

Comparative Genome Analysis of Three Eukaryotic Parasites with Differing Abilities To Transform Leukocytes Reveals Key Mediators of *Theileria*-Induced Leukocyte Transformation

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ABSTRACT We sequenced the genome of *Theileria orientalis*, a tick-borne apicomplexan protozoan parasite of cattle. The focus of this study was a comparative genome analysis of *T. orientalis* relative to other highly pathogenic *Theileria* species, *T. parva* and *T. annulata*. *T. parva* and *T. annulata* induce transformation of infected cells of lymphocyte or macrophage/monocyte lineages; in contrast, *T. orientalis* does not induce uncontrolled proliferation of infected leukocytes and multiplies predominantly within infected erythrocytes. While synteny across homologous chromosomes of the three *Theileria* species was found to be well conserved overall, subtelomeric structures were found to differ substantially, as *T. orientalis* lacks the large tandemly arrayed subtelomere-encoded variable secreted protein-encoding gene family. Moreover, expansion of particular gene families by gene duplication was found in the genomes of the two transforming *Theileria* species, most notably, the TashAT/TpHN and Tar/Tpr gene families. Gene families that are present only in *T. parva* and *T. annulata* and not in *T. orientalis*, *Babesia bovis*, or *Plasmodium* were also identified. Identification of differences between the genome sequences of *Theileria* species with different abilities to transform and immortalize bovine leukocytes will provide insight into proteins and mechanisms that have evolved to induce and regulate this process. The *T. orientalis* genome database is available at <http://totdb.czc.hokudai.ac.jp/>.

IMPORTANCE Cancer-like growth of leukocytes infected with malignant *Theileria* parasites is a unique cellular event, as it involves the transformation and immortalization of one eukaryotic cell by another. In this study, we sequenced the whole genome of a nontransforming *Theileria* species, *Theileria orientalis*, and compared it to the published sequences representative of two malignant, transforming species, *T. parva* and *T. annulata*. The genome-wide comparison of these parasite species highlights significant genetic diversity that may be associated with evolution of the mechanism(s) deployed by an intracellular eukaryotic parasite to transform its host cell.

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This paper is dedicated to the memory of Junichi Watanabe, who contributed to the full-length cDNA analysis and annotation performed as part of this study.

Theileria spp. are tick-borne intracellular parasites that belong to the phylum Apicomplexa and infect domestic and wild ruminants, including cattle, Asian water buffalos, sheep, goats, and African buffalos. Although infection by some *Theileria* species is asymptomatic or persists as a chronic infection, *Theileria parva* and *Theileria annulata* can be highly pathogenic to cattle and *Theileria lestoquardi* can cause significant disease in sheep. These three species are among the “transforming *Theileria*” species because of their ability to transform and induce indefinite proliferation of infected host leukocytes (1–4). The resulting disease syndromes can be described as lymphoproliferative disorders, which often culminate in disorganization and destruction of the host lymphoid system. Although detailed information has been generated for a number of host cell signal transduction pathways that are perturbed during leukocyte transformation, parasite molecules responsible for the initiation or regulation of the host cell transformation event have yet to be identified or fully validated (5, 6).

A comparative analysis of the *T. parva* and *T. annulata* genome sequences was reported in 2005 (7, 8). Despite the identification of a number of *Theileria* genes that could be involved in the transformation process, the selectivity of the approach was compromised by a high number of hypothetical proteins of unknown function and the high number of shared genes that exists across the genomes of these two closely related species. One way in which the discriminatory power of a comparative genomic approach could be increased would be to conduct bi- and trilateral genome comparisons with *Theileria* and *Babesia* parasites that lack the ability to transform host leukocytes but otherwise show strong similarity over the rest of their parasitic life cycle (9).

Theileria orientalis, an intraerythrocytic parasite of cattle, is a member of the nontransforming group of *Theileria* species that proliferate in the bovine host as an intraerythrocytic form and can generate anemia and icterus but rarely cause fatal disease (10). This parasite has frequently been referred to as *T. sergenti*, but this specific name is now considered invalid (11). Bovine piroplasmiasis caused by this species causes enormous economic losses in the livestock industry in Japan (12–14). *T. orientalis* is often classified into two major genotypes, the Chitose type and the Ikeda type, which are distinguishable on the basis of diversity in the small-subunit rRNA and major piroplasm surface protein (MPSP) gene sequences (15). The *T. orientalis* Ikeda type is limited to eastern Asian countries, including Japan, South Korea, the northeastern part of China, and Australia (16), and it is present in areas where livestock succumb to severe clinical cases of theileriosis and serious production losses. In contrast, *T. orientalis* Chitose is found throughout the world and is usually associated with benign infection (15, 17). Thus, even though it is believed to be relatively mild compared to the transforming *Theileria* species, *T. orientalis* can be an important pathogen in its own right and many researchers have been looking forward to the derivation of the genomic sequence to provide an important resource for further studies.

Unlike transforming *Theileria* species, the macroschizonts of nontransforming *Theileria* parasites are only transiently found in cells within lymph nodes or the spleen following the invasion of host cells by the infective sporozoite, and no evidence for proliferation of infected cells has been reported *in vivo* or *in vitro*. Indeed, *in vivo* studies indicate that the schizont undergoes continual enlargement over the course of 4 to 8 days before generating multiple merozoites that are released upon host cell destruction. A

lack of host proliferation is indicated by a substantial increase in host cell size, but it is unknown whether the parasite manipulates the cell at the molecular level or inhibits an apoptotic response to infection (13, 18). Free merozoites subsequently invade erythrocytes and, unlike the case with transforming species, undergo significant rounds of proliferation in red blood cells, similar to the proliferation observed with *Babesia* parasites. Clinical signs, when observed, are associated primarily with anemia and icterus. In addition to the schizont stage, the intraerythrocytic stage of *T. annulata* can also cause anemia.

In this study, we focused primarily on a comparative analysis of the genome of the *T. orientalis* Ikeda type relative to the genomes of the transforming *Theileria* species *T. parva* and *T. annulata* and a closely related hemoparasite species, *Babesia bovis*. The main goals of this analysis were to provide supportive data on existing candidate genes and/or identify novel candidate genes that enable the transformation of bovine leukocytes upon infection with *T. annulata* and *T. parva*.

RESULTS AND DISCUSSION

Structure of the *T. orientalis* genome. Whole-genome shotgun sequence data on *T. orientalis* (Ikeda strain) were assembled, and physical gaps between scaffolds were manually closed, resulting in the complete sequence of all four chromosomes. The derived sequence has been deposited in the DNA Data Bank of Japan (DDBJ) under project accession numbers AP011946 to AP011951. In addition to the nuclear genome, partial sequences of the apicoplast and mitochondrial genomes were also obtained. The complete genome sequence of the mitochondria has already been published (accession number AB499090) (19).

