



Complete Mitochondrial Genome of the Chagas Disease Vector, *Triatoma rubrofasciata*

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Abstract: *Triatoma rubrofasciata* is a wide-spread vector of Chagas disease in Americas. In this study, we completed the mitochondrial genome sequencing of *T. rubrofasciata*. The total length of *T. rubrofasciata* mitochondrial genome was 17,150 bp with the base composition of 40.4% A, 11.6% G, 29.4% T and 18.6% C. It included 13 protein-coding genes, 22 tRNA genes, 2 rRNA genes and one control region. We constructed a phylogenetic tree on the 13 protein-coding genes of *T. rubrofasciata* and other 13 closely related species to show their phylogenetic relationship. The determination of *T. rubrofasciata* mitogenome would play an important role in understanding the genetic diversity and evolution of triatomine bugs.

Key words: *Triatoma rubrofasciata*, mitochondrial genome, protein-coding gene, phylogeny

Chagas disease, also known as American trypanosomiasis, caused by the protozoan parasite, *Trypanosoma cruzi*, infects 8 million people worldwide, mostly in Latin America. In the past decades, it had been increasingly detected in the United States of America, Canada, and many European and some Western Pacific countries. This is mainly due to population mobility between Latin America and the rest of the world [1].

This disease is transmitted mainly through the biting of triatomine (reduviid) bugs, also called as 'kissing bug'. The sub-family Triatominae is divided into 149 species, that composed of 5 tribes: Alberproseniini, Bolboderini, Cavernicolini, Rhodniini and Triatomini, which is distributed from the southern USA to Patagonia, and a few species of Triatomini are in India and Australia [2]. So far, information of only 2 mitochondrial (mt) genomes of triatomine bugs has been published. One is *Triatoma dimidiata*, mainly distributed from Mexico to Northern South America [3], the other is *Triatoma infestans*, the main vector of Chagas disease in South America [4]. In China, information of triatomine bugs was far from complete. Only a few reports show that *T. rubrofasciata* and *Triatoma sinica* are the 2 main native triatomine bugs [5]. It was used to distribute in

Fujian, Guangxi, Hainan and Taiwan province in 1980s [6,7]. In 2017, *T. rubrofasciata* from Guangdong province had been characterized by 16S rRNA sequence [5]. However, complete information of its mt genome is still unknown.

For better understanding the biology of this species, the complete mt genome of *T. rubrofasciata* was determined and analyzed in this study. Furthermore, we constructed a phylogenetic tree on the 13 protein-coding genes of *T. rubrofasciata* and other 13 closely related species that show *Triatoma* species cluster together as a sister clade of Stenopodainae. Our findings will shed light on molecular taxonomy and population genetic studies of this important disease transmitting vector.

A single female adult of *T. rubrofasciata* collected from the rural area of Baoting County (18°39'N 109°41'E) Hainan Province was used. The sample was stored in 100% ethanol at -20°C. The total DNA was extracted using the Holmes-Bonner buffer [8] and stored at -20°C until use. The complete mt genome sequence of *T. rubrofasciata* was amplified in 5 overlapping fragments, using 5 pairs of oligonucleotide primers designed based on the conserved regions from published complete mt genome of *T. dimidiata* (NC_002609.1) (Table 1). All PCR amplification reactions were carried out in a final volume of 25 µl, which included 9.5 µl of H₂O, 12.5 µl of 1-5™ 2X High-Fidelity Master Mix (Molecular Cloning Laboratories MCLAB, South San Francisco, California, USA), 1 µl of each primer (10 µM), and 1 µl of DNA extract. Thermocycler conditions were as follows: 98°C for 2 min; 25 to 35 cycles of 98°C for 10 sec, 45-68°C for 15 sec,

