Evaluating the serological applications of Toxoplasma gondii rhoptry protein 1 (ROP1) antigen

Fatemeh Keshavarzi1*, Parviz Ashtari2, Pirooz Ebrahimi3

1. Department of Biology, Sanandaj Branch, Islamic Azad University, Sanandaj, Iran
2. Radiation Application Research School, Nuclear Science and Technology Research Institute, Tehran, Iran
3. Sapienza University of Rome, Yas Women Hospital, Tehran

*Corresponding author: Tel: +98 9183704918 Fax: +98 8333288661
Address: Department of Biology, Sanandaj Branch, Islamic Azad University, Sanandaj, Iran
E-mail: gol.keshavarzi@gmail.com
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Abstract

Introduction: Toxoplasma gondii (T. gondii) is distributed worldwide and infects most species. The serious incidence and severe or fatal injury caused by T. gondii infection clearly indicates the necessity for the event of a vaccine. The current study goals were to evaluate serological applications of Toxoplasma gondii rhoptry protein 1 (ROP1) antigen.

Materials and methods: We created a polymer vaccine by using the eukaryotic plasmid, pROP1. Purification by one-step metal affinity chromatography allowed recovery of milligram amounts of purified recombinant proteins per liter of culture. The quality of this matter for diagnosing of human infections was provided and tested on 77 serum samples which were obtained during routine diagnostic tests. A panel of 20 serum samples from patients with acute toxoplasmosis was compared to a panel of 35 serum samples from individuals with chronic toxoplasmosis.

Results: Results of the study indicated that antibodies detected from patients with acute and chronic infections were 96% and 17%, respectively, by using of pROP1 recombinant antigen.

Conclusion: According to the present study an immunoglobulin G antibody against ROP1 antigen is made throughout the acute stage of toxoplasmosis infection, but not in the chronic phase of toxoplasmosis.

Keywords: Toxoplasma gondii, ROP1, Recombinant antigen, Acute toxoplasmosis

Introduction

Toxoplasma gondii (T. gondii) is a parasite belonging to the Apicomplexa (1, 2). This parasite is distributed across worldwide and infects most species of animals and humans. Toxoplasmosis is immune competent in adults and is usually asymptomatic.

Toxoplasmosis is a main cause of fatality in congenitally infected patients and immune compromised patients (3-5). The live vaccine is dangerous because of the reversing to pathogenic phenotype and low shelf life. Additionally, DNA vaccines can exert potent, long-lasting humeral and cell-mediated immunity (6, 7). T. gondii infections usually promote to activate CD4+ and CD8+ T cells, simultaneously (7, 8). Like others unicellular organisms T. gondii is composed of various antigens. Somatic and excreted/secreted antigens are the most important Toxoplasma antigens that the secreted ones can activate potent immune response (9, 10). In the present study, we constructed a plasmid expressing the ROP1 antigen (pROP1), then examined their expression in eukaryotic cells and evaluated the usefulness of the T. gondii recombinant r-ROP1 antigen for diagnostic purposes. The antigenicity of recombinant antigen against human serum samples was
evaluated by Western blotting and ELISA analysis.

Material and methods

Animals and T. gondii strains: Some old female BALB/c mice were purchased and kept under sterile in animal laboratory of the Sanandaj Azad University. A highly virulent strain of T. gondii preserve (RH strain) in BALB/c mice was used for production of tachyzoites.

DNA extraction and polymerase chain reaction: About 5 \times 10^7 tachyzoites (100μl) were concentrated by centrifugation and washed with phosphate buffer saline and then lysed in 900μl lysis buffer and then treated with 10 μl proteinase K (100μg/ml) at 55°C for 2 h. The proteins were removed by adding an equal volume of phenol/chloroform to lysate. The lysate tube was centrifuged (13000 RPM /15 min) and an equivalent volume of chloroform was added to the supernatant. DNA pellet was obtained after adding equal content of 100% ethanol and 1/10 volume of 3M sodium acetate to this supernatant and centrifugation at 13000 RPM for 10 min and then washing with 70% ethanol. The DNA pellet dissolved in sterile distilled water and stored at -20°C until use. Extracted DNA products were examined on 0.8% agarose gel. The sequence provided by digestion of BglII and HindIII was designed for cloning. From the GenBank database (http://www.ncbi.com) we took the sequences of the gene encoding ROP1 antigen of T. gondii (accession no. M71274).