At 9.0 Mb, the genome size of *T. orientalis* is approximately 8% larger than the reported genome sizes of *T. parva*, *T. annulata*, and *B. bovis*. The number of predicted protein-coding genes identified in *T. orientalis* is, however, almost the same as that found in *T. parva* (Table 1). The G+C composition of the *T. orientalis* genome (41.6%) is higher than those of *T. parva* and *T. annulata* (34.1% and 32.5%, respectively) but similar to that of *B. bovis* (41.8%). The frequencies of the top 50 InterPro entries (see Table S1 in the supplemental material) are similar for the three *Theileria* species, suggesting that, in general, the three parasite species possess similar sets of gene families and encoded protein domains. For example, the InterPro domain of DUF529, known as the FAINT (frequently associated in *Theileria*) domain, described later in detail, is found frequently in all of the *Theileria* species sequenced to date. In contrast, the PEST motif, associated with rapid degradation of (nuclear) proteins, was found to be encoded by several gene families in the genomes of the two transforming *Theileria* species but was not identified in *T. orientalis*.

Synteny across all of the chromosomes of all three *Theileria* species is generally conserved, except for the subtelomeric regions, and several internal inversions were identified for each chromosome (Fig. 1). Most large-scale inversions were found when comparing *T. orientalis* versus *T. annulata* (Fig. 1, lower half of each panel) and were not present in the *T. annulata*-versus-*T. parva* comparison (Fig. 1, upper half of each panel), suggesting that these structural changes occurred following the speciation of *T. orientalis* and a common ancestor of *T. annulata*/*T. parva*. However, a large inversion of approximately 113,000 bp in chromosome 3 of *T. annulata* may have occurred after the speciation of *T. annulata* and *T. parva* (Fig. 1, upper right panel, indicated by a

TABLE 1 Comparison of genome characteristics of *T. orientalis*, *T. parva*, *T. annulata*, and *B. bovis*

Nuclear genome feature	<i>T. orientalis</i>	<i>T. annulata</i>	<i>T. parva</i>	<i>B. bovis</i>
Size (Mbp)	9.0	8.4	8.3	8.2
No. of chromosomes	4	4	4	4
Total G+C content (%)	41.6	32.5	34.1	41.8
No. of protein-coding genes	3,995	3,792	4,035	3,641
% of genes with introns	78.3	70.6	73.6	61.5
Mean gene length (bp)	1,861	1,606	1,407	1,514
% Coding	68.6	72.8	68.4	70.2
Mean intergenic length (bp)	390	396	402	589
% G+C composition of exons	44.5	37.6	35.9	44.0
% G+C composition of intergenic regions	35.2	22.5	24.9	37.0
% G+C composition of introns	38.1	22.2	23.6	35.9
No. of tRNA genes	47	47	47	44
No. of 5S rRNA genes	3	3	3	NA ^b
No. of 5.8S, 18S, and 28S rRNA units	2	2	2	3
Mitochondrial genome size (kb)	2.5	6	6	6
Apicoplast genome size (kb)	26.5	NA	39.5	33
Gene density ^a	2,249	2,202	2,059	2,228

^a Genome size/number of protein-coding genes.

^b NA, not available.

double-headed arrow). A striking difference between the genomes is that a number of gene families show evidence of expansion and diversification specific to the genomes of the transforming *Theileria* species, while few instances of *T. orientalis*-specific gene family expansion were recorded. In addition, several lineage-specific genes were identified at microsynteny breakpoints. Finally, the subtelomeric regions of all four *T. orientalis* chromosomes are markedly different from those of *T. annulata* and *T. parva* because they completely lack the largest subtelomeric gene family reported for *T. annulata* and *T. parva*, which encodes subtelomere-encoded variable secreted proteins (SVSPs) (see Fig. 2) of unknown function (20).

Metabolic pathways. To reconstruct KEGG metabolic pathways of *T. orientalis*, we assigned 263, 263, 273, and 264 KEGG orthology (KO) identifiers (21) to the predicted proteomes of *T. orientalis*, *T. parva*, *T. annulata*, *T. orientalis*, and *B. bovis*, respectively (see Fig. S1 in the supplemental material). These four species had 255 KOs in common, indicating no significant differences in known metabolic pathways between nontransforming and transforming *Theileria* and *Babesia* species, despite the known preference to proliferate in different host cell types (leukocytes versus erythrocytes).

K00626 is the only KO common to *B. bovis* and *T. orientalis* and not identified in *T. parva* and *T. annulata*. It codes for a putative thiolase that catalyzes the conversion of acetyl coenzyme A (acetyl-CoA) into acetoacetyl-CoA. This enzyme is known to function in a variety of metabolic pathways, including fatty acid metabolism, nucleotide metabolism, and amino acid degradation. Phylogenetic analysis has indicated that the two transforming *Theileria* species diverged from *T. orientalis* after the speciation of *Theileria* and *Babesia* (22). Therefore, the acetoacetyl-CoA thiolase might represent an example of a reduction of metabolic capacity due to an increasing host cell metabolite-scavenging ability/dependence of *Theileria* species.

Gene families. Expansion of gene families specific to different *Theileria* species could offer a valuable insight into how these parasites have evolved and adapted to their different host environments, including the acquisition of leukocyte transformation capability. To examine the expansion processes of gene families in

the *Theileria* lineages in detail, we constructed gene families composed of sequences representing the three *Theileria* species, *B. bovis*, and two *Plasmodium* species (*Plasmodium falciparum* and *P. vivax*) on the basis of the ortholog clustering framework of OrthoMCL (23), as well as additional computational and manual curations. We assigned 3,419 orthologous groups in which at least one *Theileria* species was included. While 1,740 of these orthologous groups consisted of single-copy genes across all six species, 223 orthologous groups possessed *Theileria* paralogs (see Data set S1 in the supplemental material). We focused on several family groups in the *Theileria* lineage that showed evidence of marked expansion that could be associated with acquisition of the ability to generate the proliferating, transformed, infected leukocyte.

Expansion of gene families in the genomes of transforming *Theileria* species. Three gene families showed a striking association with the genomes of the two transforming *Theileria* species. PiroF0100022 (Tar/Tpr family), PiroF0100037 (SVSP family), and PiroF0100038 (TashAT/TpHN family) are all significantly expanded within or unique to the genomes of the host cell-transforming *Theileria* lineage and are composed of genes predicted to encode proteins possessing FAINT domains. The TashAT family of *T. annulata* contains 17 tandemly arrayed genes, some of which have been shown to encode proteins that are translocated to the host nucleus, bind DNA, and alter gene expression and protein profiles of transfected bovine cells (24, 25). An orthologous cluster of 20 genes (TpHN) has also been identified in *T. parva* (25). In sharp contrast, only a single TashAT/TpHN-like gene, TOT0100571, was identified in the genome of *T. orientalis*. Reciprocal best hits using BLASTP indicate that the *T. orientalis* gene is likely to be the ortholog of Tash-a (TA03110) and TP01_0621 in the transforming *Theileria* species. Both of these genes are located at the 3' ends of their respective clusters in the *T. annulata* and *T. parva* genomes (Fig. 3A).