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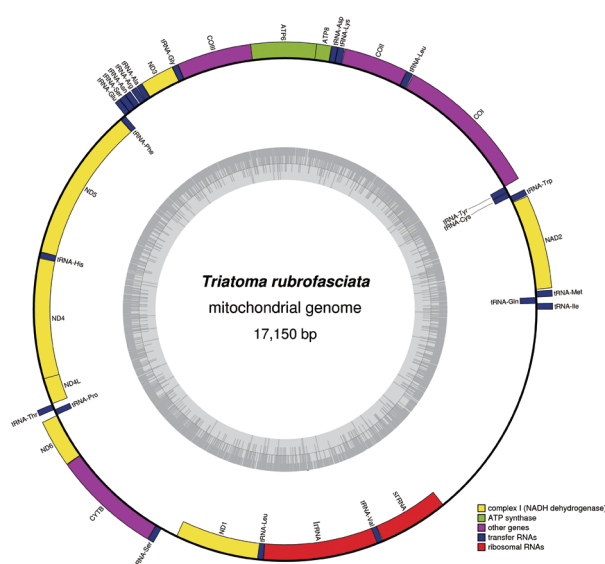
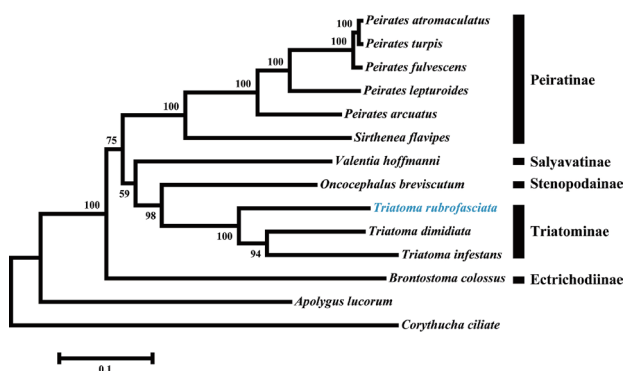
Table 1. Primer sequences used to amplify PCR fragments of *Triatoma rubrofasciata*

Primer name	Sequences (5'-3')	Size (kb)
Tr-1	ACCGCCTATTAATTCAGCCACTT GCGTGTTCCTAGTCGAAGACTTGTT	~7k
Tr-2	ACACCCGCGAGTAACCAAAGTAGAAG AGGATGTAAGGTTCTTCAGCAGGAC	~4k
Tr-3	ACCATCTCGTCCGTTATCTTCTTCT GCGGTTATACAATAGGAGCGAGTGA	~4k
Tr-4	AAACGAGAGTGACGGGCGATATG GTTATCCTGTTCTGCTCCTCTT	~4k
Tr-5	TGGAAATGATGTCTTGGTTGCTA TCAGAAAGACCTATGATCCTAAGAA	~2k

72°C for 30-90 sec depending on fragment lengths, followed by 5-15 s/kb, then a final extension at 72°C for 5 min. These amplicons were sequenced by Shanghai Tsingke (Shanghai, China) using primer walking in both directions.

The annotation of mt genome for *T. rubrofasciata* was performed using web-based services MITOS (<http://mitos.bioinf.uni-leipzig.de/help.py>) [9]. Protein-coding genes were annotated by checking manually for consistent start/stop codons, open reading frames. Then, tRNAs were identified by the proposed clover-leaf secondary structure and predicted using web-based tRNA-scan SE 1.21 program (<http://lowelab.ucsc.edu/tRNAscan-SE/>) [10] with default search mode. The rRNA genes were annotated by extending until adjacent tRNAs [11]. Comparing with other mt genomes, the 5' end of the srRNA gene was confined. Moreover, the base composition and codon usage were assessed using MEGA version 6.0 [12]. Finally, the graphical map of mt genome was drawn using web-based OrganellarGenomeDRAW (<http://ogdraw.mpimp-golm.mpg.de/>) [13] (Fig. 1).

Previously published mt genome under other Reduviidae species were obtained from GenBank, including *Brontostoma colosus* (KM044501), *Oncocephalus breviscutum* (KC887527), *Peirates arcuatus* (KF752445), *Peirates atomaculatus* (KF913538), *Peirates fulvescens* (KF913537), *Peirates lepturoides* (KF913541), *Peirates turpis* (KF913540), *Sirthenea flavipes* (HQ645959), *T. dimidiata* (AF301594), *T. infestans* (KY640305), and *Valentia hoffmanni* (FJ456952). Furthermore, the mt genome from *Apolygus lucorum* (HQ902161) and *Corythucha ciliate* (KC756280) were used as outgroup. Then, protein-coding genes were aligned using ClustalW in MEGA 6.0 [12] with default settings. Thus, the sequences from the 13 protein-coding genes alignments were concatenated to single multiple sequence alignment for phylogenetic analysis. The phylogenetic relationships were constructed us-

