



The Complete Mitochondrial Genome of *Rondotia menciana* (Lepidoptera: Bombycidae)

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RESEARCH

The complete mitochondrial genome of *Rondotia menciiana* (Lepidoptera: Bombycidae)Weiying Kong¹ and Jinhong Yang

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ABSTRACT. The mulberry white caterpillar, *Rondotia menciiana* Moore (Lepidoptera: Bombycidae) is a species with closest relationship with *Bombyx mori* and *Bombyx mandarina*, and the genetic information of *R. menciiana* is important for understanding the diversity of the Bombycidae. In this study, the mitochondrial genome (mitogenome) of *R. menciiana* was amplified by polymerase chain reaction and sequenced. The mitogenome of *R. menciiana* was determined to be 15,301 bp, including 13 protein-coding genes (PCGs), 2 ribosomal RNA genes, 22 transfer RNA genes, and an AT-rich region. The A+T content (78.87%) was lower than that observed for other Bombycidae insects. All PCGs were initiated by ATN codons and terminated with the canonical stop codons, except for *coxII*, which was terminated by a single T. All the tRNA genes displayed a typical clover-leaf structure of mitochondrial tRNA. The length of AT-rich region (360 bp) of *R. menciiana* mitogenome is shorter than that of other Bombycidae species. Phylogenetic analysis showed that the *R. menciiana* was clustered on one branch with *B. mori* and *B. mandarina* from Bombycidae.

Key Words: mitogenome, Bombycidae, diversity, phylogeny

Insect mitochondrial genomes (mitogenomes) are typically circular molecules 14–19 kb in length that contain 13 protein-coding genes (PCGs), 2 ribosomal RNA (rRNA) genes, 22 transfer RNA (tRNA) genes (Wolstenholme 1992, Boore 1999), and an A+T-rich region, which contains initiation sites for transcription and replication (Zhang et al. 1995, Zhang and Hewitt 1997, Taanman 1999).

Mitogenome sequences, which exhibit very low levels of recombination, are widely used in population genetics, comparative and evolutionary genomics, reconstruction of phylogenetic relationships, and evolutionary biology (Avice 1987, Ballard 2000, Ballard and Rand 2005, Cameron and Whiting 2008, Hao et al. 2012). The silk-producing insects in the lepidoptera with economic value belong to two families of moth, Bombycidae and Saturniidae (Mahendran et al. 2006). The complete mitogenomes of *Bombyx mori* and *Bombyx mandarina* of Bombycidae (Yukuhiro et al. 2002; Hu et al. 2010; Li et al. 2010a; Liu et al. 2013), and *Antheraea pernyi* (Liu et al. 2012b), *Antheraea yamai* (Kim et al. 2009), *Eriogyna pyretorum* (Jiang et al. 2009), *Samia cynthia ricini* (Kim et al. 2012), *Actias selene* (Liu et al. 2012a), and *Caligula boisduvalii* (Hong et al. 2008) of Saturniidae have been sequenced. The origin of bombycidae insects had been studied more according to the mitogenomes (Hu et al. 2010; Li et al. 2010a).

The mulberry white caterpillar, *Rondotia menciiana* Moore (Lepidoptera: Bombycidae) is a silk-producing insects from Bombycidae and has been exploited since the Yangshao culture period (approximately 5,500–6,000 years ago). As all the other insects from lepidoptera, *R. menciiana* is a bivoltine insect that exhibits four molts and a dormant period after the formation of resting eggs, too (Xu et al. 1994). The number of chromosomes (22) in *R. menciiana* differs from that of *B. mori*, (28) or *B. mandarina* (27 or 28) (Deng and Xiang 1993), and, thus, the genetic information of *R. menciiana* is important for understanding the diversity of the Bombycidae. *R. menciiana* larvae feed on mulberry leaves and can, in serious cases, defoliate trees. So, the natural *R. menciiana* populations have been decreasing, due to effective control of the insect by the Chinese government to prevent destruction of mulberry trees in recent years. At the same time, the research on genetic or the other aspects about *R. menciiana* was rare. In this study, the complete mitogenome sequence of *R. menciiana* was obtained (GenBank accession number: KC881286), and the phylogenetic

analyses based on the mitogenome of the selected insects from lepidoptera were performed using the maximum-likelihood (ML) method.

