Direct genotypic characterization of Toxoplasma gondii strains associated with congenital toxoplasmosis in Tunisia (North Africa).
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Direct Genotypic Characterization of *Toxoplasma gondii* Strains Associated with Congenital Toxoplasmosis in Tunisia (North Africa)

Sonia Boughattas, Rym Ben-Abdallah, Emna Siala, Olfa Souissi, Karim Aoun, and Aïda Bouratbine*

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**Abstract.** Here, we determined the *Toxoplasma gondii* genotype in amniotic fluid, placenta, and cerebrospinal fluid samples from 14 congenital toxoplasmosis cases in Tunisia, North Africa. Direct genotypic characterization of *T. gondii* strains was performed by polymerase chain reaction (PCR) amplification of six genetic markers (3' *SAG2*, 5' *SAG2*, *SAG3*, *BTUB*, *GRA6*, and *APICO*) and thereafter, was analyzed by restriction fragment-length polymorphism (RFLP). Samples were sequenced to resolve strain type whenever there were unclear enzyme digestion results. Multilocus analysis revealed that only one specimen harbored the type I allele in all studied loci, whereas the 13 others gave mixed genotype results with different alleles at different markers. Seven specimens produced RFLP profile of the recombinant strains I/III, and three produced a profile of I/II recombinant strains. The last three specimens produced complex digestion patterns. In these cases, sequence analysis revealed double peaks at known polymorphic sites, indicating the presence of multiple alleles.

**INTRODUCTION**

Toxoplasmosis is a common parasitic disease caused by the protozoan parasite *Toxoplasma gondii*. Its prevalence in humans varies from region to region depending on ecological and cultural factors. In Tunisia, North Africa, an average seroprevalence rate of 58% with a progressive rise from 24% at 10 years of age to 70% at 30 years of age was reported. This epidemiological feature suggests that even if infection is a frequent event in childhood, many women at childbearing age stay susceptible to toxoplasmosis.

The possible relationship between congenital toxoplasmosis and *T. gondii* genotype has been approached in several studies. In France, where a systematic diagnosis of congenital toxoplasmosis cases in Tunisia, North Africa. Direct genotypic characterization of *T. gondii* strains was performed by polymerase chain reaction (PCR) amplification of six genetic markers (3' *SAG2*, 5' *SAG2*, *SAG3*, *BTUB*, *GRA6*, and *APICO*) and thereafter, was analyzed by restriction fragment-length polymorphism (RFLP). Samples were sequenced to resolve strain type whenever there were unclear enzyme digestion results. Multilocus analysis revealed that only one specimen harbored the type I allele in all studied loci, whereas the 13 others gave mixed genotype results with different alleles at different markers. Seven specimens produced RFLP profile of the recombinant strains I/III, and three produced a profile of I/II recombinant strains. The last three specimens produced complex digestion patterns. In these cases, sequence analysis revealed double peaks at known polymorphic sites, indicating the presence of multiple alleles.

**MATERIAL AND METHODS**

**Clinical samples.** Fourteen cases of congenital toxoplasmosis diagnosed at the Department of Parasitology of Pasteur Institute of Tunis between 2004 and 2008 were included in the study. Congenital infections were identified through maternal pre-natal screening on the basis of pre-natal diagnosis (real-time quantitative PCR; genes B1 and AF487550) and/or neonatal screening (PCR on placenta; serology in newborns) as previously described. Parasite DNA used for genotyping was obtained from amniotic fluid (AF) in 11 cases, placentas (PL) in 2 cases, and cerebrospinal fluid (CSF) in 1 case. No clinical data were available for newborns in 11 cases. Two newborns had no toxoplastic clinical signs, and one newborn developed chorioretinitis (Table 1).

**Experimental samples.** Reference strains used for each of the clonal lineages (provided by Pr. Marie Laure Dardé, Biologic Resources Center (BRC) Toxoplasma; http://www .tioxbrcc.com/) were: RH (type I), Prugniaud (type II), and NED (type III). Parasites were grown and maintained in Swiss mice by intraperitoneal inoculations. Tachyzoites were harvested from the peritoneal cavities (RH strain) or brains (NED and Prugniaud strains) in phosphate-buffered saline (PBS); then, they were pooled, centrifuged, and washed two times at 2,000 g for 10 minutes. The parasite pellets were processed for DNA extraction using Qiamp DNA Mini Kit (Qiagen, GmbH, Germany).

