Original Article

Cloning, Expression, Purification, and Oligomeric Characterization of the AopB-C-terminus Domain in T3SS Major Translocator Protein of Aeromonas hydrophila

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Abstract: Type three secretion system (T3SS) is found exclusively in gram-negative pathogens such as Yersinia spp., Escherichia coli, Salmonella spp., Shigella spp., Pseudomonas spp., Vibrio parahaemolyticus, and Aeromonas hydrophila. The translocon pore of T3SS comprises major and minor translocator proteins that assemble to provide passage of effectors through the host cell membrane. Major translocator protein AopB from Aeromonas hydrophila plays an important role in translocon pore formation. Despite tremendous efforts, structural information regarding the C-terminus domain of major translocator AopB remains elusive. In this study, the DNA fragment encoding for the C-terminus domain of the AopB from Aeromonas hydrophila AH-1 was cloned into pET-M expression vector and expressed in Escherichia coli BL21 (DE3) host cells. The recombinant AopB-C-terminus domain was successfully purified using immobilized nickel affinity chromatography as a soluble form. Crosslinking analysis among AopB-C-terminus molecules in solution showed that this domain existed as a mixture of tetramer, trimer, dimer, and monomer forms. The three-dimensional structure model of AopB-C-terminus oligomerization was built by SWISS-MODEL and PyMol. The oligomeric model of AopB-C-terminus can be used for structural studies of the AopB-C-terminus domain which can contribute to the elucidation of the structure of the type III secretion system.

Keywords: Aeromonas hydrophila, affinity chromatography, AopB-C-terminus domain, gene expression, oligomerization.

1. Introduction

Aeromonas hydrophila is a gram-negative bacterium that causes disease in many organisms such as fish, shrimp, and humans leading to mass death in fish in aquaculture farms. Similar to other gram-negative bacteria, Aeromonas hydrophila uses a type three secretion system (T3SS) to deliver toxins into the host [1, 2]. T3SS consists of complex macromolecular machinery, translocator and effector proteins, chaperone, and other accessory proteins [3]. In T3SSs, the translocator proteins
play an important role in the pore formation on the host cell membrane and help bacterial toxins to enter the host [1, 4]. Many translocators of bacterial families have been identified such as AopB/AopD (Aeromonas sp.), IpaB/IpaC (Shigella spp.), YopB/YopD (Yersinia spp.), PopB/PopD (Pseudomonas spp.), and SipB/SipD (Salmonella spp.) [2].

To date, structure information of major translocators are limited to a short N-terminal peptide (9-13 residues) [5-8] of translocator PopB with chaperon PcrH in 2012 [9], the structure of IpaB/IpaC [10], or IpaD [11]. Especially, in 2015 Nguyen et al. showed the structure of the N-terminal parts of the translocator AopB (1-264) with its chaperon AcrH [12]. Currently, there is no structural information available to show how the C-terminus domain of major translocators interacts with other translocators. Here, we cloned the AopB-C-terminus domain (residue from 265 to 347) into a pET-M expression vector. The recombinant protein was expressed in E. coli BL21 (DE3) cell and purified by nickel affinity chromatography. We showed that AopB used the C-terminus domain to interact and form oligomeric stages which are necessary for functional T3SS formation.

2. Materials and Methods

The pET-DUET-1 AcrH-AopB vector, the pET-M expression vector, the E. coli DH5α, and the E. coli BL21 (DE3) were from the Molecular Cell Biology lab, in the Center of Life Science, Faculty of Biology, and VNU University of Science. Chemical using in this experiment were bought from international companies as Bio-rad, Sigma, Merck, Thermo Fisher Scientific (United States), Bio Basic (Canada), or Serva (Germany).

