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Sequence Variation in Superoxide Dismutase Gene of Toxoplasma gondii among Various Isolates from Different Hosts and Geographical Regions

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Abstract: *Toxoplasma gondii*, an obligate intracellular protozoan parasite of the phylum Apicomplexa, can infect all warmblooded vertebrates, including humans, livestock, and marine mammals. The aim of this study was to investigate whether superoxide dismutase (SOD) of *T. gondii* can be used as a new marker for genetic study or a potential vaccine candidate. The partial genome region of the SOD gene was amplified and sequenced from 10 different *T. gondii* isolates from different parts of the world, and all the sequences were examined by PCR-RFLP, sequence analysis, and phylogenetic reconstruction. The results showed that partial SOD gene sequences ranged from 1,702 bp to 1,712 bp and A + T contents varied from 50.1% to 51.1% among all examined isolates. Sequence alignment analysis identified total 43 variable nucleotide positions, and these results showed that 97.5% sequence similarity of SOD gene among all examined isolates. Phylogenetic analysis revealed that these SOD sequences were not an effective molecular marker for differential identification of *T. gondii* strains. The research demonstrated existence of low sequence variation in the SOD gene among *T. gondii* strains of different genotypes from different hosts and geographical regions.

Key words: Toxoplasma gondii, PCR-RFLP, superoxide dismutase, sequence variation, phylogenetic analysis

INTRODUCTION

Toxoplasma gondii, the direct pathogenic factor of toxoplasmosis, is an obligate intracellular protozoan parasite of the phylum Apicomplexa which can infect all warm-blooded vertebrates, including humans, livestock, and marine mammals [1]. Although most infections are clinically asymptomatic, the parasite can cause severe disease in immunocompromised populations and congenitally infected individuals [1,2]. In addition, infections in domestic animals may result in economic losses as well as bring enormous psychological troubles, since it can cause abortion, stillbirth, and neonatal loss [3].

Unlike many other Apicomplexa parasites which exhibit stronger host specificity, *T. gondii* faces vastly numerous hosts and can adapt to various environmental conditions during its complex life cycles, which can be contributed to many differ-

© 2015, Korean Society for Parasitology and Tropical Medicine This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0) which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited. ent *T. gondii* strains and genotypes [4]. It was popularly believed that *T. gondii* had a clonal population structure with 3 predominant lineages, namely types I, II, and III [5-7]. Besides these isoforms, it also exists in atypical and recombinant strains [8,9]. To better understand the population genetics and molecular epidemiology of this parasite, and to develop more strategies for vaccination, diagnosis, and treatment of toxoplasmosis, it is necessary to study the genetic diversities in *T. gondii* [10,11].

Superoxide dismutase (SOD), an important enzyme that widely exists in many organisms, including animals, plants, and microorganisms, can promote the conversion of superoxide (O^{2-}) into hydrogen peroxide and oxygen [12,13]. In view of SOD that can eliminate extra superoxide (O^{2-}) anion in the cells and protect cells from oxidative damages, it has potential applications in medicine, food industry, and agriculture [12,14,15]. In *T. gondii*, limited studies have shown that SOD is a typical FeSOD and its activity might be essential for the in-tracellular growth of both bradyzoite and tachyzoite forms [16]. To our knowledge, no report has described the sequence variation of SOD gene in different *T. gondii* strains. We hereby examined sequence variation of SOD gene among 10 *T. gondii*

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isolates from different hosts and geographical regions (different countries), and assess SOD could be used as a new marker for genetic study or a potential vaccine candidate against *T. gondii*.

MATERIALS AND METHODS

T. gondii strains

A total of 10 genotyped *T. gondii* strains were utilized as shown in Table 1, and the genomic DNA was prepared as described previously [17].

PCR amplification

The SOD gene was amplified by PCR from genomic DNA of *T. gondii* with 1 pair of primes: 5'-ATGGTATTCACTTTGCCCCC-GCT-3' (forward prime) and 5'-TCATTTCAAGGCATTCTC-CAAG-3' (reverse primer). The design of primers was based on SOD gene of *T. gondii* RH isolate available in GenBank database

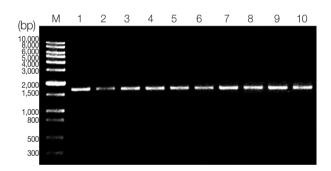


Fig. 1. Analysis of PCR products of SOD gene from 10 examined *T. gondii* strains using electrophoresis. Lane M; DNA size marker 10,000. Lanes 1-10; *T. gondii* type I (GT1, RH), type II (PTG, PRU), type III (CTG) strains and other strains (TgCgCa1, MAS, Tg-CatBr5, TgCatBr64, and TgToucan), respectively.

