Fusion expression of *Helicobacter pylori* neutrophil-activating protein in *E. coli*

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**INTRODUCTION**

*H. pylori* is now recognized as one of the most widespread human pathogens. Stomach mucosa colonized by *H. pylori* is commonly accompanied with inflammatory infiltrates, consisting mainly of neutrophils and monocytes[3,4]. There is a good correlation between the degree of mucosal damage and neutrophil infiltration[5,6]. Several studies have proved evidence for the presence of protein components in *H. pylori* water extracts capable of attracting and activating neutrophil adhesion to endothelial cells[7,8]. It is termed *H. pylori* neutrophil-activating protein (HP-NAP). HP-NAP is localized in bacterial cytosol and released upon autolysis. It can bind to the external surface of the outer membrane, similar to what was found for urease[7,8]. In such a location, HP-NAP can mediate HP-NAP binding to the cell surface, thus, HP-NAP is supposed to be a major virulence factor and vaccine candidate[9]. To study the pathogenesis of *H. pylori* infection and screen potential antigens for *H. pylori* vaccine development, neutrophil-activating protein gene of *H. pylori* (HP-NAP) was amplified by PCR from *H. pylori* MEL-HP27 strain, which was isolated from a clinical patient in Henan Province of China (preserved in Molecular Epidemiology Laboratory, Zhengzhou University). The napA sequence data have been published on GenBank (Accession No: AY366361) by the authors. Software analysis indicated that HP-napA gene was a highly conserved prokaryotic gene that could encode a 15-kD polypeptide. The aim of this study was to produce recombinant protein of *H. pylori* in *E. coli* TB1 and to screen vaccine candidates. So the recombinant plasmid pMAL-c2x-napA was constructed, human recombinant mannose-binding protein (rMBP)-NAP expressed in *E. coli* TB1 was used as an antigen to immunize rabbits. Experiment with animal, *H. pylori* human anti-serum and software were used to evaluate the immunogenicity and immunoreactivity of rMBP-NAP.

**MATERIALS AND METHODS**

**Materials**

*E. coli* TB1, pMAL-c2x, amyllose resin, and goat anti-rabbit IgG-HRP, goat anti-human IgG-HRP were purchased from NEB Company. pNEB-napA was constructed by the authors, 1 kb DNA ladder was from Sangon. *Sal I*, EcolI, T4 DNA ligase, IPTG, Xgal were from Takara. Freund’s adjuvant was from Sigma; NC Western blot membrane was from Beijing Yili Company. DAB was from Roche Company, rabbits were from Laboratory Animal Center of Henan Province, *H. pylori* human anti-sera were preserved in Molecular Epidemiology Department of Zhengzhou University.

**Construction of recombinant plasmid**

Common DNA manipulation was done according to the methods described by Sambrook *et al*.[10]. In brief, 0.5 µg pNEB-napA and 0.5 µg pMAL-c2x were digested respectively in 20 µL 1×H buffer with 1 µL SalI and 1 µL EcoR. Then, 2 µL pMAL-c2x digest and 1 µL napA fragment were mixed after purification, and 14 µL deionized H₂O (Milli-Q) was added. DNA mixture was heated at 45 °C for 5 min and then cooled on ice, and 2 µL

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The quantity was tested by Bradford assay.

Evaluation by gel image analysis system (SynGene, USA) and purified by amylose affinity chromatography. The purity was for 30 min at 4 °C overnight. Sonicated in ice-water bath, centrifuged at 9 000 r/min overnight, and incubated at 16 °C for 3 h.

Cells of 2×10^7/mL containing amp was inoculated with 1 mL of the cultured TB1 cells. Cells of 2×10^7/mL rich glucose broth were cultured in 5 mL broth containing amp at 37 °C overnight, sonicated in ice-water bath, centrifuged at 9 000 g for 30 min at 4 °C. Soluble rMBP-NAP in the supernatant was purified by amylase affinity chromatography. The purity was evaluated by gel image analysis system (SynGene, USA) and the quantity was tested by Bradford assay.

Expression and purification
The positive clones of TB1(pMAL-c2x-napA) on the master plate were cultured in 5 mL broth containing amp at 37 °C, 200 r/min overnight. One hundred mL rich glucose broth containing amp was inoculated with 1 mL of the cultured TB1 cells. Cells of 2×10^8/mL(Ao≈0.5) were cultured at 37 °C, IPTG was added to a final concentration of 0.3 mmol/L, and induced for 3 h. The expression of induced fusion protein was identified on SDS-PAGE. The cells were harvested and frozen at -20 °C overnight, sonicated in ice-water bath, centrifuged at 9 000 g for 30 min at 4 °C. Soluble rMBP-NAP in the supernatant was purified by amylase affinity chromatography. The purity was evaluated by gel image analysis system (SynGene, USA) and the quantity was tested by Bradford assay.

