Genotypic Diversity of *Theileria orientalis* Detected from Cattle Grazing in Kumamoto and Okinawa Prefectures of Japan

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**ABSTRACT.** *Theileria orientalis* is a benign protozoan species that is widely distributed in Japan, yet sometimes causes serious economic losses in the livestock industry. In this study, we conducted a molecular survey based on genes encoding the major piroplasm surface protein (MPSP) and p23 for *T. orientalis* detected in cattle grazing in southern areas of Japan. In the Kyushu and Okinawa areas, the southern prefecture of Japan, and the southern islands of Japan consisting of 2 farms in Kumamoto prefecture (Aso and Kuma districts) and 3 farms in Okinawa prefecture (Ishigaki, Iriomote, and Yonaguni Islands). High prevalence rates of *T. orientalis* infection were observed in all the cattle populations using the diagnostic MPSP- and p23-PCR assays. Phylogenetic analyses revealed 4 MPSP genotypes and 3 p23 genotypes. Furthermore, MPSP genotype-specific PCR methods were developed in this study and wide distributions of 5-district genotypes of *T. orientalis* were observed for the examined farms. Our results indicate that at least 5 types of *T. orientalis* exist in Kumamoto and Okinawa prefectures of Japan and that genotype-specific PCR assays are highly applicable for the quarantine of transported cattle and for epidemiological surveys of bovine theileriosis in Japan.

**KEY WORDS:** cattle, genotype-specific PCR, Japan, MPSP gene, *Theileria orientalis*.

*Theileria orientalis*, a tick-transmitted, intraerythrocytic protozoa belonging to the phylum Apicomplexa [4], is the causative organism of bovine theileriosis in Japan. *T. orientalis* is a member of the relatively benign *Theileria* group (*Theileria sergenti/buffeli/orientalis*) that is widely distributed throughout Japan [17]. Generally, it shows a lower pathogenicity in cattle compared to *Theileria parva* and *Theileria annulata*; however, it sometimes causes clear symptoms, including fever, anemia, and anorexia [13]. Flumethrin pour-on treatment is currently used to control the parasite in grazing cattle by reducing the number of vector ticks during pasture [11]. However, many farmers in Japan experience losses in livestock production because of the consequences of bovine theileriosis [7, 12]. In the Kyushu and Okinawa areas, the southern islands of Japan, the livestock industry for cattle has prospered; however, the practice of pasturing cattle to reduce rearing costs is associated with a high risk of contracting theileriosis.

Major piroplasm surface protein (MPSP) and p23 of *T. orientalis* are major immunodominant proteins expressed on the surface of the parasite during its intraerythrocytic (piroplasm) stage and show significant sequence diversities among field isolates of *T. orientalis* [4, 5, 7, 9, 10, 19]. *T. orientalis* populations are currently known to consist of 8 genotypes worldwide, based on all registered MPSP gene sequences [3, 6, 18]. Our recent survey carried out in Hokkaido prefecture of the northern island of Japan, demonstrated that there are at least 4 genotypes of *T. orientalis* (types 1, 2, 4, and 5) in Japan [7]. Thus, the MPSP gene has potential as a novel molecular marker for epidemiological study of *T. orientalis*.

In this study, we analyzed blood samples collected from 70 cattle (Holstein and Japanese Black) grazing in the Kyushu and Okinawa areas, consisting of 2 farms in Kumamoto prefecture (Aso and Kuma districts) and 3 farms in Okinawa prefecture (Ishigaki, Iriomote, and Yonaguni Islands) in Japan. By determining the sequences of *T. orientalis* MPSP and p23 genes that were detected in the infected cattle, we identified the genotype diversities for this parasite in the southern prefecture of Japan. Furthermore, we developed 5 different genotype-specific PCR methods and identified the district-based distribution of *T. orientalis* genotypes in each of the grazing cattle populations. Herein, we report the current molecular status of *T. orientalis* distributed in Kumamoto and Okinawa prefectures of Japan and discuss the future utility of genotype-specific PCR methods.