2.1. Cloning of AopB-C-terminus into pET-M Expression Vector

A gene encoding for the AopB-C-terminus from amino acid 265 to amino acid 347 was amplified by polymerase chain reaction using forward primer AopB265-347-F (5'-GCCggatccGTAGTTGATATTGGTACC CGGA-3') containing a sequence for BamHI cleavage and reverse primer AopB265-347-R (5'-CGgaattcTTAAATGGCTGTCGTGC CGTC-3') containing a sequence recognized by EcoRI (Table 1). The DNA template was the pET-DUET-1 AcrH-AopB vector containing the full-length sequence of AopB. The PCR reaction used Phusion High-Fidelity DNA Polymerase (Thermo Scientific) with composition and the temperature cycle according to the guideline of the manufacturer.

The PCR product of the AopB-C-terminus gene and pET-M were digested by FastDigest EcoRI and BamHI (Thermo Scientific). The DNA was inserted into the vector by T4 DNA ligase (Thermo Scientific). The ligated product was transformed into E. coli DH5α competent cells then spread on LB agar medium added with ampicillin (100 µg/ml). The plasmid containing the AopB-C-terminus gene was screened by PCR screening method. Subsequently, plasmids are extracted from E. coli cells using GeneJET Plasmid Miniprep Kit (Thermo Scientific). The foreign gene in the selected plasmid was sequenced at 1st Base (Singapore) using the Sanger method.

2.2. Expression of AopB-C-terminus Protein

The pET-M-AopB-C-terminus vector was transformed into E. coli BL21 (DE3). A single colony of transformant was cultured in 5 ml LB containing 100 µg/ml of ampicillin (LBA medium) overnight, at 37 ℃, 150 rpm. The overnight culture was transferred into 1L of LBA and shook at 37 ℃ until OD600 reached 0.6. The transcription of the foreign genes was initiated by adding 0.3 mM IPTG (isopropyl β-D-1-thiogalactopyranoside) into medium and continued culture at 25 ℃, 150 rpm for 16 hours. The cells were collected by centrifugation for 10 min at 4000 rpm and stored at -30 ℃ until purification.
2.3. Purification of AopB-C-terminus

The cell pellet was resuspended in 25 ml of lysis buffer (Tris-HCl 30 mM, pH 8.0; NaCl 300 mM; imidazole 5 mM) and were lysed by sonication on ice with 40% amplitude and for 6 rounds (3s on, 3 s off) of 5 min each. The lysate was centrifuged at 13000 rpm, 4 °C for 30 min. The collected supernatant was loaded into Econo-column (Biorad) containing 3 ml Ni-NTA bead pre-equilibrated with 20 ml of lysis buffer. The nonspecific-binding proteins were removed from the column by washing step with washing buffer (Tris-HCl 30 mM, pH 8.0; NaCl 300 mM; 30 mM imidazole). The AopB-C-terminus tagged hexahistidine were eluted with 20 ml of elution buffer (Tris-HCl 30 mM, pH 8.0; NaCl 300 mM; imidazole 500 mM). The eluted proteins were dialyzed overnight against phosphate-buffered saline buffer (140 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, and 1.8 mM KH₂PO₄, pH 7.4) and stored at -80 °C.

2.4. Chemical Crosslinking of AopB-C-terminus Domain

The crosslinking reaction was conducted with 100 µL of AopB-C-terminus (1 mg/mL protein concentration) in PBS buffer and 0.2% glutaraldehyde. The reaction was carried out at 4 °C for 30 minutes. Total 20 µL of the reaction mixture was sampled at 10 min, 20 min, and 30 min periods. The samples were added with SDS loading dye buffer and denatured immediately at 95 °C for 5 min. Then all samples were run on a 15% polyacrylamide gel SDS-PAGE.

2.5. Modelling Oligomeric State of AopB-C-terminus

The tertiary structure of AopB-C-terminus domain was modeled based on the amino acid sequence of this domain using SWISS-MODEL [13]. The oligomer structure was predicted based on the homologous structures published on PDB and analyzed by PyMOL software.