The forward (F) and reverse (R) sequences of primers were appropriately determined as follow:

F: GTCGACATCTAGGTGTCAATCTCG
R: CCAAGCTTTTTGCGATCCATCTCG

The DNA segment of rop1 (corresponding to nucleotides 252 to 1188) encoding ROP1 was obtained by PCR using above primer. DNA extracted from tachyzoites was used as a template to amplify the ROP1 gene by PCR performed in 25μl of solution containing 3μl of template DNA, 0.5μl TaqDNA polymerase, 2.5μl 10X PCR buffer, 0.75μl MγCl2, 15.75μl distilled water and 1μl each of primers under the following conditions: After an early 5 min denaturation at 94°C, each cycle consisted of 60s at 94°C, 30s at 62°C and 60sat 72°C at the end of the 32 cycles of amplification and 30 min at 72°C was final extension. The PCR products checked out by electrophoresis on a 1% agarose gel and taken a photo. The molecular weight markers used were the 1kb DNA ladders (Fermentas).

Cloning of ROP1: The pUET1 cloning vector was used and ligation was done according to our previous study (11). The ligation product used to transform and was accomplished Escherichia coli BL21 plus. Competent cell was prepared and the ligation output was a shiftin competent cell. According to protocol (12) retrieved in Luria-Bertani (LB) broth and LB broth medium free antibiotic by incubating at 37 °C for 2 hours and then were plated onto LB agar plates containing IPTG 200 mg /ml, ampicillin 100 mg/ml and X-Gal 20 mg/ml, and plates were incubated at 37°C for overnight to screening white and blue colonies. The selected colony of both color passages in LB broth or LB agar incubated at 37 °C for an overnight. Based on the protocol, DNA plasmid was extracted and the cloned ROP1 was verified by PCR, nucleotide sequencing and restriction digestion.

Sub-cloning of the pT-ROP1 inpcDNA3eukaryote expression vector: The digested pT-ROP1by EcoR1 and HindIII inserted into the digested pcDNA3 vector and the inserted product was transformed into CHO cells (competent cells). The plasmid was extracted and identify by electrophoresis, PCR amplification and restriction digestion (11).

SDS-PAGE and Western blot analysis: After 48h, pellets of transfected cells were suspended in 150ml SDS-PAGE sample buffer and then sonicated and heated (100°C/ 5 min). Lysates were separated on a 12% polyacrylamide gel, transferred to
polyvinylidene difluoridecoats, and then blocked with Tris-buffered saline (20 mM Tris, 137 mM NaCl, pH 7.6) containing 0.1% Tween 20 and 5% skim milk. Membranes were washed with buffer and then for 2 hours at room temperature mixed with primary antibody of mouse anti-FLAG. Interested antibody was washed out with TBST. Peroxidase activity was a study using enhanced chemiluminescence (ECL) Western blotting detection system.

**ELISA assay:** MaxiSorp multi well plates were coated with 0.1 ml of r-ROP1 protein at a final concentration of 1 g/ml for antigen in a coating buffer (0.05 M carbonate buffer, pH 9.6). Control plates were incubated at 4°C with the extract recombinant proteins as mentioned above. After overnight, controls were washed (PBS–0.1% Triton X-100) and blocked (37°C for 1 hour) by using blocking solution (1% bovine serum albumin–1% Triton X-100–PBS). Afterward, they were covered (1 hour at 37°C) with human serum specimens diluted 1:100 in the blocking solution. Plates were washed with PBS–0.1% Triton X-100 and added to each well from the conjugates of anti-human IgM peroxidase-labeled (1 mg/ml), diluted 1: 4,000 in blocking solutions. Next, the plates were incubated 35 min at 37°C and then washed. Finally, by incubating the plates with o-phenylene diamine dihydrochloride chromogenic substrate was revealed enzymatic activity. After 40 min at 37°C in darkness, by adding 0.1 ml of 1 M sulfuric acid was stopped the color development reaction and the color intensity was measured using a Microtiter plate reader at 490 nm. Each serum sample was examined and calculated of the optical density (OD) reading.

**Serum samples:** A total of 77 healthy human serum samples was analyzed and based on serological profiles divided into three classes. The class I consisted of 20 human serum specimens from patients in the acute phase of toxoplasmosis. The presences of specific IgM, antibodies were measured by IgM-ELISA Vistas'. All serum samples had positive IgG antibodies. Class II consisted of 35 human serum samples from patients with chronic toxoplasmosis. By an absence of specific IgM antibodies and an antibody avidity test evaluated all those serum samples had positive IgG antibodies high avidity. Class III (the control group) included 22 human serum samples from serious negative individuals. For more, statistical analysis of the ELISA results was evaluated with the Microsoft Excel schedule for assessment of the 2 test.

**Results**

The sequence analysis verified that PCR-created change and the resulting vector included of 1686 base pairs (accordance with Gene Bank, Z36906.1) from ROP1 gene. ROP1 indicated in vitro by transfected CHO cells (competent cells) ROP1 was synthesized during an organism system (Figures 1, 2 and 3).

