PCR detection of *Toxoplasma gondii* *B1* gene in women suffering from abortion

Aseel S. Mahmood  
Sabeeha A. Al-Sarray  
Abdul-Kareem A. Al-Kazaz

Biotechnology Department / College of Science / University of Baghdad

E-mail: aseel78nm@yahoo.com

**Abstract**

Background: Primary infection of maternal with toxoplasmosis during gestation and this infection transmission to the fetus continue to be the cause complex disease in offspring.

Objective: This study was conducted to test the utility of nested Polymerase Chain Reaction (nPCR) assay to detect recent infections with *Toxoplasma* in abortive women.

Material and methods: *Toxoplasma gondii* DNA was detected by using *B1* gene as a target for amplification which was highly specific for *T. gondii* and is well conserved among all of the tested strains. Blood from 60 abortive women and 25 apparently healthy pregnant women with no history of abortion (as control group) were taken in this current study.

Results: The results revealed that nPCR was positive in 48(80%) subjects and negative in 12(20%), Chi-square- $\chi^2$ for patients and control was (13.82, 15.75) respectively.

Conclusion: It can be concluded that nPCR assay in blood has advantage in detection of recent and active toxoplasmosis.

Keywords: *Toxoplasma gondii*, *B1* gene, nested PCR.

**Introduction**

The *toxoplasma gondii*, is an obligated intracellular parasite. It can infect humans and several vertebrates leading to toxoplasmosis. This mostly benign affection cause severe life-threatening disease, immunocompromised particularly in patients and congenitally affected children (1,2).

Infection of healthy adults *T. gondii* is asymptomatic in most cases. Function of Immune system shall prevent parasite replication and destroy any bradyzoites that are dormant tissue released cysts. However, tachyzoites can cross the placenta if a woman is infected during pregnancy this lead fetus infect (2). The infection course and symptoms are depend on many factors inclusive inoculation factors, particular organism virulence, gestational age at time of infection, genetic factors, and immune status of the mother and fetus (3).

Serological diagnosis act as most widely used approach to know the infection stage, whether current, recent or past (4). In spite of its high sensitivity these tests can supply vague results. For a definitive diagnosis direct detection of the parasite is necessary. Several PCR based assay have been developed for the detection of *Toxoplasma* DNA with *B1* repetitive sequence which has been proven to be more sensitive when compared to other target e.g. *P30* gene and rDNA (5). This is probably because *B1* is repeated DNA sequence with a higher copy number than the single – copy *P30* gene (6). The *B1* gene is a 35-fold repetitive gene sequence with unknown function, PCR amplifying this target has shown high specificity for DNA detection (7).

**Materials and Methods**

This study was carried out on 60 abortive Iraqi women and 25 control group of apparently healthy pregnant woman in ranged age between 18-34 years. Samples were obtained from subjects in two hospitals included, AL– ALwia and Eben AL Balady.

**Blood samples and isolation of DNA**

Five ml was obtained of venous blood sample from each patient and control subjects in EDTA tube and stored at -20°C until DNA extraction. DNA extraction from blood than stored at -20°C until used by AccuPrep® Genomic DNA Extraction Kit Bioneer.
Nested PCR assay

Nested PCR was complete all DNA samples to amplify a fragment from the B1 gene, which is present in 35 copies and is conserved in the T. gondii genome, as described by (8). The primers used in the first round of the PCR (inner primers) were (5'-GGAACCTGCATCGTCTCGTCT-3') and (5'-TTCTTAAACGCGTTCTGTCGTC-3'), which correspond to nucleotides 694–714 and 887–868, respectively at 193 bp. Then primers used in the second round (outer primers) were (5'-TGCATAGGTTGCAGTCACTG-3') and (5'-GGCGACCAATGTCGAATAGACCC-3'), which correspond to nucleotides 757–776 and 853–831, respectively at 96 bp. The 5 µL of DNA template were added to a final volume of 20 µL of PCR and mixture consisting of master mix, Then 1 µL forward and reverse primers in concentration (10 pmol), and 13 µL of nuclease-free water added. Then amplification in the thermal cycler PCR shows in Table (1).