3. Results and Discussion

3.1. Cloning of AopB-C-terminus Gene into the pET-M Expression Vector

The AopB-C-terminus gene, which is containing BamHI and EcoRI restriction enzyme sites at 5' and 3' end respectively, was amplified by Phusion DNA polymerase (Thermo Scientific). The result was shown in Figure 1(A). The AopB-C-terminus gene of 265 bp was specifically amplified by PCR (Lane 2, Figure 1A). The PCR product was digested by the BamHI and EcoRI and inserted into pET-M expression vector to generate a recombinant plasmid pET-M-AopB-C-terminus (Figure 1B).

The AopB-C-terminus gene in the recombinant vector was confirmed by Sanger sequencing.

![Figure 1. Analysis of PCR product of AopB-C-terminus gene (A) and extracted recombinant plasmid pET-M-AopB-C-terminus (B) by electrophoresis on agarose gel 0.8%. 1A: DNA marker 100 bp (iNtRON); 2A: The PCR product. 1B: DNA marker 1 kb (iNtRON); 2B: Plasmid pET-M; 3B: pET-M-AopB-C-terminus.](image-url)
3.2. Expression of AopB-C-terminus Domain in *E. coli* BL21 (DE3)

The result of SDS-PAGE electrophoresis in Figure 2 showed that AopB-C-terminus domain with the length of 10.97 kDa was expressed in *E. coli* BL21 (DE3) cells when the cells were cultured in the medium containing 0.3 mM IPTG at 25 °C (Lane 3, Figure 2) and this band was not observed in the control sample (Lane 2, Figure 2).

3.3. Purification of His-tag AopB-C-terminus

The protein AopB-C-terminus was purified using a Ni-NTA affinity column. The results (Figure 3) showed that AopB-C-terminus domain was successfully purified with a thick specific band of about 10.97 kDa in lane 7. The AopB-C-terminus was dialyzed in PBS buffer overnight at 4 °C. Dialyzed protein was aliquoted into 1ml tubes and stored at -80 °C (Figure 3).

3.4. Determination of Oligomeric states of the AopB-C-terminus

In order to determine the oligomeric states of *AopB-C-terminus*, we performed chemical crosslinking between the AopB-C-terminus molecules in solution. Only the molecule closely interacts with each other would cross-link and result in higher molecular weight on denatured protein electrophoresis (SDS-PAGE) analysis. In Figure 4, lane 3, there were 4 bands with corresponding sizes for a dimer (21.94 kDa), trimer (32.91 kDa), and tetramer (43.88 kDa) form of AopB-C-terminus domain (Figure 4).
Figure 4. The result of chemical crosslinking reaction. Lane 1: Protein ladder (Lonza); Lane 2: Monomeric AopB-C-terminus without chemical crosslink; Lane 3: crosslinked AopB-C-terminus domain.

3.5. Oligomeric Model of AopB-C-terminus

We modeled the three-dimensional structure of AopB-C-terminus using SWISS-MODEL software. The monomer and oligomer structure of the AopB-C-terminus domain was built based on a template with SWISS-MODEL Template Library ID (SMTL ID) of 1o5h.1.A. The result in Figure 5A showed the monomer form of AopB-C-terminus domain, mainly alpha helices.

The oligomer state of this protein was predicted based on the monomeric structure as well as the chemical crosslink analysis as shown in Figure 5B, which consists of parallel alpha-helices. Based on the published structure of the AopB N-terminal domain with the AcrH chaperon, we predict that the oligomerization of the C-terminus helps align the N-terminus region together to form the transmembrane region.

4. Conclusion

In this research, we cloned the AopB-C-terminus domain gene into the pET-M vector and induced the expression in E. coli BL21 (DE3). The AopB-C-terminus domain was purified by His-tag affinity chromatography. The highest oligomeric state of AopB-C-terminus domain could be at least tetramer form based on the result of the chemical crosslinking reaction. The structure models of AopB-C-terminus were predicted by SWISS-MODEL software. In the future, proteins with high purity can be used for further studies to understand their structure and function.

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