A preliminary Study: Expression of Rhopty Protein 1 (ROPl) *Toxoplasma gondii* in Prokaryote System

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**Background:** *Toxoplasma gondii* is an obligatory intracellular protozoan parasite, which infects human beings. Since the current antigens used for diagnosis or vaccination are contaminated with non parasitic material in which the parasite is grown, it is tried to produce recombinant antigens to design vaccines against toxoplasmosis, or make diagnostic kits. Choosing the type of antigen to produce recombinant vaccine or diagnostic kits is considerably important. The rhoptry protein 1 is one of the excretory-secretary antigens of Toxoplasma which seems to be an appropriate candidate in production of recombinant vaccines and diagnostic kits.

**Objectives:** The current study aimed to produce rhoptry protein 1 from Iranian strains of *Toxoplasma* for further investigations.

**Materials and Methods:** Genomic DNA was isolated from tachyzoite of parasite by phenol chloroform method and gene fragment was amplified by polymerase chain reaction. The polymerase chain reaction products were ligated into restriction enzymes sites of pTZ57R/T cloning vector. The product was sub-cloned into a prokaryotic expression plasmid (pET32a). The recombinant expression vector containing rhoptry protein 1 sequence was transformed into *Escherichia coli* BL21 pLysS and was induced by isopropyl β-D-1-thiogalactopyranoside. The sodium dodecyl sulfate polyacrylamide gel electrophoresis and western blotting methods were used to confirm the production of protein.

**Results:** The result showed that the desired molecular weight protein is produced. Recombinant protein was confirmed by western blot using Ni-NTA-conjugate.

**Conclusions:** The result of this study showed that recombinant ROPl *Toxoplasma* was produced successfully. In the present study, unlike other studies, the goal was to express full length of ROPl. Also this protein was produced alone not fused with other proteins.

**Keywords:** Toxoplasma gondii; ROPl Protein; Purification

1. Background

*Toxoplasma gondii* is an obligatory intracellular protozoan parasite, which infects human beings when eating undercooked meat or contaminated vegetable. *Toxoplasma* may cause different clinical forms such as asymptomatic or severe symptoms like abortion, congenital defects and death (1). Since the current antigens that are used for diagnosis or vaccination are contaminated with non parasitic material in which the parasite is grown, a lot of efforts are made to produce recombinant antigens to design vaccine against toxoplasmosis or make diagnostic kits (2). Choosing the type of antigen to produce recombinant vaccine or diagnostic kits is considerably important because it should be immunogen, trigger of cellular imm

2. Implication for health policy/practice/research/medical education

The results of this study allow us to design future studies to evaluate the diagnostic value of this protein in human toxoplasmosis. We can also evaluate the ability of this protein to induce host immune response.
tein could be used as a specific marker for differentiation of acute and chronic toxoplasmosis (3,17-19).

2. Objectives

The current study aimed to produce rhoptry protein 1 from Iranian strains of *Toxoplasma* for further investigations.

3. Materials and Methods

3.1. Parasite

The RH strain of *T. gondii*, was provided by Department of Parasitology, Faculty of Health, Tehran Medical Sciences University. Tachyzoites of parasite were harvested from peritoneal fluid of Balb/c mice. These mice had been infected 3 - 4 days earlier.

3.2. Cloning of ROP1 Gene in Bacterial Expression Vector

Standard methods were used for DNA extraction (20). Authors designed a pair of primers according to the ROP1 gene sequence (Accession number M71274) with EcoR1 and SalI restriction sites at 5’ end of forward and reverse primer respectively (Forward: 5’-GGAATTCACCATGGAGCAAAGGCTG-3’ and Reverse: 5’-GCGTCGACTTATTGCGATCATCATCC-3’). The ROP1 gene of *Toxoplasma* was amplified by PCR. The purified PCR products were ligated into pTZ57R/T plasmid vector (20). The gene fragment was sub-cloned in a pET32a expression plasmid (prepared from Microbiology Department, School of Medical Science, Arak University).