**PCR–RFLP analysis.** Strain typing was performed by PCR amplification of six genetic markers (3' *SAG2*, 5' *SAG2*, *SAG3*, *BTUB*, *GRA6*, and *APICO*) and thereafter, was analyzed by RFLP. First, the lineage type was determined using two nPCRs...
amplifying separately the 5′ and 3′ ends of the SAG2 gene as previously described (Table 2). 13 Thereafter, a multiplex nPCR with a set of three-way markers (SAG3, BTUB, GRA6, and APICO) was used (Table 2). 14,18 The initial round of amplification with external primers for the four different markers was carried out in 25 μL of mixture containing 1× PCR buffer, 3 mM MgCl2, 400 μM each of the deoxyribonucleoside triphosphates (dNTPs), 1 μg bovine serum albumin (BSA), 0.2 μM each of the forward and reverse primers, 1.5 units of Hot Start DNA polymerase (Qiagen), and 4 μL of DNA sample. Amplification was conducted at 94°C for 15 minutes followed by 40 cycles of 94°C for 45 seconds, 55°C for 1 minute, and 72°C for 1 minute. The last extension step was at 60°C for 10 minutes. PCR products were diluted and used for second-round amplification of each marker separately in a 25-μL volume mixture containing 1× PCR buffer, 2.5 mM MgCl2, 200 μM each of the dNTPs, 0.2 μM of each primer, and 0.75 units of Hot Start DNA polymerase (Qiagen), and 4 μL of DNA sample. Amplification protocol was 94°C for 15 minutes followed by 40 cycles of 94°C for 45 seconds, annealing temperature for 45 seconds, and 72°C for 1 minute. The overextension step was at 72°C for 5 minutes. The annealing temperature was 50°C for Apico, 55°C for GRA6, and 62°C for BTUB and SAG3. PCR products were examined by electrophoresis on 2% agarose gel stained with ethidium bromide and visualized under ultraviolet (UV) light. The amplified fragments of PCR products were used to screen for possible new polymorphisms. nPCR products and sequencing was also performed on selected samples to determine strain type whenever there were unclear enzyme digestion results, and sequencing was also performed on selected samples to distinguish each type strain. The digested PCR products were resolved by 3% agarose gel stained with ethidium bromide.

To avoid possible contamination, several measures were taken—separate spaces were used to set up PCRs, filter tips, and UV radiations; also, tests were run with few samples in the same run. Different negative controls (no DNA, uninfected sample, and extracted no DNA) were also used. Controls were carried out from RH, Prugniaud, and NED strains. The polymorphism within each locus was analyzed by RFLP patterns used to distinguish each type strain. The digested PCR products were resolved by 3% agarose gel stained with ethidium bromide.

**DNA sequencing.** Samples were sequenced to resolve strain type whenever there were unclear enzyme digestion results, and sequencing was also performed on selected samples to screen for possible new polymorphisms. nPCR products from T. gondii strains as well as clinical samples were used for DNA sequencing. The fragments were sequenced with both internal primers SAG3 and GRA6 to obtain forward and reverse sequences. Sequencing was done three times with different PCR products from the same isolate to avoid any contamination. The kit ABI Prism BigDye Terminator Cycle Sequencing Reaction (Applied Biosystems, Branchburg, NJ) was used, and electrophoreses were run on a polyacrylamide gel POP7 in a four-capillary Applied Biosystems 3130 Genetic Analyzer. Nucleotide sequences were aligned for comparison using “Clustal W” from Bio-Edit Sequence Alignment Editor. The sequences were deposited in GenBank, and the accession numbers of the nucleotide sequences are cited later.