To gain further insight into the species-conserved Tash-a gene relative to the other members of the TashAT cluster, we obtained microarray data to examine whether gene expression of the different TashAT genes is associated with proliferating, macroschizont-infected leukocytes (26). Analysis of the normalized dataset showed that, in general, TashAT family expression is consistently

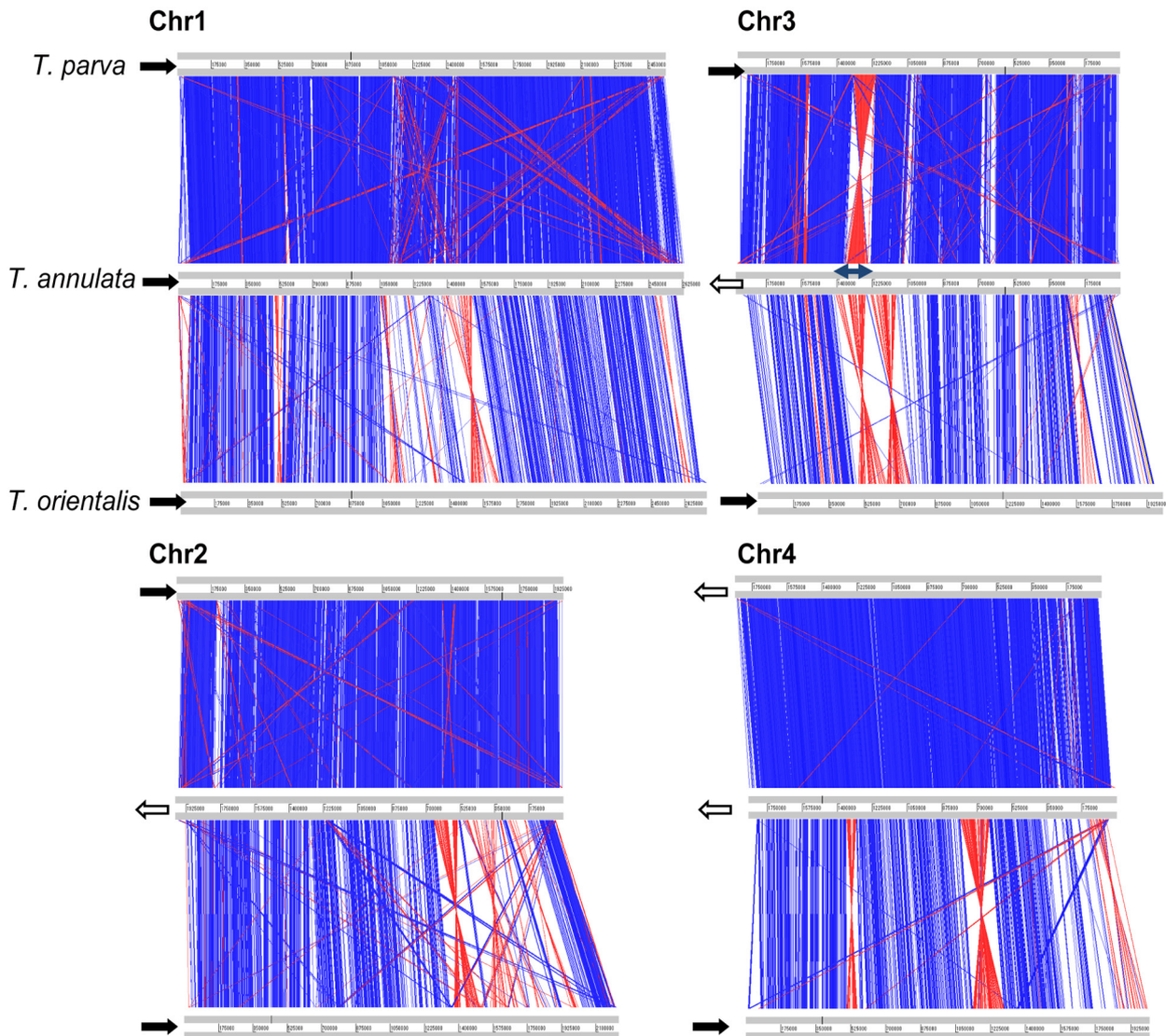


FIG 1 Genome scale synteny among three species of *Theileria* chromosomes. Shown is an Artemis Comparison Tool (65) plot of *T. orientalis* (bottom) versus *T. annulata* (middle) and *T. parva* (top). Blue bars indicate matching regions in the same orientation, while red bars indicate inverted matching. The direction of the chromosome is shown by arrows. Chromosome 3 (Chr3) of *T. parva* has a large gap due to the complexity of the Tpr locus; two contigs (AAGK01000005 and AAGK01000006) were connected with gaps.

downregulated as the macroschizont undergoes differentiation to the merozoite and host cell proliferation subsides, as demonstrated previously for a number of individual family members (25). In marked contrast, transcripts representing Tash-a were found to be significantly upregulated during the differentiation process (see Fig. S2A in the supplemental material). This result may indicate a requirement for synthesis of the protein during merozoite production. This postulation was supported by an indirect fluorescent-antibody test (IFAT) using serum raised against a Tash-a fusion protein (see Fig. S2B) and colocalization of Tash-a staining with a merozoite rhoptry antigen (see Fig. S2C). We conclude that the Tash-a protein performs a function that is required during or following merozoite production and that the temporal expression and location of the protein are distinct from those of other members of the family. Phylogenetic analysis suggests that Tash-a and its orthologs represent ancestral members of the

TashAT and TphN clusters (see Fig. 3B). In addition, we did not find any obvious TashAT orthologs in *B. bovis* or two *Plasmodium* species genomes. We propose that Tash-a diverged after the separation of *Theileria* from a common ancestor of *Theileria* and *Babesia* and that gene duplication and functional diversification of the TashAT and TphN clusters has then occurred as *Theileria* species of the transforming lineage evolved. Whether expansion of the cluster coincided with acquisition of a transforming capability is unknown.