(AF029915). The amplification reaction was performed in a volume of 20 µl containing 2 µl template DNA, 10 µl 2×1 Step buffer (0.5 U *Taq* polymerase), 1 µl of each primer, and 6 µl RNase-free dH₂O. The target DNA was amplified under the following conditions: 94°C for 30 sec, 63.6°C for 1 min, and 72°C for 1 min. The PCR amplification products were confirmed by electrophoresis in a 1.5% agarose gels and staining with ethidium bromide followed by visualization under UV.

The analysis of PCR-RFLP

The SOD PCR amplification products from representative *T. gondii* strains were digested with restriction enzymes *Xba* I and *EcoR* I, respectively, and incubated at 37°C for 3 hr. The restriction fragments were separated by electrophoresis in 1.5% aga-

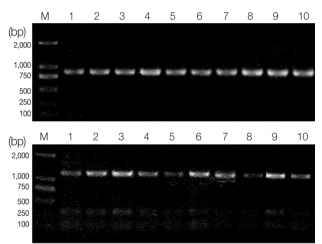


Fig. 2. PCR-RFLP analysis of SOD genes using restriction endonucleases *EcoR* I (upper) and *Xba* I (bottom). Lane M; DNA size marker 2,000. Lanes 1-10; *T. gondii* type I (GT1, RH), type II (PTG, PRU), type III (CTG) strains and other strains (TgCgCa1, MAS, TgCatBr5, TgCatBr64, and TgToucan), respectively.

Table 1. Toxoplasma gondii strains subjected to SOD gene sequence analysis

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Strain	Host	Geographical origin	Genotype ^a		
RH	Human	France	Reference, Type I, ToxoDB #10		
GT1	Goat	United States	Reference, Type I, ToxoDB #10		
PTG	Sheep	United States	Reference, Type II, ToxoDB #1		
Prugniaud (PRU)	Human	France	Reference, Type II, ToxoDB #1		
CTG	Cat	United States	Reference, Type III, ToxoDB #2		
TgCgCa1	Cougar	Canada	Reference, ToxoDB #66		
MAS	Human	France	Reference, ToxoDB #17		
TgCatBr5	Cat	Brazil	Reference ToxoDB #19		
TgCatBr64	Cat	Brazil	Reference ToxoDB #111		
TgToucan (TgRsCr1)	Toucan	Costa Rica	Reference, ToxoDB #52		

^aBased on genotyping results of Su et al. [17].

rose gel and staining with ethidium bromide, followed by visualization under UV.

Sequence analysis and reconstruction of phylogenetic relationships

The SOD PCR products were purified with the DNA purification kit (TransGen Biotech, Beijing, China) and ligated with the pEASY-T1 vector (TransGen Biotech) according to the manufacturer's instructions, and then transformed into *Escherichia coli* DH5α competent cells. The transformed cells carrying the insert were successively selected by blue-white screening, PCR, and restriction enzyme digestion. The positive colonies were sequenced by Beijing Genomics Institute Company (Beijing, China). The obtained SOD gene sequences from dif-

Table 2. Nucleotide polymorphisms of the SOD gene coding region within Toxoplasma gondii strains