Initially, the gene fragment was separated from previous cloned plasmid by restriction cleavage and then purified using gel-extraction kit (Fermentas Inc. Canada). In addition, pET32 was cultivated in an antibiotic-containing media. After plasmid extraction, its restriction cleavage sites for EcoR1 and SalI were enzymatically cleaved and purified. Then, resulting plasmids were used for ligation and transformation of gene fragment. The recombinant pET32a expression vector containing ROP1 gene (pET32a-ROP1) was confirmed by PCR, restriction analysis and sequencing.

3.3. Protein Expression and Purification

Escherichia coli BL21 (DE3) pLysS was transformed with pET32a-ROP1 and grown in LB broth supplemented with ampicillin (100 µg/mL) and chloramphenicol (34 µg/mL) (Invitrogen, USA) and shaken at 200 rpm at 37°C. Also *E. coli* BL21 (DE3) pLysS was not transformed grown as control. The plasmid promoter was induced by 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) for 6 hours. Sampling was done before and after induction in 2-hour intervals. The bacterial cells were harvested by centrifugation at 8000 rpm for 5 minutes. The expressed protein was analyzed by SDS-PAGE and western blotting (12). The expressed protein was purified using Ni-NTA column according to manufacture instructions (Qia-gen, Germany).

3.4. SDS-PAGE and Western Blot Analysis

Recombinant protein was assayed by 12% gradient SDS-PAGE and western blot. SDS-PAGE was performed according to the method of Laemmli (21). Western blot was performed by Ni-NTA conjugate. At first, SDS-PAGE gel was blotted on to Polyvinylidene fluoride (PVDF Membrane, Roche Diagnostic, Germany) membrane using transfer buffer (12). Membrane was incubated 1 hour in 1% skimmed milk and washed 3 times with washing buffer (Tris-buffer-saline and tween 20 or TBS-tween). Membrane was incubated 1 hour in TBS-tween buffer containing a 1/1000 dilution of Ni-NTA conjugate. The blots were then washed three times with TBST and reactions were developed by diaminobenzidine (DAB) Solution (Roche Diagnostic Germany).

4. Results

The genomic DNA of *T. gondii* was prepared and used as template for amplification of the gene encoded ROP1. The amplified fragment had the expected size of 1183 bp comparing to Molecular weight marker (Figure 1).

![Figure 1. ROP1 Gene Amplification by PCR](image)

Lane 1: Molecular weight marker (1 kb ladder); Lanes 2 and 3: amplified ROP1 with 1183 bp band.
The recombinant plasmid was analyzed by restriction enzyme and the resulting fragments were compared with molecular weight marker (Figure 2).

**Figure 2.** Cleavage of ROP1 Fragments From Prokaryotic pET32a Expression Plasmid

Lane 1: 1Kbp molecular weight marker; Lane 2: recombinant pET32a-ROP1 before cleavage; Lane 3: recombinant pET32a-ROP1 after cleavage showing a 1183 bp fragment; Lane 4: cleavage of non-recombinant pET32a and absence of 1183 ROP1 fragment

The recombinant plasmid (pET32a-ROP1) was sequenced. The sequencing result was confirmed by comparing with the databases and using basic local alignment search tool (BLAST) Software (Figure 3).

**Figure 3.** Alignment of Amino Acids Sequence of *T. gondii* ROP1 Fragment Compared with Amino Acid Sequence of ROP1 Accession Number AY661790.1 and M71274.1 (Respectively)

Expression of pET32a-ROP1 in *E. coli* BL21 (DE3) pLysS was induced and the expressed protein was purified by Ni-NTA column. SDS-PAGE and western blot analyses showed the expected molecular mass near 60 kDa recombinant protein (Figures 4 and 5).

**Figure 4.** Expression of Recombinant ROP1 Protein in SDS-PAGE gel

L. protein size marker SM0671; Lane 1: pET32a-ROP1 before induction without expected protein (~ 60 kDa); Lane 2, 3: pET32a-ROP1 after induction that expressed expected protein (~ 60 kDa).

**Figure 5.** Western Blot Analysis of the Recombinant ROP1 Protein Using Ni-NTA Conjugate

Protein size marker SM0671 (Fermentas Inc.). Lane 1: expressed ROP1 4 hours after induction with expressed protein (~ 60 kDa); Lane 2: expressed ROP1 6 hours after induction (~ 60 kDa); lane 3: recombinant plasmid before induction.