Polypeptides encoded by the subtelomeric SVSP gene family (PiroF010037) are a major component of the predicted macroschizont secretome of *T. annulata* and *T. parva*, and a number of SVSPs have been predicted to translocate to the nucleus of the infected cell. Most SVSP genes are coexpressed in cultures of macroschizont-infected cells, and the SVSP family shows a high level of amino acid sequence diversity (20). Further work is

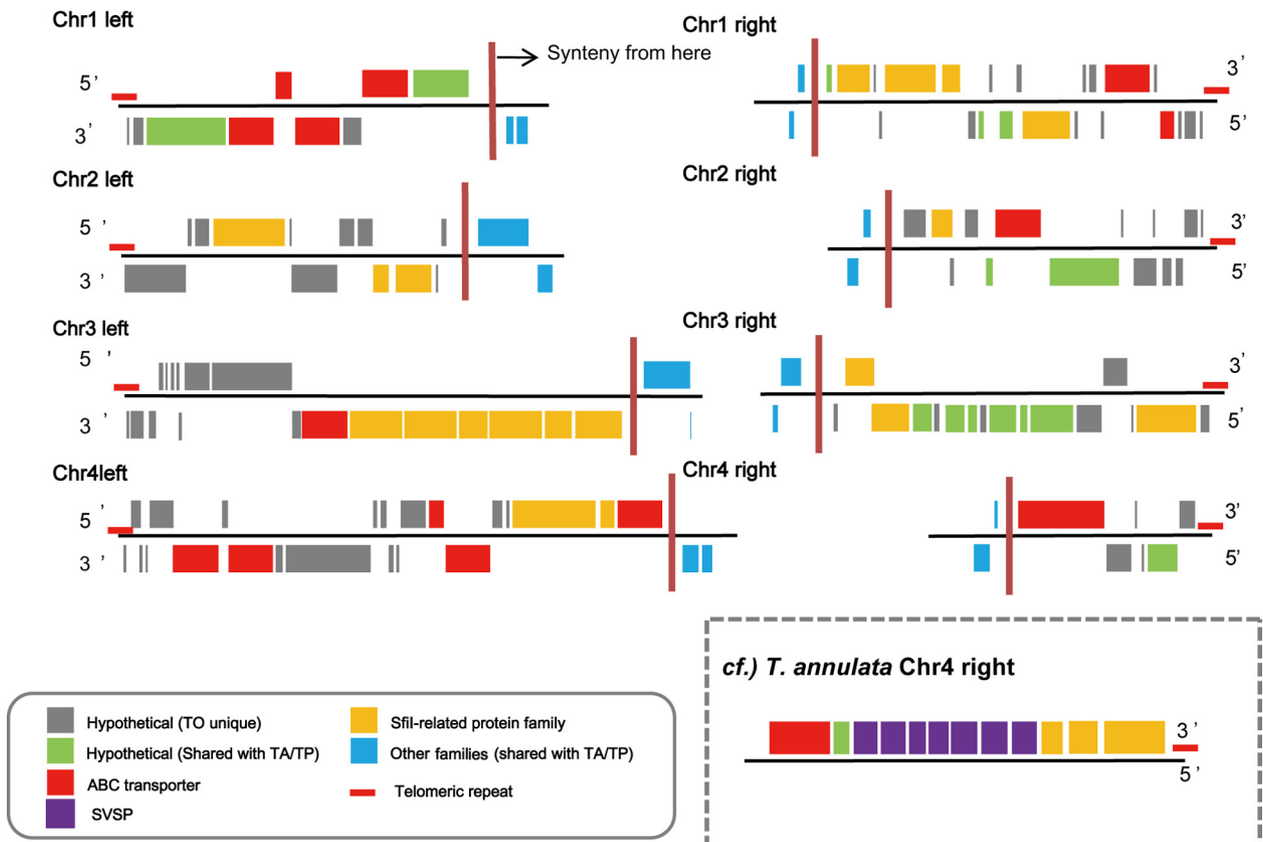


FIG 2 Genes in the subtelomeric region of each chromosome. Annotation of the subtelomeric regions of *T. orientalis* chromosomes identified mainly ABC transporter family or SfiI-related family genes but not SVSP genes, while subtelomeric regions of *T. annulata* or *T. parva* chromosomes are characterized by tandem-arrayed SVSP genes, SfiI-related family genes, and ABC transporter family genes. For comparison, the structure of chromosome 4 (Chr4) in *T. annulata* is shown within the dotted box.

needed to determine the function of SVSPs, whether they contribute directly to the transformation of the host cell or play a role in subverting the bovine immune response. Some of the SVSPs contain bioinformatically detectable signal peptides, suggesting secretion into the host cell cytoplasm. Though the expression patterns of *T. parva* SVSPs appear complicated and their involvement in phenotypic changes in host leukocytes remains unclear, the fact that some SVSPs encode functional nuclear localization signals (NLSs) in addition to a predicted signal sequence for secretion suggests that they might be transported to the host nucleus and modulate signaling pathways (20). In this context, the absence of SVSP loci in *T. orientalis* is noteworthy. Thus, like the TashAT/TpHN clusters, SVSP gene expansion in *T. annulata*/*T. parva* appears to be associated with species of the transforming *Theileria* lineage and may provide an as-yet-unknown function that promotes the establishment or maintenance of proliferating macroschizont-infected leukocytes.

In addition to the SVSP and TashAT clusters, the Tar/Tpr (PiroF0100022) family of orthologous genes showed evidence of significant expansion in the transforming *Theileria* lineages, as only five genes dispersed over the four chromosomes were detected in *T. orientalis*, compared with the 69 dispersed Tar genes in *T. annulata*. The function of the proteins encoded by Tar/Tpr genes is unknown. They lack a FAIN domain, and the presence of multiple transmembrane domains predicts a membrane location.

Transcriptome studies indicate that copies of Tpr genes dispersed throughout the *T. parva* genome are expressed in the macroschizont stage (27), while those organized in a tandem array of 28 genes are expressed by the intraerythrocytic piroplasm (28).

The CD8 T cell response is considered to play a key role in immunity to *T. parva*/*T. annulata* (29). Of the macroschizont antigens that are recognized by CD8 T cells from immune animals (30, 31), one, TA9/TP9 (TA15705/TP02_0895), is encoded by a member of a small orthologous gene family (PiroF0100041) in the genomes of transforming *Theileria* species. The family consists of five and six members in *T. annulata* and *T. parva*, respectively, all of which encode predicted proteins with a signal peptide for secretion by the parasite. Expressed sequence tag (EST) data and microarray data indicate that one of the TA9 family members (TA15705) is expressed in a specific manner by the transforming macroschizont stage (see Fig. S3C in the supplemental material), and it has been reported that the protein can be detected in the host cell cytosol (32). In the *T. orientalis* genome, a single gene (TOT020000921) showing weak homology in the signal peptide region and C-terminal region with the TA9/TP9 family was found in a syntenic region of chromosome 2 (see Fig. S3). The data indicate that the TA9/TP9 gene family has expanded uniquely in the transforming *Theileria* species. A role for TA9-encoded polypeptides in the transformation of the host cell requires further investigation.