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<tr>
<th>Steps</th>
<th>Temperature</th>
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<tr>
<td>Initial Denaturation</td>
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<td>Denaturation</td>
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The PCR product were used for electrophoresis is 10 µL on 2% gm agarose gel and detect 193 fragment of B1 gene. five µL from PCR product was used as template was used to the second run PCR in a total volume of 20 µL of PCR mixture consisting of master mix, 1 µL of Forword and Reverse in the concentration 10 pmol from each primer, completed by 11 µL of nuclease-free water. Conditions of PCR second run were temperature 94°C at 3 minutes, 40 cycles at 94°C at 30 seconds, 50°C at 45 seconds and 72°C at 45 seconds and final extension at 72°C for 10 minutes. A 10 µL from PCR product was used into electrophoresis in 2.5% agarose gel stained by ethidium bromide to detect 96 fragment of B1 gene.

Result and Discussion

A parasite DNA disclosure by PCR has extremely amended diagnosis, especially in prenatal diagnosis to congenital disease (9).

A results are depend on amplification two DNA fragments (193 bp and 96 bp) of B1 gene by Analysis of nPCR in infected women blood sample. The consequence have showed that clear band of DNA fragment of PCR product these signal a presence of B1 gene in 49(81%) with highly significant difference and that clarified the existence of infection, while 11(19%) showed absence of amplified DNA fragment indicated that not found the B1 gene no infection Figure (1) and Figure (2) as well as Table (2).

![Figure (1): First run of nested PCR (amplification of 193bp) fragment of B1 gene of T. gondii. M marker 100bp, 1-5 positive sample and other negative samples](image-url)
Toxoplasma gondii is compose parasite protozoan intracellular that worldwide spreading. Humans infection is commonly wanting incidental, but it ability apparent itself in a several form in cases of congenital toxoplasmosis and in immune compromised individuals(10). In this study results of statical analysis the chi-square $\chi^2$ of patients have shown $(13.82)$ and control $(15.75)$ are significant in nPCR mean to high infection in Iraqi women.

Some studied presented that a single T. gondii parasite could be detected by PCR (8) agreement with this study. A top exposition to increase rate of positive PCR results that the amplification of $B1$ gene may be due to samples containing DNA parasite but it no viable pathogens (11).

A highly sensitive and specific of PCR is in order to a single one can be detected in a clinical sample (5). Numerous studies were reported that parasitaemia could detect by PCR in a few weeks prior to the appearance of any clinical symptoms (12, 13). These advantages have proportional speed, A detect potential very low numbers of pathogens or may more precisely, particular sequences of nucleic acid of pathogens and the ability to strictly distinguish at the species or sub-species level (14).

Disclosure T. gondii by DNA with minimizes PCR the problems notable when employ serology or cultured based assays. It economize labor and time display the high advantages of sensitivity and specificity. Used PCR to demonstrate the existence of Toxoplasma in diverse clinical samples such brain, amniotic fluid and cerebrospinal fluid (CSF), lymph nodes, whole blood and aqueous humor (15). PCR is consider ultimate importance in diagnosing infection of T.gondii in cases of immunosuppressive therapy or in patients with AIDS (14).
The abortive women with positive was found Toxoplasma DNA in 48 (80.00%), this could be coincide to a very neoteric infection when serological leading to scanty production of immunoglobulin not detected by serology, or other explanation that those patients are not able to produce specific antibodies, representing a state of immunodeficiency. Molecular tests that could detect the presence of circulating parasites would be of extreme application in this scenario. A positive serological result is direct detection of T. gondii in blood or elucidate the presence of parasite leading to the primary diagnosis.

(16)

References