mM EDTA, 0.2% TritonX-100, 2 M urea] for three times at 10min/times and centrifuged at 12000g for 10 min. The pellets were dissolved by denaturation buffer [50 mM Tris-Hcl (PH8.0), 2 mM 2-mercaptoethanol, 8 M urea] and supernatant were collected after centrifugation. The supernatant was treated with renaturation buffer [50 mM Tris-Hcl (PH8.0), 0.1 mM oxidized glutathione, 1 mM reduced glutathione, 0.5 M urea] for overnight at 4°C and filtrated through 0.45 μM filtration membrane. Then the solution was purified on a column packed with Ni-NTA His-Bind superflow according to the manufacture’s instruction (Merck, Darmstadt, Germany). The eluate from each imidazole concentration were collected and analyzed by SDS-PAGE.

Western Blot Analysis

Purified proteins were separated by SDS-PAGE, then the proteins were transferred to nitrocellulose membrane with 0.45μm pore size (Millipore Corp., USA) at 15V for 1.5 h with a Bio-Rad Transblot Cell as described in the manufacturer’s specifications. The membrane was then blocked for 90 min with milk buffer (20mM Tris–HCl pH 8.0, 150mM NaCl, 0.05% Tween 20, 5% skimmed dry milk) at 37°C, and then washed with Tris-buffered saline with Tween 20 (TBST) buffer (20mM Tris-HCl PH 8.0, 150mM NaCl, 0.05 Tween-20) for three times and incubated with mouse antiserum diluted 1:100 in 0.5% bovine serum albumin (BSA) /PBS for 60 min at 37°C. The membrane was washed with TBST, then incubated with hors eradish peroxidase (HRP) -labeled sheep-anti-mouse IgG (zhongshan Goldenbridge Biotechnology Co., LTD, Beijing, China) for 60 min at 37°C. Target proteins were visualized using 3,3'- diaminobenzidine (Tiangen, Beijing, China).

Development of an apxIVA-ELISA

A 96-well microtiter plate (Costar, USA) was coated with 100 μl of purified apxIVA and incubated at 4°C overnight. The plate was then blocked for 90 min with milk buffer (5% skimmed dry milk in PBS) at 37°C and washed twice with PBS. Subsequently, 100 μl of mice sera were added and incubated at 37°C for 60 min. The samples were washed, and then incubated for 45 min with 100 μl of anti-mouse IgG diluted 1:10000 in 0.1% BSA/PBS at 37°C, washed again, and detected with 100 μl of 3,3',5,5'- tetramethyl benzidine (TMB) for 20 min at room temperature (RT). The reaction was then stopped by the addition of 35 μl of 2 M H2SO4. The optical density (OD) value was read at 450 nm, using a Bio-Rad model 860 plate reader (Bio-Rad, CA, USA).

Checker board method was used to determine the optimal antigen coating concentration, serum dilutions and HRP labeled rabbit anti-mouse IgG dilutions. For determine the cut-off value, twenty-four sera samples from mice uninfected with *A. pleuropneumoniae* were used as negative sera, the cut-off value was calculated using the formula: mean of the negative serum values plus three standard deviations (SDs). To determine the specificity of this method, rabbit antisera of HCV, PRRSV and PM (obtained from Dr. Qigui Yan, Sichuan Agricultural University, China) were detected.
according to the method above, except that the second antibody was anti-rabbit IgG (Zhongshan, Beijing, China) diluted 1:10000 in 0.1% BSA/PBS. Finally, the method was used to detect the sera of inactivated *A. pleuropneumoniae* immunized and live *A. pleuropneumoniae* infected mice.

**RESULTS**

**Construction of an apxIVA Prokaryotic Expression Vector**

The *apxIVA* gene was isolated from *A. pleuropneumoniae* ATCC 263 by PCR as described in Materials and Methods. The oligonucleotides used in the PCR reaction were designed to allow the 1152 bp long fragment in 5′ terminal part of *apxIVA* gene to be cloned into pET-32a(+) in frame with the poly-His purification signal. PCR and RE analysis showed that prokaryotic expression plasmid PET32a-apxIVA was successfully constructed. (Fig. 1).