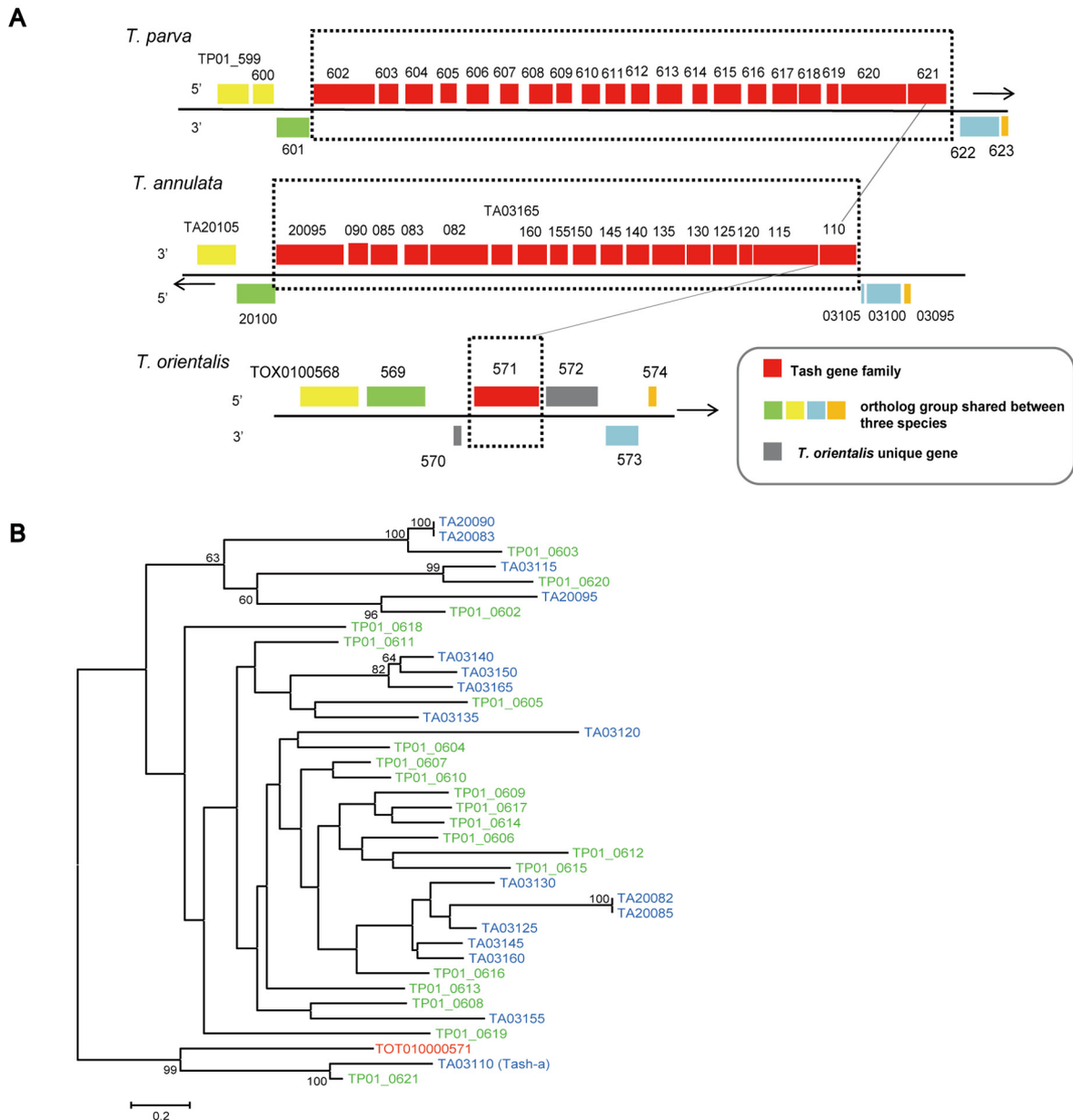


FIG 3 Genomic and phylogenetic structures of the TashAT gene family (PiroF0100038). (A) Schematic representation of the TashAT clusters in *T. parva* and *T. annulata* and the corresponding locus in *T. orientalis*. Genes in the same orthologous group are represented by the same color. Bars indicate direct orthologous gene pairs as inferred by phylogenetic analysis. (B) Phylogenetic trees of the TashAT/TpHN (PiroF0100038) family. Proteins representative of *T. orientalis*, *T. annulata*, and *T. parva* are indicated in red, blue, and green, respectively. Bootstrap percentage values (>60) are shown at the nodes.

Evolution of the FAINT domain superfamily. As observed for *T. annulata* (8) and *T. parva* (7), a large number of genes whose predicted polypeptides encode DUF529 domains (IPR007480 in InterPro), alternatively called FAINT domains, were found in *T. orientalis* (see Table S1 in the supplemental material). Previous analysis revealed that ~900 copies of FAINT domains are present in the genomes of *T. annulata* and *T. parva* (8). With our pipelines for InterPro annotation, 686 FAINT copies were identified in 137 predicted *T. orientalis* proteins, and 913 and 725 copies were identified in 126 *T. annulata* and 142 *T. parva* putative proteins, respectively. This suggests that expansion of FAINT domain-containing polypeptides (FAINT superfamily) is likely to have

occurred in the common ancestor of the three *Theileria* species. In addition, ortholog clustering indicated that different FAINT families have been expanded in *T. orientalis* than in *T. parva* and *T. annulata*. For example, the FAINT superfamilies of PiroF0001942 and PiroF0001943 are specifically expanded in *T. orientalis* (see Table S2 in the supplemental material). In contrast, the PiroF0100056 orthologous group of SfiI-related genes showed greater expansion in *T. parva* and *T. annulata* (see Table S2). A protein of the FAINT superfamily was also found in *T. equi* (8), which has been considered to be an outlier species in the genus *Theileria* (33). This indicates that FAINT domain polypeptides were present in early ancestral species of the *Theileria* genus and

have subsequently been subjected to differential expansion or contraction pressures as the different species evolved.

Many of the FAINT superfamily members in *T. parva* and *T. annulata* are inferred to be secretory proteins (5). Out of 137 proteins of the FAINT superfamily identified in *T. orientalis*, signal peptides were found in 103, indicating that members of the FAINT superfamily are significantly enriched for proteins with a predicted signal peptide ($P = 5.97 \times 10^{-55}$, Fisher's exact test). Thus, the differential expansion and diversification of FAINT domain proteins could be associated with the adaptation of different *Theileria* species to preferential host niches that require specific host-parasite interactions. Comparison of additional genome sequences derived from both nontransforming and transforming *Theileria* species may be informative.