5. Discussion

*T. gondii* is an obligatory intracellular parasite which has a complicated life cycle and attacks almost all nucleated cells. The high incidence and severe or lethal damages of toxoplasmosis clearly indicate the need for the develop-
ment of a more effective vaccine (22). Moreover, in recent years, researchers are trying to improve the serological diagnosis of toxoplasmosis using recombinant proteins. Therefore, it is critical to generate specific antigens. Recombination is an easy method to access this specific protein (23).

Available commercial tests to detect anti-Toxoplasma antibodies were usually prepared based on whole antigens derived from tachyzoites of parasite. This stage of parasite must be harvested from infected mice or cell cultures because antigens prepared by the traditional method are not composed of defined parasite antigens. They are inevitably contaminated with at least some host tissue, thus could result in false-positive reactions and reducing the clinical value of serological test (24). Many researchers have emphasized the detection of specific antibodies against excreted-secreted antigens (ESA) of Toxoplasma in diagnosis of acute toxoplasmosis is worth (24). T. gondii ESA constitutes 90% of the circulating antigens in infected humans, thus this parasite is one of the first targets of the host immune response (24).

ROP1 gene is one of the major antigens of T. gondii with unusual gene structure. The major part of this gene has similar sequence in genomic DNA, cDNA and mRNA so it is possible to directly isolate gene fragment from DNA (11). This protein is presented in tachyzoites, bradizoites and sporozoites forms of parasite and plays a role in invasion to host cells. The ROP1 is expressed in many phases of parasitic life cycle, it confers a suitable target for production of clinical diagnostic kits and recombinant vaccines (2). Gue et al. cloned a 760bp ROP1 gene fragment into pBV220 expression plasmid and assessed the immune responses of Balb/c mice vaccinated by this plasmid (13). In another study carried out by this group, encoding plasmids for a 760bp fragment of ROP1 gene and a 600bp fragment of IFN-α gene were generated and immunologically assessed in Balb/c mice (15).

Chen et al. cloned a 750bp fragment of T. gondii ROP1 gene into pUC18 plasmid and then transferred it to pCDNA3 eukaryotic expression plasmid (11). Another group directed by Chen constructed a fusion-based recombinant plasmid consisting of two fragments of ROP1 (760bp) and SAG1 genes (pSAG1-ROP1). Recombinant plasmids were injected to Balb/c mice via liposome-entrapped particles and immune responses were analyzed (16). Eslamirad et al. cloned a 760bp fragment of T. gondii ROP1 gene into eukaryotic expression plasmid. In the study, the used vector was pTZ57R, which was different from vectors used in other studies. Also the used gene fragment could only encode part of ROP1 protein that was expressed at very low levels in eukaryotic CHO cells and was hardly traceable in vivo. However, the results reflected relatively satisfactory immunogenicity of plasmids in animal models (20).

Aubert et al. produced a number of T. gondii antigens such as ROP1 in fusion with cks protein of E. coli and used these recombinant antigens to identify anti-toxoplasmosis specific IgM and IgG antibodies in human serum (17). Meek et al. cloned fragments of ROP1 and SAG1 genes into pRPl261 expression plasmid and used the resulting expressed proteins to serological diagnosis of toxoplasmosis in patients suffering from acute and chronic toxoplasmosis (18).

In brief, the current study expressed recombinant ROP1 in E. coli expression system as full length. This protein was produced alone. It was prepared for use in the diagnosis of toxoplasmosis by ELISA system and for evaluation as a recombinant vaccine. In the current study, unlike other studies, the goal was to express full length of ROP1. Also this protein was produced alone not fused with other proteins. This will provide the chance to design future studies to evaluate the diagnostic value of this protein in human toxoplasmosis. One can also evaluate the ability of this protein to induce host immune response.

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Authors’ Contribution
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References
7. Garcia JL, Gennari SM, Navarro JT, Machado RZ, Sinhorini IL. Toxoplasma gondii: isolation of tachyzoites rhoptries and incorpora-