	RH	CTG	GT1	MAS	PRU	PTG	TgCatBr5	TgCatBr64	TgCgCa1	TgToucan
1403	А	*	*	*	*	*	*	С	*	*
1437	А	*	*	*	*	*	*	Т	*	*
1480	G	*	*	*	*	*	*	С	*	*
1501	G	*	*	*	*	*	-	*	-	*
1503	G	*	*	*	*	*	Т	*	*	*
1504	G	*	*	*	*	*	*	*	Т	*
1518	А	*	*	*	*	*	Т	*	*	*
1519	С	*	*	*	*	*	А	*	*	*
1520	Т	*	*	*	*	*	С	*	*	*
1521	G	*	*	*	*	*	Т	*	*	*
1532	С	*	*	*	*	*	Т	*	*	*
1535	С	*	*	*	*	Т	*	*	*	*
1545	А	*	*	*	*	*	*	*	G	*
1547	G	*	*	*	*	*	*	С	*	*
1549	А	*	*	*	*	*	*	*	G	*
1555	С	*	*	*	*	*	Т	*	*	*
1556	А	*	*	*	*	*	Т	*	*	*
1557	Т	*	*	*	*	*	С	*	*	*
1558	С	*	*	*	*	*	Т	*	*	*
1566	А	*	*	*	*	*	G	*	*	*
1567	А	*	*	*	*	*	*	*	Т	*
1568	С	*	*	*	*	*	*	*	Т	*
1574	G	*	*	*	*	А	*	*	*	*
1575	G	*	*	*	*	*	А	*	*	*
1584	Т	*	*	*	*	*	*	С	*	*
1585	С	*	*	*	*	*	*	-	*	*
1586	G	*	*	*	*	*	*	-	*	*
1587	А	*	*	*	*	*	*	G	*	*
1590	G	*	*	*	*	А	А	*	*	*
1599	G	*	*	*	*	*	*	*	А	*
1601	С	*	*	*	*	*	*	*	-	*
1614	G	*	*	*	*	А	*	*	*	*
1616	Т	*	*	*	*	G	*	*	*	*
1618	G	*	*	*	*	Т	*	*	*	*
1620	G	*	*	*	*	*	*	*	А	*
1628	Т	*	*	*	*	А	*	*	*	*
1635	G	*	*	*	*	*	*	*	*	С
1637	С	*	*	*	*	*	-	*	*	*
1668	A	*	*	*	*	*	*	*	-	*
1669	С	*	*	*	*	*	А	*	*	*
1679	G	*	*	*	*	*	A	*	*	*
1685	C	*	*	*	*	А	*	*	*	*
1793	C	*	*	*	*	*	-	*	*	*

Numbers in the left column indicate positions of variable nucleotides. Asterisks (*) indicate identical nucleotides related to the sequence of RH (first column). Dashes (-) indicate deletions.

ferent *T. gondii* strains were aligned using Multiple Sequence Alignment Program Clustal ×1.83. Phylogenetic reconstructions based on the sequences of SOD gene among different *T. gondii* strains were performed using 3 methodologies, namely maximum parsimony (MP), Bayesian inference (BI), and maximum likelihood (ML) [18,19].

RESULTS

PCR-RFLP and sequence analysis

The amplification of the SOD gene resulted in a single product of approximate 1,700 bp in length on agarose gel for all tested *T. gondii* strains (Fig. 1). There are no obvious differences in all tested *T. gondii* strains after digestion of the amplified SOD products with *EcoR* I and *Xba* I, revealing that subtypes I, II, and III could not be differentiated in this condition (Fig. 2). Then, the amplicons of all isolates were sequenced. The sequencing results showed that the SOD gene was 1,712 bp in length for the TgCatBr64, 1,709 bp in length for PTG, 1,706 bp in length for RH, 1,702 bp in length for TgCgCa1, and 1,707 bp for the remaining 6 strains. Besides, the A + T contents of these sequences ranged from 50.1% to 51.1% among all isolates. The alignment of all 10 sequences revealed nucleotide polymorphisms at 43 positions, with an intraspecific variation of 0-1.0% (Table 2). Of these variable nucleotide positions, there were 29 transitions (C \leftrightarrow T, T \leftrightarrow G, A \leftrightarrow C, and A \leftrightarrow G), 7 transversions (A \leftrightarrow T and C \leftrightarrow G), and 7 deletions in all the sequences. These results showed 97.5% sequence similarity of SOD gene among all the tested isolates, suggesting considerable sequence diversity. Amino acid polymorphisms at 24 positions were detected among 10 *T. gondii* strains, including 20 amino acid position substitutions and 4 amino acid position deletions (Table 3).

Phylogenetic reconstruction

Phylogenetic analysis of the 10 examined *T. gondii* isolates based on the SOD gene sequences demonstrated 2 major clusters (Fig. 3). TgCatBr5, TgCatBr64, and TgCgCa1 strains were clustered in 1 clade, and PRU, MAS, GT1, CTG, TgTgucan, PTG, and RH were clustered in the other clade. Overall, the to-

Table 3. Amino acid changes of the SOD gene coding region among ten Toxoplasma gondii strains