**MATERIALS AND METHODS**

**Blood samples:** Cattle examined in this study were grazed on each of 5 farms located in 2 areas of Kumamoto prefe-
ture (Aso and Kuma districts) and 3 areas of Okinawa prefecture (Ishigaki, Iriomote, and Yonaguni Islands) in Japan (Fig. 1). Farms were identified as *T. orientalis*–prevalent areas by field veterinarians. Holstein cattle aged from 8 months to 2 years were grazed for the first time on the Kuma district farm, while Japanese Black cattle from the Aso district farm were of various ages (3 to 14 years) and grazing histories. Japanese Black cattle in Okinawa prefecture also were of various ages (Ishigaki Island, from 8 months to 14 years; Iriomote Island, 7 months to 2 years; Yonaguni Island, from 7 years to 19 years) and had various grazing histories. All cattle were originally born in the same prefectures, except for three cattle in Aso district of Kumamoto prefecture that were born in other neighbor prefectures (either of Kagoshima or Miyazaki prefecture in Kyushu area) and then introduced to the Aso district. Three cattle in Iriomote Island were male and all other cattle were female. A total of 70 blood samples were collected from the grazing cattle late in the pasturage period during 2007 (Table 1).

**Diagnoses of parasitemia and anemia:** Approximately 2 mL of blood was collected from the tail veins of cattle and added to ethylenediaminetetraacetic acid (EDTA) to a final concentration of 0.15–0.22%. Thin blood smears were made and then stained with Giemsa solution for classic microscopic diagnosis as described previously [7]. In addition, the hematocrit (HCT) values of the respective blood samples were determined and anemia was diagnosed if the HCT value was <24% [2].

**DNA extraction and plasma preparation:** Genomic DNA was extracted from 100 μl of blood samples using a QIAamp® DNA Blood Mini Kit (Qiagen, Hilden, Germany) and 200 μl of eluted DNA (approximately 6 μg/μl) were obtained and stored at 30°C for subsequent PCR analyses. Blood plasma fractions were obtained by centrifugation of blood samples at 700 × g for 15 min at 4°C and then stored at −30°C for enzyme-linked immunosorbent assay (ELISA).

**PCR:** A classic universal PCR (product size: 857 base pairs [bp]) assay, which had been previously designed for the MPSP (*p*32/33/34) gene of *T. orientalis* [14], and 2 kinds of recently developed PCR assays, MPSP-PCR (776 bp) and p23-PCR (601 bp) assays [7], were used for the diagnostic detection of *T. orientalis* genes, according to the previously described procedures.

Here, we designed 5 new primer pairs by aligning all of the registered sequences of *MPSP* genes listed in Fig. 2. These primer pairs were designed to specifically amplify the target DNA fragments for each of the 5 genotypes of *MPSP* genes (types 1–5 in Fig. 2) in the respective PCR analyses. Primer sequences, annealing temperatures and the resultant product sizes are summarized in Table 2. Each of the type-specific primer pairs specifically anneal to the conserved regions of *MPSP* gene sequences categorized only in the corresponding genotype groups. The PCR methods were evaluated as new diagnostic assays to survey actual type-
DIVERSITY OF *T. ORIENTALIS* IN JAPAN

Table 1. Diagnostic history of grazing cattle in southern areas of Japan

<table>
<thead>
<tr>
<th>Prefectures</th>
<th>Districts (Species)</th>
<th>Date&lt;sup&gt;a&lt;/sup&gt; (Species)</th>
<th>Total cattle</th>
<th>Microscopic test&lt;sup&gt;b&lt;/sup&gt;</th>
<th>ELISA&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Anemia&lt;sup&gt;a&lt;/sup&gt;</th>
<th>PCR Universal&lt;sup&gt;b&lt;/sup&gt;</th>
<th>p23&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kumamoto</td>
<td>Aso (Japanese Black)</td>
<td>12/18 (15)</td>
<td></td>
<td>15 (100.0)</td>
<td>15 (100.0)</td>
<td>15 (100.0)</td>
<td>15 (100.0)</td>
<td>15 (100.0)</td>
</tr>
<tr>
<td></td>
<td>Kuma (Holstein)</td>
<td>11/19 (35)</td>
<td></td>
<td>28 (80.0)</td>
<td>1 (3.0)</td>
<td>31 (88.6)</td>
<td>30 (85.7)</td>
<td>33 (94.3)</td>
</tr>
<tr>
<td>Okinawa</td>
<td>Ishigaki (Japanese Black)</td>
<td>9/4,14 (10)</td>
<td></td>
<td>10 (100.0)</td>
<td>0 (0.0)</td>
<td>6 (60)</td>
<td>9 (90.0)</td>
<td>10 (100.0)</td>
</tr>
<tr>
<td></td>
<td>Iriomote (Japanese Black)</td>
<td>11/28 (5)</td>
<td></td>
<td>5 (100.0)</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
<td>5 (100.0)</td>
<td>5 (100.0)</td>
</tr>
<tr>
<td></td>
<td>Yonaguni (Japanese Black)</td>
<td>12/5 (5)</td>
<td></td>
<td>5 (100.0)</td>
<td>0 (0.0)</td>
<td>4 (80.0)</td>
<td>4 (80.0)</td>
<td>5 (100.0)</td>
</tr>
</tbody>
</table>

<sup>a</sup> Month/Date.
<sup>b</sup> Number (%) of cattle showing positive reactions (of total cattle).
<sup>c</sup> Number (%) of cattle showing clinical anemia in a group of MPS-P PCR-positive cattle.

specific distributions of *T. orientalis* in each cattle population.