Immunization of rabbits and Western blotting
Rabbits (3-5 kg) were immunized with purified rMBP-NAP four times. The primary immunization by hypodermic injection consisted of 100 μg rMBP-NAP and 0.5 mL complete Freund’s adjuvant. Three enhancement immunizations were performed 4 wk after the first injection, each dose consisting of 50 μg rMBP-NAP and 0.25 mL incomplete Freund’s adjuvant at weekly intervals. The sera were collected 3 wk after the last immunization. The negative sera were collected before the vaccination. Purified fusion protein was transferred to nitrocellulose (NC) membranes by semi-dry transfer cell (TRANS-BLOT.SD, BIORAD) at 15 V for 20 min after SDS-PAGE. Fusion protein imregnated NC strips were incubated with anti-rMBP-NAP rabbit serum (1:500) and anti-H pylori human serum (1:50) as the first antibody respectively, incubated with goat anti-rabbit IgG-HRP and goat anti-human IgG-HRP after washing. The secondary antibody was detected by reaction with DAB.

Antigenic features of rMBP-NAP evaluated by Omiga2.0
HP-napA and MBP-NAP gene sequence were inputted into Omega2.0 program. The sequences were translated into proteins and their antigenic features were calculated.

RESULTS
Construction of recombinant plasmid
Recombinant plasmid restriction enzyme digestion is shown in Figure 1. The target fragment of 435 bp napA was inserted into the pMAL-c2x between EcoRI and sal I.

rMBP-NAP expression and purification
SDS-PAGE analysis of expressed and purified rMBP-NAP is shown in Figure 2. According to the gel image analysis system, the molecular weight of rMBP-NAP was about 57 kD, and the soluble fusion protein accounted for 37.55% of the total protein in the sonicated supernatant. The yield of fusion protein after affinity purification was 100 mg/L culture by Bradford assay.

Immunoreactivity and immunogenicity of rMBP-NAP
Western blotting showed that the purified rMBP-NAP could be recognized with the sera from rabbits immunized with rMBP-NAP (Figure 3) and anti-H pylori sera from clinical patients infected with H pylori. In contrast, the result of the control sera from rabbits before vaccination and healthy people was negative.

Antigenicity of rMBP-NAP analyzed by Omega 2.0.

Figure 1 Restriction analysis of pMAL-c2x-napA. Lane 1: 1 kb DNA ladder; Lane 2: pMAL-c2x-napA digested by EcoRI; Lane 3: pMAL-c2x-napA digested by EcoRI and SalI; Lane 4: pMAL-c2x digested by EcoRI; Lane 5 HP-napA PCR fragment.

Figure 2 SDS-PAGE analysis of rMBP-NAP. Lane 1: empty plasmid induced by IPTG; Lane 2: TB1(pMAL-c2x-napA) induced by IPTG; Lane 3: lower molecular weight protein marker; Lane 4: affinity chromatography purified rMBP-NAP.

Figure 3 Western blotting of purified rMBP-NAP.

Figure 4 Antigenicity of rMBP-NAP analysed by Omega 2.0.
**DISCUSSION**

*H. pylori* infection is the major cause of chronic active gastritis and most peptic ulcer diseases\(^1\). It is also closely related to gastric cancers such as adenocarcinoma, mucosa-associated lymphoid tissue (MALT) lymphoma. Approximately half the population in the world is infected with *H. pylori*. Vaccination might be the most effective and economic way to control this organism. Vaccine development is a process to identify the unique structures capable of generating immunological protection when formulated as a vaccine. So antigen and adjuvant selection is critical in developing *H. pylori* vaccines. But so far the achievement in this area is still unsatisfied. Several studies have focused on identifying major virulence factors of *H. pylori* such as CagA\(^2\), urease and heat shock proteins. The toxicity of these natural proteins has limited their application and heat shock proteins might cause host immuno-cross reactions. A vaccine candidate must meet the following criteria: antigenic shock proteins might cause host immuno-cross reactions. A of these natural proteins has limited their application and heat shock proteins might cause host immuno-cross reactions.

A vaccine candidate must meet the following criteria: antigenic conservation among clinical isolates, elicitation of functional antibodies, protection of animal models, and ultimate safety and efficacy for human use\(^3\). So one of the tactical designs of subunit vaccines is to modify the structure of the virulence factors. Dundon et al reported that the majority of *H. pylori*-infected patients were found to produce antibodies specific for HP-NAP, making it a strong vaccine candidate. We amplified HP-napA by PCR from *H. pylori* MEL-HP27 strain and sequence analysis, and found that HP-napA gene was highly conserved among prokaryotic organisms. The results of this study showed that HP-napA fusionally expressed with maltose binding protein of *E. coli* did not degrade the immunoreactivity, but the immunogenicity of the fusion protein increased when the molecule was larger than natural HP-NAP. The feature of HP-NAP could be modified added with MBP. Moreover, maltose-binding protein provided a one-step affinity purification of the fusion protein using maltose resin, which could be cleaved from rMBP-NAP by factor Xa after purification. In conclusion, DNA recombinant technique and bioinformatics can provide an alternative approach to produce novel antigens for vaccine development. The features of rMBP-NAP as a vaccine candidate and the role of HP-NAP in *H. pylori*-related pathogenesis should be further evaluated in animal models.

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