**Fig. 1.** Graphical map of the complete mitochondrial genome of *Triatoma rubrofasciata*. Genes encoded by the heavy strand were shown outside the circle, and genes encoded by the light strand were shown inside the circle, respectively. The GC content of the genome were shown in the inner circle.**Fig. 2.** Phylogenetic tree based on Maximum likelihood analysis of 13 protein-coding genes. Sequence from the present study was indicated with a blue font. *Apolygus lucorum* and *Corythucha ciliate* were used as outgroup. Bootstrap support values were displayed at each node.

ing maximum-likelihood method in MEGA 6.0, with node support was assessed with 1,000 bootstrap replicates (Fig. 2).

The complete mt genome of *T. rubrofasciata* was a typically circular molecule with 17,150 bp in length (GenBank accession no. MH934953). The size of *T. rubrofasciata* mt genome was smaller than *T. infestans* (17,301 bp) [4], whereas was larger than *T. dimidiata* (17,019 bp) [3]. The slight difference in length is mainly due to the repeat region. The mt genome of *T. rubrofasciata* consisted of 13 protein-coding genes, 22 transfer

Table 2. Annotation of the complete mitochondrial genome of *Triatoma rubrofasciata*. tRNA abbreviations follow the IUPAC-IUB three letter code. For other abbreviation see legend for Fig. 1.

Gene	Stand	Nucleotide number	Anticodon	Start codon	Stop codon
tRNA-Ile	H	1-67	33-35 GAT	-	-
tRNA-Gln	L	65-133	101-103 TTG	-	-
tRNA-Met	H	133-200	163-165 CAT	-	-
ND2	H	219-1199	-	ATT	TAG
tRNA-Trp	H	1206-1271	1236-1238 TCA	-	-
tRNA-Cys	L	1264-1325	1293-1295 GCA	-	-
tRNA-Tyr	L	1327-1392	1358-1360 GTA	-	-
COI	H	1394-2922	-	ATG	T(aa)
tRNA-Leu	H	2928-2996	2961-2963 TAA	-	-
COII	H	2997-3672	-	ATT	T(aa)
tRNA-Lys	H	3676-3745	3706-3708 CTT	-	-
tRNA-Asp	H	3745-3808	3775-3777 GTC	-	-
ATP8	H	3809-3967	-	ATC	TAA
ATP6	H	3961-4644	-	ATG	TAA
COIII	H	4631-5416	-	ATG	TA(a)
tRNA-Gly	H	5416-5478	5446-5448 TCC	-	-
ND3	H	5476-5832	-	ATA	TAA
tRNA-Ala	H	5833-5894	5862-5864 TGC	-	-
tRNA-Arg	H	5898-5961	5927-5929 TCG	-	-
tRNA-Asn	H	5973-6036	6003-6005 GTT	-	-
tRNA-Ser(AGN)	H	6036-6108	6060-6062 GCT	-	-
tRNA-Glu	H	6108-6168	6137-6139 TTC	-	-
tRNA-Phe	L	6171-6234	6201-6203 GAA	-	-
ND5	L	6234-7946	-	ATT	TAA
tRNA-His	L	7944-8009	7975-7977 GTG	-	-
ND4	L	8012-9340	-	ATG	TAA
ND4L	L	9334-9627	-	ATG	TAA
tRNA-Thr	H	9630-9692	9660-9662 TGT	-	-
tRNA-Pro	L	9693-9757	9626-9628 TGG	-	-
ND6	H	9761-10261	-	ATG	TAA
CytB	H	10261-11391	-	ATG	TAG
tRNA-Ser(UCN)	H	11393-11459	11422-11424 TGA	-	-
ND1	L	11641-12573	-	ATT	TAA
tRNA-Leu	L	12559-12624	12593-12595 TAG	-	-
16S rRNA	L	12625-13890	-	-	-
tRNA-Val	L	13879-13949	13915-13917 TAC	-	-
16S rRNA	L	13952-14722	-	-	-
Control region		14723-17150	-	-	-