Type-specific PCR methods were performed using 1 µl of stored DNA template extracted from blood samples, mixed with 9 µl of reaction buffer consisting of 0.05 µl of Ex Taq DNA polymerase (Takara, Tokyo, Japan), 0.1 µl of 10 µM each primer, 1 µl of 10 × Ex Taq Buffer (Takara), 0.8 µl of dNTP mixture (2.5 mM each, Takara), and 6.95 µl of double distilled water (DDW). Reactions were carried out using an iCycler Thermal Cycler (Bio-Rad Laboratories, Hercules, CA, U.S.A.) for 35 cycles. After an initial denaturation step for 5 min at 94°C, amplification consisted of denaturation for 30 sec at 94°C, annealing for 45 sec at the indicated temperature (Table 2) and extension for 1 min at 72°C, followed by an additional 7 min at 72°C. The amplified PCR products were subjected to 2.0% agarose gel electrophoresis, stained with ethidium bromide and then visualized under an ultraviolet light.

ELISA: Classic serodiagnostic ELISA was performed using a crude parasite antigen to assess the presence of *T. orientalis*-specific antibodies in the collected blood plasma fractions, as described previously [7].

DNA sequencing and phylogenetic analyses: DNA sequencing analyses of PCR fragments were carried out to examine the diversities of MPS-P and p23 genes detected from the blood samples, as described previously [7]. At least 10, 12, and 15 blood samples were randomly selected from the diagnostic PCR-positive samples from Aso, Kuma, and Okinawa farms, respectively. DNA fragments were amplified by another PCR step for subsequent DNA sequencing of MPS-P and p23 genes [7]. In detail, PCR was performed using 2 µl of the selected DNA template, mixed with 18 µl of reaction buffer, consisting of 0.2 µl of Expand HiFi Plus Enzyme Blend (Roche Applied Science, Basel, Switzerland), 0.2 µl of 10 µM of each primer, 4 µl of 5 × Expand HiFi Plus Reaction Buffer with 7.5 mM MgCl₂, 0.4 µl of 10 mM PCR Grade Nucleotide Mix (Roche Applied Science) and 13 µl of DDW. All PCR cycles consisted of an initial denaturation step for 2 min at 94°C, followed by 35 cycles for 1 min at 94°C, annealing for 30 sec at 58°C and extension for 1 min at 72°C, followed by an additional 7 min at 72°C. The amplified PCR products were cloned into a pCR2.1-TOPO vector, according to the manufacturer's instructions for the TOPO TA cloning kit (Invitrogen, Carlsbad, CA, U.S.A.). DNA sequencing of the inserts was performed commercially (Sigma-Aldrich, Tokyo, Japan). The CLUSTAL X program (University College Dublin, Dublin, Ireland) was used to align the obtained sequences and a neighbor joining method using an NJplot program [8] was used to construct 2 phylogenetic trees based on MPS-P and p23 genes. The accuracy of the phylogenetic tree branches was confirmed by a bootstrap method [1]. Representative sequences obtained in this study were registered in the GenBank database (National Center for Biotechnology Information, National Institutes of Health) (Figs. 2 and 3).

RESULTS

Detection of *T. orientalis* from grazing cattle: Although only 1 cattle showed anemia (Holstein in Kuma farm) among all the examined animals, results of the diagnostic MPS-P and p23-PCR assays, together with classic microscopic test and serodiagnostic ELISA, demonstrated high prevalent rates of *T. orientalis* in all populations examined (Table 1). As the case of a previous report [7], diagnostic MPS-P and p23-PCR assays were proven to show higher sensitivities for the detection of *T. orientalis*, as compared to the classic universal PCR assay. In addition, Japanese Black cattle in Iriomote Island gave a negative reaction when tested with the serodiagnostic ELISA, although the parasite was detected in all cattle, even using microscopy.