**Prokaryotic Expression and Purification of apxIVA**

Prokaryotic expression vector PET32a-apxIVA was induced by addition of IPTG to produce recombinant *apxIVA* toxin at different time ranged from 0, 2, 3, 4, 5 to 6 h and at different IPTG concentration ranged from 0, 0.2, 0.4, 0.6, 0.8, 1.0 to 1.2 mmol L⁻¹, the results showed that a protein about 62Kd was expressed (Fig. 2) since 1h and stabilized on 4 h post culturation at a most appropriate IPTG concentration of 0.6 mM (data not show ). The recombinant *apxIVA* was expressed mainly in the form of inclusion body and Ni-NTA superfowl was used to purify this his-tagged *apxIVA*. For the purified protein, a single objective band was detected by SDS-PAGE (Fig. 2).

**Western Blot Assay**

To confirm the reactivity of his-tagged *apxIVA* with antibodies from *A. pleuropneumoniae* infection, the purified protein was exposed to a Western blot assay by using serum from *A. pleuropneumoniae*. The result showed that the fusion protein can react positively with serum from live *A. pleuropneumoniae* infection (Fig. 2), while negatively with serum from inactivated *A. pleuropneumoniae* immunization.(data not show).

**Indirect ELISA Based on Recombinant ApxIVA**

The results of checker board experiment showed that a concentration of 1.0 μg/ml of apxIVA-His peptide and 1:40 serum produced the best resolution of the positive reference serum from the negative reference serum. Then 24 negative mice sera was selected randomly to react with *apxIVA*-His peptide, the average χ was 0.143, standard deviation(SD) was 0.028. The positive critical value was χ+3SD=0.226. Means that while the OD value of sample ≥0.226, the result was positive; while the OD value of sample <0.226, the result was negative. For the results of specificity test, positive sera of HCV, PRRSV and Pm were negative in the *apxIVA*-ELISA. indicating the specificity of this *apxIVA*-ELISA (Table 1).

**Table 1. The Results of Specificity Test of apxIVA-ELISA**

<table>
<thead>
<tr>
<th>OD₄₅₀ of apxIVA-ELISA</th>
<th>PRRSV</th>
<th>HCV</th>
<th>Pm</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Positive Sera</strong></td>
<td>Sera Number</td>
<td>0.151</td>
<td>0.083</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0.095</td>
<td>0.162</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0.133</td>
<td>0.098</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.125</td>
<td>0.087</td>
</tr>
</tbody>
</table>

PRRSV: Porcine reproductive and respiratory syndrome virus; HCV: Hog cholera virus; Pm: Pasteurella multocida.

For the results of discrimination of infection and inactivated vaccine immunization, Sera of four mice immunized with inactivated *A. pleuropneumoniae*, four mice
infected with live serotype 1, four mice infected with live serotype 5 and three mice infected with live serotype 7 A. pleuropneumoniae gave positive results (Table 2).

**DISCUSSION**

Porcine pleuropneumonia was an respiratory disease hazardous to China pig industry. The virulence of the 15 serotypes of *A. pleuropneumoniae* is mainly determined by three major RTX toxins apx I, apx II and apx III [7], which are secreted by the different serotypes in various combinations, but apxIV can be produced by all 15 serotypes only during infection, but not under *in vitro* conditions [4].

The diagnosis of *A. pleuropneumoniae* infections is important for the identification of farms that are latently infected with *A. pleuropneumoniae*, detection of different *A. pleuropneumoniae* serotypes, inspection of the animals on their importation into a farm, or eradication of the infection. It is done routinely by serological examination of pig herds or by bacterial agent cultivation from nasal swabs or lungs with *pleuropneumonia* lesions or by PCR. Serological examination were usually carried out by detecting antibodies provoked by Apx toxin or outer membrane protein (OMP) or lipopolysaccharides (LPS) etc. OMP are common to all *A. pleuropneumoniae* serotypes [8], while LPS are serotype-specific and used to serologically distinguish infections with different serotypes of *A. pleuropneumoniae* [9], LPS was also found to give serological cross-reaction with non-pathogenic *Actinobacillus* species [10]. PCR-based methods were suitable for the rapid detection of *A. pleuropneumoniae* [11].