![Figure1](https://via.placeholder.com/150)

**Figure1.** Figure shows the result of electrophoresis pUET1 and pT-ROP1 plasmids. Lane 1: Plasmid extracted from pT-ROP1 and Lane 2: Plasmid extracted from pUET1.

The CHO cells (transfected and non-transfected control cells) were produced for 48 h following the transaction and list in sample buffer. The ROP1 antigen described in detail as soluble protein with calculated molecular weight 39 kDa. The immune action of the purified antigen r-ROP1 was tested by Western blot analysis with 4 human serum samples selected from each of the three serum sample groups that we were assayed (Figure 4). IgG antibodies present in serum group I samples in the serious stage of toxoplasmosis efficiently identified the r-ROP1 antigen. In contrast,
IgGs from patients group II in the continuous stage of infection reacted weakly with Rop1. Moreover, immunoreactions of r-ROP1 antigen with serum samples from healthy patients (group III) was not studied.

![Figure 2](image1.png)

**Figure 2.** Digestion of extracted pT-ROP1 after RANs formation. Lane1: ladder 1kb, Lanes 2: (ROP1 757bp, pUET12876 bp).

![Figure 3](image2.png)

**Figure 3.** Detection of pcDNA3-ROP1 by enzyme Digestion. Lane1: ladder 1kb, 2: pcROP1, 3: pc-ROP1 after digestion EcoRI enzyme, 4: pc-ROP1 after digestion with HindIII and 5: pcROP1 after digestion with each other enzymes (ROP1~ 760 bp).
Figure 4. Expression of Recombinant ROP1 Protein in SDS-PAGE gel. L: protein size marker; Lane 1: p-ROP1 before induction without expected protein, Lane 2 and 3: p-ROP1 after induction that expressed expected protein.

Discussion

Chen et al. cloned a 750bp fragment of *T. gondii* ROP1 sequence into PUC18 cellular inclusion so transferred it to pcDNA3 organism expression cellular inclusion (12). Another group constructed a fusion-based recombinant plasmid (pSAG1-ROP1). By liposome-entrapped particles Recombinant plasmids were injected into Balb/c mice and immune responses were analyzed (13). The recombinant antigenic proteins are encouraging tools that can be used in diagnostic tests for the between serious and constant infections. Aubert et al. produced ROP1 in fusion with cks protein of *E.coli* and to identify specific IgM and IgG antibodies of toxoplasmosis in human serum used these recombinant antigens (14). The fragments of ROP1 and SAG1 genes into pRP261 expression vector cloned and used the resulting explicit proteins to serological assessment of toxoplasmosis in patients suffering from acute and chronic toxoplasmosis by Meek et al. (15).

In this research reported the bacterial production of *T. gondii* soluble r-ROP1 antigen protein by the use of the effective expression system developed in our laboratory (11). This expression system and the method of disinfection by a one-step metal attraction chromatography led to the production of 14 mg of r-ROP1 per liter of culture (11). In this work, we appraised the diagnostic avail and worthiness of *T.gondii* recombinant antigenic protein r-ROP1 and then tested r-ROP1 restarts by the use of two separate classes of human serum parable show serious and constant infections. Both groups were analyzed by ELISA techniques and Western blot. IgG antibodies in serum samples from patients in the serious stage of toxoplasmosis (group I; serum samples with positive IgM and IgG at low enthusiasm) respond with the recombinant antigens more potent than those in serum specimens from patients in the constant state of toxoplasmosis (group II; serum samples with positive IgG at high acting and negative IgM. Thus, our results with r-ROP1 antigen imply that these recombinants can be used as serological indicator of recently obtained infection. The r-ROP1 antigen was shown before to be sensitive in its capability to discover IgG antibodies in serum samples from patients with toxoplasmosis (12-15). The usefulness of this antigen for diagnosis of human
infections was provided and tested on 77 serum samples which are obtained during routine diagnostic tests. A panel of 20 serum samples from patients with acute toxoplasmosis was compared to a panel of 35 serum samples from individuals with chronic toxoplasmosis. The results indicated that ROP1 recombinant antigen detected antibodies further in individuals with acute infections (96%) than in individuals with chronic infections (17%). The results of this study show that special IgG anti-ROP1 antibodies are present in serum samples from patients at the serious stage of T. gondii infection. Successful use of recombinant antigenic proteins for the finding of anti-T. gondii- special antibodies has been reported by a number of authors (15-16).

Conclusion

These data imply that ROP1 antigen is specifically encouraging candidates for efficient vaccine against toxoplasmosis. Additionally, results suggest that an immunoglobulin G antibody against ROP1 antigen is created during the acute stage of infections but not generally in the persistent phase of the sickness.

Acknowledgment

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References