Phylogenetic analyses of isolated MPS-P and p23 genes: For DNA sequencing of MPS-P and p23 genes, we randomly selected the PCR-positive blood samples from Aso, Kuma,
Fig. 2. Phylogenetic tree of the MPSP genes based on the results from blood samples collected in Kumamoto and Okinawa prefectures, together with previously registered sequences from the GenBank database. MPSP gene sequences determined in this study are shown (in boldface), together with the number of collected samples. The representative sequences of isolated MPSP genes refer to the GenBank accession numbers, as indicated at the end of each branch. The length of the horizontal bar indicates the number of nucleotide substitutions per site. Numbers shown at the branch nodes indicate the bootstrap values.
Table 2. Primer sequences for *T. orientalis* genotype-specific PCR assays

<table>
<thead>
<tr>
<th>Primer Names</th>
<th>Sequences</th>
<th>Annealing (°C)</th>
<th>Product sizes</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPSP1</td>
<td>Fa) 5'-ttg cct agg ata ctt cet cet cg-3'</td>
<td>64</td>
<td>559</td>
</tr>
<tr>
<td></td>
<td>Rb) 5'-tgc ggt gta ttt ggc ctt c-3'</td>
<td>58</td>
<td>550</td>
</tr>
<tr>
<td>MPSP2</td>
<td>F 5'-cgc atc aag aca ctc aag gtc-3'</td>
<td>58</td>
<td>550</td>
</tr>
<tr>
<td></td>
<td>R 5'-cac tgt tca tgg cgt gca-3'</td>
<td>66</td>
<td>228</td>
</tr>
<tr>
<td>MPSP3</td>
<td>F 5'-agt cgc cca cag aat taa gca ta-3'</td>
<td>56</td>
<td>424</td>
</tr>
<tr>
<td></td>
<td>R 5'-ttt ccc gta cta ctt cgt gca-3'</td>
<td>55</td>
<td>326</td>
</tr>
<tr>
<td>MPSP4</td>
<td>F 5'-aga agt cga cgg ggg att teg cat c-3'</td>
<td>56</td>
<td>424</td>
</tr>
<tr>
<td></td>
<td>R 5'-ctg cga tga atg cgt cga cca ccc aac-3'</td>
<td>55</td>
<td>326</td>
</tr>
<tr>
<td>MPSP5</td>
<td>F 5'-tga agg atc ace acc atc cag-3'</td>
<td>55</td>
<td>326</td>
</tr>
<tr>
<td></td>
<td>R 5'-agt cgg tga cgt cag ctt ctt cca ccc aac-3'</td>
<td>55</td>
<td>326</td>
</tr>
</tbody>
</table>

*a*) Forward primer. *b*) Reverse primer.

Fig. 3. Phylogenetic tree of the *p23* genes based on the results from blood samples collected in Kumamoto and Okinawa prefectures, together with previously registered sequences from the GenBank database. The *p23* gene sequences determined in this study are shown (in boldface), together with the number of collected samples. The representative sequences of isolated *p23* genes refer to the GenBank accession numbers, as indicated at the end of each branch. The length of the horizontal bar indicates the number of nucleotide substitutions per site. Numbers shown at the branch nodes indicate the bootstrap values.
and Okinawa farms. Consequently, 47 and 38 PCR fragments derived from MPSP and p23 genes, respectively, were isolated and complete sequences determined. From the phylogenetic analysis of identified MPSP genes, we detected 4 genotypes of MPSP genes that were categorized as types 1, 2, 3, and 5 (Fig. 2). In Kuma farm, only type 2 (Ikeda type) of MPSP genes was isolated, while 3 district types of MPSP genes, including the type 2 gene, were detected from Aso and Okinawa farms. In contrast, all 3 genotypes of the p23 gene, consisting of Buffeli, Chitose, and Ikeda types, were successfully detected in this study, based on phylogenetic analysis of previously identified p23 genes (Fig. 3). In Kuma farm, only the Ikeda type of p23 gene was isolated, while all 3 types of the p23 gene were detected from Aso and Okinawa farms. Consequently, 4 and 3 types of MPSP and p23 genes, respectively, were detected in this study.