Vaccination is potentially an effective tool for the control of Porcine pleuropneumonia, although natural or experimental infection generally elicits at least partially protection against reinfection with another serotype [12]. Killed whole cell bacterins were commercially used in China now, and a major drawback of vaccination, is that traditional serological tests including hemmaglutination test [13, 14] and ELISA [15], cannot distinguish immunization of killed whole cell bacteria from field exposition. As apxIVA can be expressed by all 15 serotypes only *in vivo* conditions, therefore, antibodies against apxIVA should be the target of serological differentiation between immunization with killed whole cell bacteria vaccine and infection with live *A. pleuropneumoniae*.

Although an indirect ELISA based on recombinant apxIVA of *A. pleuropneumoniae* serotype 1 have been reported previously [16], but indirect ELISA based on recombinant apxIVA of serotype 5 have not been reported now. The serotype 5 is highly pathogenic, and the sequence of apxIVA of serotype 5 was different from those of serotype 1 and 3 in length and homology [17]. So we develop an indirect ELISA based on recombinant apxIVA of serotype 5 to see whether it can also detect antibodies against *A. pleuropneumoniae* infection. After the bioinformatic analysis of apxIVA gene sequence of serotype 5 strain CVCC 263, it was found that the 5'-terminal of apxIVA gene was highly antigenic, and subsequently a 1152 bp fragment in this region was amplified and expressed to establish an indirect ELISA to detect the antibody against the apxIVA of all serotypes. The results showed that the sera from inactivated vaccine immunized mice were negative, while the sera from live *A. pleuropneumoniae* infected mice were positive. So this indirect ELISA could detect *A. pleuropneumoniae* infection in a mouse model. The validation of this apxIVA-ELISA to detect *A. pleuropneumoniae* infection in pigs will be further undertaken.

**ACKNOWLEDGEMENTS**

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**ABBREVIATIONS**

- **BSA** = Bovine serum albumin
- **ELISA** = Enzyme-linked immunosorbent assay
- **HRP** = Horseradish peroxidase
- **i.p.** = Intraperitoneally
- **IPTG** = Isopropy 1-β-D-thiogalactoside
- **LPS** = Lipopolysaccharides
- **NAD** = Nicotinamide adenine dinucleotide
- **OD** = Optical density
- **OMP** = Outer membrane protein
- **PBS** = Phosphate-buffered saline
- **PCR** = Polymerase Chain Reaction

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**Table 2. Detection of Antibodies Against *A. pleuropneumoniae* Infection and Inactivated *A. pleuropneumoniae* Immunization**

<table>
<thead>
<tr>
<th>OD₄₅₀ of apxIVA-ELISA</th>
<th>Animal Numbers</th>
<th>Inactivated App Serotype 5 Vaccinated Mice</th>
<th>Live App Serotype 1 Infected Mice</th>
<th>Live App Serotype 5 Infected Mice</th>
<th>Live App Serotype 7 Infected Mice</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>0.171</td>
<td>1.145</td>
<td>1.297</td>
<td>1.175</td>
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<tr>
<td></td>
<td>2</td>
<td>0.133</td>
<td>0.899</td>
<td>1.183</td>
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<tr>
<td></td>
<td>3</td>
<td>0.201</td>
<td>1.013</td>
<td>1.215</td>
<td>0.933</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.145</td>
<td>1.235</td>
<td>1.033</td>
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</table>

*App: A. pleuropneumoniae.*
Indirect ELISA for Detecting A. pleuropneumoniae Infection

RE = Restriction enzyme
RT = Room temperature
SDS-PAGE = Sodium dodecyl sulfate polyacrylamide gel electrophoresis
TBST = Tris-Buffered Saline with Tween 20
TSB = Tryptic Soy Broth

REFERENCES