RNAs, 2 ribosomal RNAs and a control region (Table 2). The genome organization was consistent with *T. infestans*, *T. dimidiata* and other reduviid genomes, such as *Sirthena flavipes* [14] and *Oncocephalus breviscutum* [15]. However, there was 23 tRNA instead of 22 in *Brontostoma Colossus* [16]. Moreover, heavy strand (H strand) encoded 23 genes, whereas light strand (L strand) encoded the remaining 14 genes, and the overlap sequences were found at 16 locations. Most overlap sequence was only 1 bp and the longest overlap was 15 bp between ND1 and tRNA-Leu. In *T. rubrofasciata*, several intergenic

sequences were observed, however the intergenic spacer function was still unknown. The most longest intergenic spacer located between tRNA-Ser and ND1 was similar to the intergenic spacer of *Apis mellifera* that thought to be another origin of replication [3]. There was a bias toward A and T nucleotides in mt genome nucleotide composition. The overall nucleotide composition of H-strand was 34.9% A, 34% T, 12.9% G, and 18.2% C, thus biased toward A+T (68.9%).

The canonical start codons of protein-coding genes were ATN, GTG and TTG in invertebrate [17]. In *T. rubrofasciata*, the

start codon of 13 protein-coding genes was ATN, instead of GTG and TTG. The start codons for most of the coding sequences was ATT (ND2, COII, ND5, ND1) and ATG (COI, ATP6, COIII, ND4, ND4L, ND6, CytB), except that ATC for ATP8 and ATA for ND3. Almost all start codons of protein-coding genes were consistent with *T. infestans* and *T. dimidiata*. However, in *T. rubrofasciata*, ND5 start codons was ATT instead of GTG that was found in *T. infestans* and *T. dimidiata*. Most coding sequences were terminated with TAA (ATP8, ATP6, ND3, ND5, ND4, ND4L, ND6, ND1). The stop codons of ND2 and CytB were TAG. The COI and COII were stopped with incomplete T (aa) stop codons, and COIII stop codon was TA (a). The incomplete TA (a) and T (aa) stop codons were the deletion version of canonical TAA stop codon. It was common in metazoan mt genomes [11].

The 22 tRNA genes typically found in metazoan mt genomes were also identified in *T. rubrofasciata* ranging in length from 61 bp to 73 bp (Table 2). Most tRNA genes had the typical secondary structures except tRNA-Ser (AGN). It may be due to the formation of the dihydrouridine arm. The large rRNA subunit gene (lrRNA) was 1,266 bp long and the A+T content was 75.2%. The small rRNA subunit gene (srRNA) was 771 bp long and the A+T content was 70.8%.

The control region in *T. rubrofasciata* was 2,428 bp in length. It was smaller than *T. infestans* (2,653 bp), whereas was larger than *T. dimidiata* (2,166 bp). The control region was the most variable region between Reduviidae species [16]. This region in *T. rubrofasciata* was divided into 4 different parts. The first part was 467 bp in length and located downstream of srRNA gene and ended with 11 guanines. The second region (436 bp) was located next to first region, heavily biased toward A+T (78.4%). The third region was the longest region (1,357 bp) and composed of 2 different tandem repeat units. One was 760 bp in length with 5 consecutive 152 bp repetitions. The other had 2 consecutive 198 bp repetitions. The fourth part of the control region was 168 bp, ended at the beginning of the tRNA-Ile, which had the potential to form stem-and-loop structure.

We constructed a phylogenetic tree based on the analysis of 13 protein-coding genes (Fig. 2). As expected, *T. rubrofasciata* was clustered together with *T. dimidiata* and *T. infestans* that grouped in a highly supported clade. *Triatoma* species and *Oncocephalus breviscutum* (Stenopodainae) also grouped together in an extremely close clade. The close relationship between both subfamilies had been confirmed using morphological characteristics and phylogenetic analyses of the mitochondria

16S and nuclear 18S and 28S ribosomal DNA gene sequences [18,19].

In conclusion, our study details the complete sequencing and annotating of the mt genome of *T. rubrofasciata*. The complete mt genome contains 37 genes, including 13 protein-coding genes, 2 ribosomal RNA genes and 22 transfer RNA genes. Phylogenetic analyses using the amino acid sequences of the 13 protein-coding genes showed that the *Triatoma* species cluster together as a sister clade of Stenopodainae. This mt genome provides a unique genetic marker for studying the molecular biology, genetics and systematics of *T. rubrofasciata*.

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CONFLICT OF INTEREST

There is no conflict of interest related to this work.

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