Wide distributions of 5 district genotypes of *T. orientalis* in cattle populations: Four genotypes of MPSP gene (types 1, 2, 3, and 5) were detected in the southern prefectures of Japan, in contrast to the different set of MPSP genotypes (type 1, 2, 4, and 5) that we reported in a previous study for the northern area [7]. These findings indicate that at least 5 genotypes of *T. orientalis* (types 1–5) currently exist in Japan. Therefore, we designed 5 genotype-specific PCR methods that can identify the 5 genotypes of MPSP gene (Table 2), based on all the registered sequences of MPSP genes listed in Fig. 2. The specificities of the genotype-specific PCR methods were evaluated by using the respective plasmids containing the 5 genotypes of MPSP gene as templates. Type 1 PCR method amplified a 559-bp DNA band specific to the type 1 (Chitose type) MPSP gene and did not amplify any other MPSP genotypes (Fig. 4). The other type-specific PCR assays also exhibited the same specificities to the target types of MPSP gene. PCR evaluation of all plasmid samples, which we had constructed for the 5 types of MPSP gene in the present and previous studies (Fig. 2) [7], showed no change in the high specificities of genotype-specific PCR methods (data not shown).

Finally, we surveyed the actual distributions of the 5 genotypes of *T. orientalis* in each of the cattle populations using the developed genotype-specific PCR assays (Fig. 5). In Kumamoto prefecture, all 5 genotypes were highly detectable in the Aso population, while a dominant infection of *T. orientalis* type 2 was observed in the Kuma population. In contrast, only types 1 and 2 were detected in the Iriomote population, while all 5 genotypes were distributed in the Ishigaki and Yonaguni populations. Overall, the type-specific
PCR assays demonstrated wide distributions of the 5 district genotypes of *T. orientalis* in Kumamoto and Okinawa prefec-
tures of Japan.

**DISCUSSION**

In this study, at least 5 genotypes of *T. orientalis* (types 1–5) were shown to currently exist in the southern areas of Japan. Combined with previous reports [6, 18], this study indicates that a total of 7 genotypes of *T. orientalis* possibly exist in Japan, based on *MPSP* gene sequences (types 1, 2, 3, 4, 5, 7, and 8). The type 6 genotype of this parasite seems to form a distinct phylogenetic group from the family of benign *T. sergenti/buffeli/orientalis* group and has been previously detected in China, Thailand, and the United States [4]. In contrast, previous phylogenetic analysis of the *p23* gene indicated the presence of only 3-district genotypes, consisting of Buffeli, Chitose, and Ikeda types [9]. All 3 genotypes were successfully isolated in this study. Although the *p23* gene exhibited a relatively lower diversity than the *MPSP* gene, a greater phylogenetic complexity may become apparent when more detailed information on *p23* gene is accumulated from the field in the future. How and why such high diversities of *MPSP* and *p23* genes occur among field isolates of *T. orientalis* remain unclear and further epidemiological studies are required to understand the molecular diversity and wide distribution of *T. orientalis*.

Infection rates detected during our study do not reflect all farms in Kumamoto and Okinawa prefectures as we focused on farms that were suspected to be prevalent for *T. orientalis*. However, it is clear that *T. orientalis* is distributed throughout these prefectures, with a total of 5 *MPSP* genotypes identified, taking into account the results of our previous study [7]. Furthermore, with the aid of cloned plasmids, we successfully constructed 5 genotype-specific PCR methods that have the potential as diagnostic assays for the identification of actual infection rates for each genotype in each population. The results of the genotype-specific PCR analyses demonstrate the original distributions of 5 district genotypes of *T. orientalis* in the examined farms. Interestingly, dominant distributions of 1 or 2 genotypes were observed in the Kuma and Iriomote farms, while all genotypes of *T. orientalis* were widely distributed in the Aso, Ishigaki, and Yonaguni farms. Almost all Japanese Black cattle were infected with multiple parasite genotypes.

Terada et al. [15] reported that the indigenous Japanese Black breed of cattle was more resistant to *T. orientalis* infection than the exotic Holstein. Our previous report [7] also showed that the Japanese Black cattle showed no sign of anemia, although the exotic Holsteins and Herefords showed a high rate of anemia in the grazing field. Additionally, we recently reported significantly lower immunological responses of the Japanese Black cattle against *T. orientalis* infection, as compared to Holsteins [16]. Such unique characteristics of the Japanese Black cattle are supported by this breed in Iriomote Island giving a negative reaction in the serodiagnostic ELISA. The possibility that Japanese Black cattle resistant to *T. orientalis* might become
infectious carriers of multiple genotypes of *T. orientalis* is also highly interesting, but further epidemiological survey using the genotype-specific PCR assays is required to prove this.

This study indicated that at least 5 types of *T. orientalis* exist in Kumamoto and Okinawa prefectures of Japan and that genotype-specific PCR assays have useful applications in the quarantine of transported cattle and epidemiological surveys of bovine theileriosis in Japan.

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