Candidate genes responsible for *Theileria*-induced host cell transformation. Comparative genomic analysis of *T. orientalis* and *T. annulata*/*T. parva* provides a tool for identifying candidate genes responsible for *Theileria*-mediated host cell transformation. This premise is based on the assumption that transformation-related genes are unique to the *T. annulata*/*T. parva* lineage, as there is no evidence that *T. orientalis* can transform leukocytes into proliferating infected cells. It can also be predicted that molecules that regulate the transformation event are likely to be secreted or localized to the macroschizont membrane, since *Theileria* parasites have direct contact with the host cell cytoplasm (34). In the course of ortholog classification analysis, we applied both of these criteria and identified 97 ortholog groups present in the *T. parva* and *T. annulata* lineages that were absent from *T. orientalis*, *B. bovis*, *P. falciparum*, and *P. vivax*. Of these lineage-specific ortholog groups, 29 are predicted to encode polypeptides with an endoplasmic reticulum signal sequence (several of which also contain a GPI anchor motif), indicating potential interaction with the host cell compartment (Table 2). The majority of these genes encode hypothetical proteins and do not show any similarities to known cancer-related genes, although several domains are predicted in the InterPro entries. We propose that genes placed within these 29 groups, plus the TashAT/TpHN family, can be considered candidates for involvement in the transformation process.

Identification of candidate genes as host cell phenotype manipulators has been reported previously (5, 35). The predicted proteins have signal sequences, protein kinase properties, phosphatase properties, NLSs, or DNA binding motifs, or they show identity with higher eukaryotic proteins that are involved in neoplasia. We searched for these genes in the genome of *T. orientalis* and found that all of them, with the exception of TashAT and SVSP family genes, are conserved across the three *Theileria* species (see Table S3 in the supplemental material). However, four *T. orientalis* genes lack the signal sequence or NLS that is predicted in each of the *T. annulata*/*T. parva* orthologs. Thus, it is possible that the function or localization of the encoded polypeptides has diverged between *T. orientalis* and the transforming *Theileria* species, and this may be worthy of further investigation.

Conclusions. This is the first genome sequence of a nontransforming *Theileria* species that occupies a phylogenetic position close to that of the transforming *Theileria* species and thus provides an ideal opportunity to analyze unique features of *Theileria* parasitism from an evolutionary viewpoint. Genome sequencing of the nontransforming *Theileria* species *T. orientalis* and comparison with the transforming *Theileria* species *T. annulata* and

T. parva highlighted lineage-specific evolutionary features. Several transforming *Theileria* lineage-specific gene family expansions were identified, including the SVSP, Tash/TpHN, Tpr/Tar, and TP9/TA9 families, that may have been coincident with development of the ability to transform host leukocytes. Additional genes identified as specific to the genomes of transforming *Theileria* species can also be considered transformation candidates. This study provides increased understanding of the evolution of transforming *Theileria* species at the genomic level and has generated a database that will serve as the foundation for future studies on *Theileria* pathobiology and parasite-host cell interaction.

MATERIALS AND METHODS

Parasite samples. *T. orientalis* (Shintoku stock) was used as the starting genomic material in this study. This stock contains two different genotypes, Ikeda and Chitose. Parasites of a single genotype (Ikeda) were selected following syringe passage of the original isolate through calves and then used to infect an animal for parasite isolation. Blood collected from the infected animal was passed through a leukocyte removal filter (Terumo), and the resulting red cells were washed three times with phosphate-buffered saline (PBS). Erythrocytes were resuspended in an equal volume of PBS and disrupted by nitrogen cavitation, and piroplasmids were purified by differential centrifugation as described previously (36). Infection of the cow was conducted in accordance with protocols approved by the National Institute of Animal Health, Japan, Animal Care and Use Committee (approval no. 2000/901). Genomic DNA was purified by proteinase K and SDS treatment, followed by phenol-chloroform extraction. Purified parasite DNA was dissolved in TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0). Confirmation that the DNA represented the Ikeda genotype was carried out by PCR targeting genes encoding small-subunit rRNA and the MPSP as previously described (37).

Genome sequencing. The complete genome sequence of *T. orientalis* was determined by a combination of the whole-genome shotgun method and fosmid end sequencing. Genomic DNA was fragmented for plasmid library construction with an average insert size of 2 to 4 kb using a HydroShear DNA Shearing Device (Genemachines). Plasmid DNA was amplified with a TempliPhi DNA amplification kit (GE Healthcare) from the bacterial culture. The fosmid library was constructed by TaKaRa Bio Inc. using a CopyControl pCC1FOS vector (Epicentre, Madison, WI). Fosmid DNA was extracted with PI-1100 plasmid isolators (Kurabo). Both ends of 40,704 plasmid inserts and 3,840 fosmid clones were sequenced with ABI 3730 sequencers (Applied Biosystems) and MegaBACE 4500 sequencers (GE Healthcare). Contigs were assembled by using 111,945 shotgun reads. Gap closing and resequencing of low-quality regions in the assembled data were performed by shotgun sequencing of fosmid clones that covered the target regions, nested deletion (38), construction of short-insert libraries (39), and primer walking on selected clones and PCR-amplified DNA fragments. The overall accuracy of the finished genome sequence was estimated to have an error rate of less than 1 per 10,000 bases. The sequence is available from DDBJ/GenBank/EMBL under accession numbers AP011946 to AP011951.

cDNA/ESTs. Six volumes of Trizol LS was added to 1 volume of parasite-infected erythrocytes and homogenized with a Polytron homogenizer. Total RNA was then isolated according to the manufacturer's protocol, and full-length cDNA libraries were produced by either the oligocapping or the vector-capping method (40). Random clones were picked from the oligocapped and vector-capped library, and inserts were amplified by PCR from the single colonies sequenced at the 5' end or both the 5' end and the 3' end. Sequences were aligned with available whole-genome sequences by using the est2genome (41) program. These sequences are available from DDBJ/GenBank/EMBL under accession numbers FS557591 to FS578553.

Gene structure prediction and annotation. All of the repetitive and low-complexity sequences in the *T. orientalis* genome sequence were masked by using RepeatMasker (<http://www.repeatmasker.org>) with