**Table 1.** Origin of samples and information about biological diagnosis of congenital toxoplasmosis.

<table>
<thead>
<tr>
<th>Code</th>
<th>Sample</th>
<th>Maternal infection</th>
<th>Pre-natal diagnosis</th>
<th>Neonatal screening</th>
<th>Post-natal follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ultrasound</td>
<td>AF puncture (PCR)</td>
<td>Placenta (PCR)</td>
</tr>
<tr>
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<td>AF06/06</td>
<td>AF</td>
<td>1st T</td>
<td>N</td>
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</tr>
<tr>
<td>2</td>
<td>AF44/05</td>
<td>AF</td>
<td>1st T</td>
<td>N</td>
<td>+</td>
</tr>
<tr>
<td>3</td>
<td>CSF01/08</td>
<td>CSF</td>
<td>3rd T</td>
<td>ND</td>
<td>ND + +</td>
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<tr>
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<td>AF08/08</td>
<td>AF</td>
<td>2nd T</td>
<td>N</td>
<td>+</td>
</tr>
<tr>
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<td>AF26/04</td>
<td>AF</td>
<td>1st T</td>
<td>N</td>
<td>+</td>
</tr>
<tr>
<td>6</td>
<td>AF08/06</td>
<td>AF</td>
<td>1st T</td>
<td>N</td>
<td>+</td>
</tr>
<tr>
<td>7</td>
<td>AF33/05</td>
<td>AF</td>
<td>3rd T</td>
<td>N</td>
<td>+</td>
</tr>
<tr>
<td>8</td>
<td>AF07/06</td>
<td>AF</td>
<td>1st T</td>
<td>N</td>
<td>+</td>
</tr>
<tr>
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<td>PL</td>
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<td>AF16/07</td>
<td>AF</td>
<td>1st T</td>
<td>N</td>
<td>+</td>
</tr>
</tbody>
</table>

AF = amniotic fluid; CSF = cerebrospinal fluid; PL = placenta; T = trimester; N = normal; ND = not done.

**Table 2.** Summary of markers used for genotyping T. gondii.

<table>
<thead>
<tr>
<th>Marker (location)</th>
<th>External primers</th>
<th>Internal primers</th>
<th>Restriction enzymes</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAG2 (VIII) 5′</td>
<td>F4: GCTACCTCGAACAGGAACAC</td>
<td>F4: GCATCAACAGTCTCTCGTGC</td>
<td>Sau 3AI</td>
</tr>
<tr>
<td>R4: GCATCAACAGTCTCTCGTGC</td>
<td>F3: TCTGGTTTCCCGAAATGACCTCC</td>
<td>R3: TCCAAAAGCTGCTATTCCGC</td>
<td>HhaI</td>
</tr>
<tr>
<td>P43S1: AACTCTCACCATTCACCACC</td>
<td>P43AS1: GCCGCTTGGTTAGACAAAGACA</td>
<td>P43AS2: CCAAGGAGACCGAGAAGGA</td>
<td>NciI</td>
</tr>
<tr>
<td>SAG3 (XI) 15′</td>
<td>GRA6-F1x: ATTTGTGTTCGAGCACTGGT</td>
<td>GRA6-F1: TTCCCGAGACGGCTGCT</td>
<td>MseI</td>
</tr>
<tr>
<td>GRA6-R1: GCACTCTGCGTGTGGTGTT</td>
<td>GRA6-R1: TCGCGGAGAGTTGACATAG</td>
<td>Btb (ext): TCCAAAATGAGAAATCGT</td>
<td>Taq I</td>
</tr>
<tr>
<td>BTUB (IX)</td>
<td>Btb (ext): TCCAAAATGAGAAATCGT</td>
<td>Btb-R: TTGTAGGAAACACCCGACGC</td>
<td>BsiE I</td>
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<tr>
<td>APICO (plastid)</td>
<td>Apico-o: TGTTTTTTAACCTTATGTTGG</td>
<td>Apico-o: TGCAATCTGGTATTGTGG</td>
<td>Afl II</td>
</tr>
<tr>
<td></td>
<td>Apico-o: AAACCGAAATTAATGAGATTGAA</td>
<td>Apico-o: AAACCGAAATTAATGAGATTGAA</td>
<td>Dde I</td>
</tr>
</tbody>
</table>

*This study.*
**RESULTS**

**PCR analysis.** Small amount or no amplification products were observed after the first step of amplification because of the low amount of parasite DNA present in clinical samples (data not shown). After the second round of amplification, positive amplicons were obtained for all samples with SAG2 (3′ and 5′), GRA6, and APICO PCRs. Only 13 and 8 samples were amplified using SAG3 and BTUB markers, respectively (Table 3). Negative controls remained free of amplified products.