	RH	CTG	GT1	MAS	PRU	PTG	TgCatBr5	TgCatBr64	TgCgCa1	TgToucan
66	N	*	*	*	*	*	*	Y	*	*
80	S	*	*	*	*	*	*	Т	*	*
87	G	*	*	*	*	*	-	*	-	*
88	G	*	*	*	*	*	С	*	V	*
93	Т	*	*	*	*	*	*	S	*	*
94	G	*	*	*	*	*	*	W	*	*
102	K	*	*	*	*	*	Ν	*	Е	*
103	E	*	*	*	*	*	*	*	G	*
105	Т	*	*	*	*	*	I	*	*	*
106	S	*	*	*	*	*	F	*	*	*
109	Ν	*	*	*	*	*	*	*	I	*
112	D	*	*	*	*	*	Ν	*	*	*
115	S	*	*	*	*	*	*	Р	*	*
116	К	*	*	*	*	*	*	Е	*	*
117	V	*	*	*	*	I	I	*	*	*
120	G	*	*	*	*	*	*	*	S	*
125	G	*	*	*	*	S	*	*	*	*
126	W	*	*	*	*	F	*	*	*	*
127	А	*	*	*	*	*	*	*	Т	*
129	L	*	*	*	*	Н	*	*	*	*
132	D	*	*	*	*	*	-	*	*	Н
143	Т	*	*	*	*	*	Ν	*	-	*
148	Т	*	*	*	*	Ν	*	*	*	*
151	Т	*	*	*	*	*	-	*	*	*

Numbers in the left column indicate positions of variable amino acids. Asterisks (*) indicate identical amino acid related to the sequence of RH (first column). Dashes (-) indicate deletions.

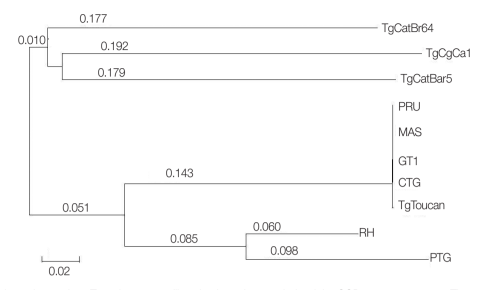


Fig. 3. Phylogenetic analyses of 10 *Toxoplasma gondii* strains based on analysis of the SOD gene sequences. The tree was built by Bayesian inference (BI) analysis. The numbers along branches indicate bootstrap values resulting from BI analysis.

pologies of all trees based on nucleotide sequences inferred by 2 different methods were similar, with only small differences of bootstrap values.

DISCUSSION

SOD widely exists in many organisms and plays a crucial role in eliminating the extra superoxide anion (O^{2-}) in the cells to avoid oxidative damages [12-15]. In *T. gondii*, SOD is an iron-containing type, which correlates with the protection of the parasite [16]. However, SOD enzymes from C and RH strain tachyzoites do not appear to be the basis for differences in virulence to mice [20]. In the present study, we cloned and sequenced the partial genome sequence of SOD gene among 10 *T. gondii* isolates from different hosts and geographical regions and examined genetic diversity of SOD locus by the techniques of PCR-RFLP, sequence analysis, and phylogenetic reconstruction. The results revealed nucleotide polymorphisms at 43 positions and amino acid polymorphisms at 24 positions, suggesting low sequence variability among all the tested isolates.

PCR-RFLP has been widely used in analysis of specific genetic loci for *T. gondii* genotyping. Multilocus PCR-RFLP marker is a high resolution for identification of parasites, although it requires a huge investment of time to test and optimize each marker [11]. By contrast, single PCR-RFLP marker is simply and more convenient. Then, studies on single marker loci (e.g. GRA5, GRA6, and ROP17) have shown signs of positive selection, and could be sufficient for genotyping of *T. gondii* isolates. [21-23].

In our study, however, 3 clonal lineages (types I, II, and III) cannot be differentiated in 10 examined *T. gondii* strains using a single PCR-RFLP marker SOD. One possible reason is that the sequence variation of SOD could not be located in restriction enzyme sites, to some degree, leading to loss of many polymorphisms in this situation [24].

The direct sequencing of genomic regions can detect small deletions and insertions (e.g., indels) and single nucleotide polymorphisms (SNPs) in the genomic regions, and hence it is capable of testing more genetic diversity compared with PCR-RFLP. Based on the partial sequences of the SOD locus, phylogenetic analysis revealed 2 major clusters with only a little difference of bootstrap values, and it cannot differentiate *T. gondii* strains to their genotypes, implying a low genetic diversity. Our results were similar to some studies such as MIC13, eIF4A [25,26] and different from other genetic markers including GRA5, GRA6, and ROP17 [21-23]. The low sequence variation (0-1.0%) in the partial SOD gene suggests that SOD gene could not be an ideal genetic marker for differentiation of the *T. gondii* strains or intraspecific phylogenetic analyses.

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CONFLICT OF INTEREST

We have no conflict of interest related to this work.

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