TABLE 2 Possible candidate transforming genes in *T. parva* and *T. annulata*

Gene family	Product ^a	TA ID	TP ID	Signal ^b	TMD ^c	GPI ^b
PiroF0100038	TashAT family	(TA03110), TA03115, TA03120, TA03125, TA03130, TA03135, TA03140, TA03145, TA03150, TA03155, TA03160, TA03165, TA20082, TA20083, TA20085, TA20090, TA20095	TP01_0602, TP01_0603, TP01_0604, TP01_0605, TP01_0606, TP01_0607, TP01_0608, TP01_0609, TP01_0610, TP01_0611, TP01_0612, TP01_0613, TP01_0614, TP01_0615, TP01_0616, TP01_0617, TP01_0618, TP01_0619, TP01_0620, (TP01_0621)			
PiroF0100041	Hypothetical protein (TA9/TP9 family)	TA15685, TA15705 (TA9), TA15710, TA15690	TP02_0890, TP02_0895, TP02_0896, TP02_0891, TP02_0894	Y (TA15705)	0	N
PiroF0100037	<i>Theileria</i> -specific subtelomeric protein, SVSP family	TA02740, TA04895, TA05540, TA05545, TA05550, TA05555, TA05560, TA05565, TA05570, TA05575, TA05580, TA09420, TA09425, TA09430, TA09435, TA09785, TA09790, TA09795, TA09800, TA09805, TA09810, TA09865, TA11385, TA11390, TA11395, TA11410, TA16025, TA16030, TA16035, TA16040, TA16045, TA17120, TA17125, TA17130, TA17135, TA17140, TA17346, TA17475, TA17480, TA17485, TA17535, TA17540, TA17545, TA17550, TA17555, TA18860, TA18865, TA18885, TA18890, TA18895, TA18950, TA19005, TA19060	TP01_0004, TP01_0005, TP01_0006, P01_0007, TP01_0008, TP01_0009, TP01_1225, TP01_1226, TP01_1227, TP02_0004, TP02_0005, TP02_0006, TP02_0007, TP02_0008, TP02_0010, TP02_0011, TP02_0953, TP02_0954, TP02_0955, TP02_0956, TP02_0958, TP02_0959, TP02_0960, TP03_0001, TP03_0002, TP03_0003, TP03_0004, TP03_0005, TP03_0498, TP03_0866, TP03_0867, TP03_0868, TP03_0869, TP03_0870, TP03_0871, TP03_0872, TP03_0873, TP03_0874, TP03_0875, TP03_0877, TP03_0878, TP03_0879, TP03_0880, TP03_0881, TP03_0882, TP03_0883, TP03_0884, TP03_0885, TP03_0886, TP03_0887, TP03_0888, TP03_0889, TP03_0890, TP03_0892, TP03_0893, TP03_0930, TP04_0001, TP04_0002, TP04_0003, TP04_0004, TP04_0005, TP04_0006, TP04_0007, TP04_0008, TP04_0009, TP04_0010, TP04_0013, TP04_0014, TP04_0015, TP04_0016, TP04_0017, TP04_0018, TP04_0019, TP04_0916, TP04_0917, TP04_0918, TP04_0919, TP04_0920, TP04_0923, TP04_0927	Y	0	N
PiroF0100039	<i>Theileria</i> -specific conserved protein	TA18755, TA18760, TA18765	TP03_0633, TP03_0634, TP03_0635, TP03_0636, TP03_0637, TP03_0638	Y	0	N
PiroF0003402	Hypothetical protein	TA20990	TP01_0378	Y	0	N
PiroF0003403	Hypothetical protein	TA20985	TP01_0379	Y	0	N
PiroF0003404	Proline-rich hypothetical protein	TA20980	TP01_0380	Y	0	N
PiroF0003405	Cysteine repeat modular protein homologue, putative	TA20781	TP01_0438	Y	0	N
PiroF0003407	Hypothetical protein	TA20615	TP01_0487	Y	1	N
PiroF0003411	Integral membrane protein, putative	TA20325	TP01_0549	Y	6	N
PiroF0003421	<i>Theileria</i> -specific hypothetical protein	TA18750	TP03_0632	Y	1	N
PiroF0003425	Hypothetical protein	TA18535	TP03_0582	Y	0	N
PiroF0003432	<i>Theileria</i> -specific hypothetical protein	TA17695	TP03_0678	Y	1	N
PiroF0003436	Hypothetical protein	TA17220	TP04_0030	Y	1	Y
PiroF0003437	Hypothetical protein	TA17215	TP04_0029	Y	1	N
PiroF0003438	Hypothetical protein	TA17210	TP04_0028	Y	0	Y
PiroF0003456	Hypothetical protein	TA16020	TP02_0952	Y	0	N
PiroF0003462	Hypothetical protein	TA15695	TP02_0888	Y	0	N
PiroF0003486	Hypothetical protein	TA13955	TP02_0065	Y	0	N
PiroF0003519	Hypothetical protein	TA11050	TP04_0896	Y	1	N

(Continued on following page)

TABLE 2 (Continued)

Gene family	Product ^a	TA ID	TP ID	Signal ^b	TMD ^c	GPI ^b
PiroF0003520	Hypothetical protein	TA11020	TP04_0585	Y	1	N
PiroF0003524	Hypothetical protein	TA10740	TP04_0642	Y	0	Y
PiroF0003546	Sfil subtelomeric fragment-related protein family member, putative	TA09140	TP04_0116	Y	0	N
PiroF0003548	Hypothetical protein	TA08935	TP04_0539	Y	2	N
PiroF0003567	Hypothetical protein	TA06680	TP01_0719	Y	0	N
PiroF0003568	Hypothetical protein	TA06675	TP01_0718	Y	1	N
PiroF0003582	Hypothetical protein	TA05315	TP03_0135, TP03_0134	Y	0	N
PiroF0003592	Hypothetical protein, conserved	TA04390	TP03_0410	Y	2	N
PiroF0003612	Hypothetical protein	TA02590	TP03_0038	Y	0	Y
PiroF0003613	Hypothetical protein	TA02580	TP03_0040	Y	0	N

^a *T. annulata* definitions.^b Y, yes; N, no.^c TMD, transmembrane domain.

Repbse. rRNA and tRNA genes were detected by using BLAST searches against Rfam (42) and the tRNAscan-SE program (43).

T. orientalis genes were first predicted computationally by using *T. orientalis* EST pair gene models and several gene prediction programs and then finally identified by genome-wide manual curation. *T. orientalis* EST sequences, identified from a full-length cDNA library made from parasite-infected erythrocytes, were mapped onto the *T. orientalis* genome. Based on EST-genome alignments using est2genome (41), EST pair gene models were constructed by merging the exon overlap on the same strand of ESTs of the same clone. We identified 544 *T. orientalis* EST pair gene models. Genes were predicted by several gene-finding software packages, including GlimmerHMM (44), GeneMark.hmm (45), GeneWise (46), and JIGSAW (47). GlimmerHMM was trained on two sets of full-length gene sequences. The first set consisted of *T. orientalis* genes (544 EST pair gene models), and the second set consisted of these *T. orientalis* genes and annotated genes of *T. parva* and *T. annulata* that were predicted to be longer than 400 amino acids. GeneWise was trained on all of the annotated genes of *T. parva* and *T. annulata*. The genes from four sets of results of genome coordinates provided by GlimmerHMM, GeneMark.hmm, GeneWise, and *T. orientalis* (544 EST pair gene models) were summarized by using JIGSAW. JIGSAW was also trained on *T. orientalis* EST pair gene models.

We essentially used annotation procedures described previously (48, 49). For each *T. orientalis* gene product, we conducted InterProScan (50). We then assigned a standardized functional annotation to each gene as illustrated in Fig. S4 in the supplemental material, based on the results of a BLASTX similarity search against the UniProtKB/Swiss-Prot, UniProtKB/TrEMBL, and RefSeq protein databases and InterProScan (48, 51). Finally, to identify the representative *T. orientalis* genes, manual curation was performed by using a custom-made annotation system named TOT-SOUP/G-integra (48, 52). The numbers of manually curated *T. orientalis* genes are summarized in Table S4 in the supplemental material. Signal peptides were inferred by SignalP 3.0 (53).