**RFLP analysis.** Genotyping of samples by SAG2 PCR-RFLP revealed that the 14 samples were infected with SAG2 Type I parasites (Table 3). SAG3 marker showed that only one specimen harbored the type I allele, whereas six harbored the type III, one harbored the type II, and five harbored a mixture of alleles (Figure 1 and Table 3). GRA6 marker showed that 10 specimens harbored the type I allele, whereas four samples harbored a mixture of alleles (Figure 1 and Table 3). BTUB marker showed that three specimens harbored the type I allele, whereas three samples harbored the type III and two samples harbored a mixture of alleles (Figure 1 and Table 3). APICO marker showed that 13 specimens harbored the type I allele, whereas one sample harbored type III (Table 3). When results from each locus were combined, only one specimen harbored the type I allele in all studied loci. All the remaining samples had genotypes consisting of different combinations of alleles seen in the clonal types. Seven specimens were shown to possess a chimerical combination of allele types I–III, and three possessed a combination of allele types I–II. The last three specimens produced complex RFLP patterns with a mixture of two to three genotypes in the same sample (Figure 1 and Table 3).

**Sequencing analysis.** To confirm SAG3 type, all restriction fragment products were subjected to direct DNA sequence determination. All sequences were submitted to Genbank (accession numbers = GU139462–GU139479). The sample was identified, because allele 1 was identical to type I TgUgCh62 strain (accession number = EF585690). The sample identified as type 2 was identical to type II TgUgCh68 strain (accession number = EF585684), and the six samples identified as type 3 gave homology of 99–100% with type III TgUgCh64 strain (accession number = EF585687). Samples with restriction profile showing a mixture of two to three archetypal alleles revealed double peaks at known polymorphic sites, indicating the concomitant presence of two alleles in the same sample (Figure 2). It was interesting to note that the last sample with restriction profile showing a mixture of three archetypal alleles also revealed the presence of two strains; the first had 99% homology with the natural recombinant I/III P-Br strain (accession number = AY187280), and the second had 98% homology with type II TgUgCh68 strain (accession number = EF585684). No new single-nucleotide polymorphisms (SNPs) were found. For the GRA6 marker, any product that did not produce a single type after restriction analysis was sequenced. In four cases, double peaks of polymorphic sites were noted, revealing a double infection. Three were caused by type I allele (identical to TgUgCh83 strain; accession number = EF585715) and type II allele (identical to TgUgCh78 strain; accession number = EF585712), and one was caused by type I allele (identical to TgUgCh83 strain) and type III allele (giving 99% homology with TgUgCh64 strain; accession number = EF585705).

**DISCUSSION**

Very few studies concerning genotypic characterization of Toxoplasma strains associated with human toxoplasmosis in Africa have been reported. Two studies genotyped the SAG2 locus and were performed on samples from Ugandan human immunodeficiency virus (HIV) patients and Egyptian female patients with abortion and intrauterine fetal death, respectively.\(^{11,12}\) Both reported the predominance of genotype II. Using the same marker, all Tunisian samples were classified as SAG2 type I. Similar results were mentioned in some Mediterranean countries, such as Spain, where strains possessing the type I allele at the SAG2 locus were found in 6 of 13 cases of congenital infection.\(^{20}\) However, this marker is not able to detect recombinant or exotic strains, causing misclassification of them as genotype I.