Ortholog clustering. Ortholog groups consisted of *T. orientalis*, *T. annulata*, *T. parva*, *B. bovis*, *P. falciparum*, and *P. vivax* proteins derived primarily from gene annotation. *T. annulata* orthologs were from GeneDB (<http://old.genedb.org/genedb/annulata/>); *T. parva*, except for the mitochondrion proteome, and *B. bovis* orthologs were from RefSeq (<http://www.ncbi.nlm.nih.gov/RefSeq/>); the *T. parva* mitochondrial proteome was from UniProt (<http://www.uniprot.org/>); and *P. falciparum* and *P. vivax* orthologs were obtained from PlasmoDB (<http://plasmodb.org/plasmo/>). Ortholog groups were generated by OrthoMCL (23) on the basis of sequence similarity by using an all-versus-all NCBI BLASTP search (54) with a bit score cutoff of <60 and default parameters. Because

E values from the BLASTP search were applied for a similarity measure, we recomputed the exact E values between closely related proteins if the E value was approximated at 0.0. We integrated the orthologous groups assumed to be duplicated in the *Theileria* lineage after separation from *Babesia* into a single group by using both automatic algorithms/software and manual integration as described below. Ortholog groups A and B were merged if any *Theileria-Theileria* gene pairs in which two genes belonging to A and B, respectively, had higher bit scores than any *Theileria-Babesia/Plasmodium* gene pairs within single ortholog group A or B. Several ortholog groups were merged by manual curation based on sequence homology and genomic location if they generated tandem arrays on the chromosomes. We also merged nonclustered genes using OrthoMCL into the ortholog groups with the same procedure. Finally, 3,502 ortholog groups were used for the following analyses; PiroF0100001 to PiroF0100062 represent the merged ortholog groups, and PiroF0000001 to PiroF0003675 represent the other ortholog groups. The ortholog clustering left 436, 112, and 293 nonclustered genes in *T. orientalis*, *T. annulata*, and *T. parva*, respectively.

KEGG metabolic pathway reconstruction. Metabolic pathways in *T. orientalis* were analyzed by KEGG metabolic pathway reconstruction. First, BLAST searches were performed for protein sequences in each orthologous cluster against the KEGG GENES database. A KO identifier was then assigned to each cluster according to the most similar hit with a KO annotation; the E value threshold was <1.0⁻⁵.

Molecular phylogenetic analysis. Amino acid sequences of each ortholog group were multiply aligned with the L-INS-I alignment strategy in MAFFT (55), and gap-rich sequences, such as truncated ones, were removed from the alignments with MaxAlign (56). Ambiguously and/or poorly aligned sites were removed by Gblocks (57), and the rest were subjected to phylogenetic analysis. Phylogenetic trees were inferred by maximum likelihood (ML) (58, 59) with a heuristic ML tree search using RAxML (60) with the WAG-F model (61). Heterogeneity of evolutionary rates among sites was modeled by a discrete gamma distribution, with optimization of gamma shape parameter alpha for each alignment set (62). Bootstrap probability (59) was calculated for each tree node with 1,000 replications.

Generation of recombinant protein and antiserum. A 1,788-bp fragment of TA03110 was PCR amplified with the C9 (genome) strain of *T. annulata* as a template. This corresponds to the full-length encoded protein minus the N-terminal signal peptide sequence and spans nucleotide positions 70 to 1,857 relative to the translation start codon. In addition to gene-specific sequences, the PCR primers incorporated *attB* adaptors to facilitate the use of Gateway Recombination Cloning Technology (Invitrogen); the forward primer was 5'-forward *attB* adaptor-

GAGGACTTGGACCTAAACTCTCC-3', and the reverse primer was 5'-reverse *attB* adaptor-AGGATTTTGATCAGTGTAAATATCG-3'. The amplicon was cloned into the pDONR221 shuttle vector and subcloned into the expression vector pDEST17, which has a six-histidine (His₆) repeat at the 5' end of the multiple cloning site. After the transformation of chemically competent *Escherichia coli* BL21 cells (Invitrogen), expression of the His₆-tagged fusion protein was induced by adding l-arabinose to a final concentration of 0.2% in LB liquid medium. Recombinant protein was purified by affinity chromatography on nickel agarose columns under denaturing conditions by using the manufacturer's protocol (Qiagen). Eluted fractions containing the recombinant protein were assessed by using SDS-PAGE before being pooled. To generate polyclonal anti-TA03110 serum, two rats were immunized a total of four times with 30 μg of recombinant protein per immunization. Immunizations were conducted under a project license issued by the United Kingdom Home Office, i.e., Animals (Scientific Procedures) Act 1986 contract immunization project license PPL 60/3464.

Parasite material and IFAT. The *T. annulata*-infected cloned cell line Ankara A₂ D7 (26) was used to provide material for the microarray experiment and for the IFAT. To stimulate differentiation from the macroschizont stage to the merozoite stage, cultures were maintained at 41°C by using a previously described protocol (26). Cytospin preparation of *T. annulata*-infected cells, paraformaldehyde fixation, and the IFAT were performed as described previously (62). The anti-Tash-a serum was used at dilution of 1:500 in cell culture medium, and the anti-His₆ tag antibody (sc-65902; Stratagene) was used at 1:200; monoclonal antibodies against a macroschizont surface antigen (1C12), the Tams1 merozoite surface antigen (5E1), and a merozoite rhoptry antigen (1D11) were used as undiluted hybridoma culture medium as previously described (63); anti-rat IgG and anti-mouse IgG secondary antibodies conjugated to Alexa 488 or Alexa 555 (Invitrogen) were used at a 1:200 dilution.

Microarray analyses. Parasite gene expression was investigated by using a custom-designed tiling microarray (Roche NimbleGen Inc., Madison, WI). Each gene in the TashAT cluster was represented by a set of 45-mer oligonucleotides that were specific to that gene. cDNA was generated from 10 μg total RNA by using an oligo(dT) primer and tagged with 3'-Cy3 dye, after which labeled cDNA was hybridized to the array. Gene expression values were calculated from a robust multiarray average-normalized dataset (64).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <http://mbo.asm.org/lookup/suppl/doi:10.1128/mBio.00204-12/-/DCSupplemental>.

- Data set S1, XLSX file, 0.4 MB.
- Figure S1, TIF file, 0.5 MB.
- Figure S2, TIF file, 5.6 MB.
- Figure S3, TIF file, 0.7 MB.
- Figure S4, TIF file, 0.4 MB.
- Table S1, PDF file, 0.1 MB.
- Table S2, PDF file, 0.01 MB.
- Table S3, PDF file, 0.01 MB.
- Table S4, PDF file, 0.01 MB.

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