The advantage of using a combination of different independent markers is that it is much more likely to detect recombinant genotypes. Recently, multiplex PCR of microsatellites for typing was developed.\(^{21}\) However, this method has some limitations; a special sequencer and analysis software are needed, and it has a low level of analytic sensitivity (at least 50 parasites in a sample). In contrast, the multilocus PCR–RFLP genotyping method developed by Khan and others\(^{19}\) offers the advantages

<table>
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<tr>
<th>Alleles</th>
<th>5′SAG2</th>
<th>5′SAG2</th>
<th>SAG3</th>
<th>Accession No.</th>
<th>GRA6</th>
<th>Accession No.</th>
<th>BTUB</th>
<th>APICO</th>
<th>Final genotype</th>
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</thead>
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<td>Sample</td>
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</tr>
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<tr>
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<td>1</td>
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<td>1</td>
<td>ND</td>
<td>1</td>
<td>Mix of I/II and I/III</td>
</tr>
</tbody>
</table>
that it is simple, can be conducted on small amounts of dif-
ferent types of samples, and has good sensitivity (detecting
between 5 and 10 parasites). Because our samples contained
small numbers of organisms (data not shown), this technique
was more suitable for their typing, and precautions were taken
to avoid contamination.

Using multilocus analysis, all Tunisian isolates examined
in the present study, except one, harbored recombinant I/II
and/or I/III strain, which is in concordance with the very few
results reported in Ugandan HIV patients and other patients
of African origin.\textsuperscript{10, 11, 21} Sequencing of different alleles revealed
a greater homology than those previously reported in African
free-ranging chickens.\textsuperscript{22, 23} Although the very limited clinical

data in the present study cannot allow hypothesis testing about
strain virulence, it is important to note the presence of strains
with close homology to the natural recombinant I/III P-Br
strain that was incriminated in severe cases in South America.\textsuperscript{6}
This suggests that this genotype could be a new emerging one
in this part of the world. In addition, other recombinant or
atypical strains may be present in Africa, which has been
proven recently by extensive sequence analysis of eight iso-
lates from Uganda.\textsuperscript{23} This study found mostly type II geno-
types, but they contained novel SNPs that suggested regional
allelic variants.\textsuperscript{23}

Perhaps, the most significant finding of our present work
compared with previously published data is the high frequency

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1}
\caption{Restriction analysis of PCR products directly amplified from clinical samples: PCR amplifications of \textit{SAG3} marker (A), \textit{GRA6} marker (B), and \textit{BTUB} marker (C). Lane PM = 100-basepair (bp) DNA ladder (Amersham, Buckinghamshire, UK); lane ND = undigested PCR product; lanes RH, PRU, and NED = digested PCR products representative of \textit{T. gondii} strain types I, II, and III, respectively; lane T = negative con-
trol; lanes 1, 5, 9, and 12 = digested PCR products of positive samples AF 06/06, AF 26/04, PL 05/06, and PL 04/06 genotyped as type I, type I/III, type
I/II, and mix of strains, respectively. Note that the last two specimens produce a complex digestion pattern with \textit{GRA6} and \textit{SAG3}, respectively.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2}
\caption{Sequence analysis of the \textit{SAG3} marker using reverse primer, which reveals double peaks at known polymorphic sites (sample AF 33/05). This figure appears in color at www.ajtmh.org.}
\end{figure}
of apparent concomitant infection in clinical samples. Mixed infection with *T. gondii* strains has been previously reported in England and Wales.24 This finding was explained by the ingestion of more than one type of parasite in food products containing meat originating from multiple animals.25 However, the high proportion of mixed infections shown in African isolates from free-ranging chickens underlines the high frequency in this part of the world of sequential infections with parasites of different types acquired as oocysts directly from the environment.22 It also suggests that intermediate hosts are infected with more than one strain. Congenital toxoplasmosis is, in most cases, the result of a primary infection acquired by an immunologically naïve patient during pregnancy. It seems that we can exclude sequential infection as an explanation for the presence of different genotypes in a significant number of patients. Similarly, it is hard to explain the observed frequency of mixed infections as a consequence of exposure to mixed oocysts.26 The possible mechanism leading to the observed mixed infection is the ingestion of tissue cysts from infected meat.27 Mutton is the meat most commonly incriminated in the transmission of toxoplasmosis in Tunisia, and it could be seen as a real source of contamination.28 Genotypic characterization of *T. gondii* in sheep in Tunisia could help to establish correlation between mixed infections in humans in Tunisia and contaminated mutton.

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