Materials and Methods

Specimen Sampling and DNA Extraction. Adult specimens of *R. menciiana* were collected from the Tsinling Mountains (106° 55'19" E, 34° 14'29" N), Shaanxi Province, China, in September 2011, preserved in 100% ethanol, and stored at –80°C until DNA extraction. Total genomic DNA was extracted from heads excised from frozen insects using the MagSi Tissue DNA Kit (Omega, GA).

Polymerase Chain Reaction Amplification and Sequencing. To amplify the entire mitogenome of *R. menciiana*, 10 primer sets (Table 1) were designed according to known mitochondrial DNA sequences from Bombycidae insects. Purified genomic DNA was amplified using the polymerase chain reaction (PCR) technique and the Taq PCR Kit (NEB, MA), under the following cycling parameters: 94°C for 3 min; 35 cycles of 30 s at 94°C, 40 s at 55–60°C, 1–3 min at 72°C; and 72°C for 10 min. The PCR products were detected by 1.0% agarose-gel electrophoresis and purified using a DNA gel extraction kit (TaKaRa, Japan). The purified PCR products were ligated into the T-vector (TaKaRa) and sequenced at least three times at Sangon.

Sequence Analysis and Gene Annotation. The BLASTN (<http://www.ncbi.nlm.nih.gov/blast>) was used to determine sequence similarity. Sequence assembly was performed using DNASTar software. The location of tRNA genes and potential stem-loop secondary structures were identified using tRNAscan-SE software version 1.21 (Lowe and Eddy 1997), specifying mito and chloroplast DNA as the source and using invertebrate mitochondrial genetic code predictors. Thirteen PCGs were identified using an open reading frame finder, using the invertebrate mitochondrial genetic code. The nucleotide composition and codon usage were calculated using MEGA 5.05 (Tamura et al. 2011) and the composition skewness to the formulas: AT skew = $[A - T] / [A + T]$; GC skew = $[G - C] / [G + C]$ (Perna and Koehler 1995). The putative control region was identified by alignment with sequences from the closely related species *B. mori* and *B. mandarina*,

Table 1. Primers used in this study

Primers	Location	Sequence(5'–3')	Mismatch
nad2-coxIF	335–354	TGATTGGDTGTTGAATTGGHYTAGAA	1
nad2-coxIR	1,952–1,927	GCTCCTAAGATTGAWGAAATACCWGC	2
coxI-coxIIF	1,830–1,850	TGGTGCAGGAACAGGATGAAC	3
coxI-coxIIR	3,791–3,771	GAGACCADTACTTGCTTTCAG	1
coxII-coxIIIF	3,673–3,692	ATTTGTGGRGCTAATCWTAG	2
coxII-coxIIIR	4,788–4,769	GGTCAAGGWCTATAATCYAC	1
coxIII-nad3F	4,511–4,528	TCGACCTGGAACCTTTAGC	1
coxIII-nad3R	5,727–5,709	TGGATCAAATCCACATTCA	1
nad3-nad5F	5,444–5,466	GAAGCAGCAGCTTGATATTGACA	2
nad3-nad5R	7,487–7,462	GCAGCTATAGCMGCTCCTACTCCWGT	1
nad5-nad4F	7,421–7,444	CCCCTGCTGTACTAAAGTTGAWG	0
nad5-nad4R	9,079–9,055	GGCTCTTTACCTTTATTAATRGAA	1
nad4-cytbF	8,892–8,914	GGAGCTTCTACATGAGCTTTGG	3
nad4-cytbR	10,906–10,885	CCCCTCAAAGWATATTTGACC	0
cytb-rrnLF	10,717–10,739	CGTACTTTCATGCWAATGGRGC	4
cytb-rrnLR	12,974–12,941	CTAATCAAYAGAAAAGWTTGCGACCTCGATGTTG	4
rrnLF	12,858–12,881	CGGTTTGAACCTCAGATCATGTAAG	0
rrnLR	13,920–13,895	TATTGTATCTTGTGTATCAGAGTTTA	0
rrnL-nad2F	13,304–13,335	ATGCTACCTTTGCACRGTCAAATACYGCRGC	1
rrnL-nad2R	588–563	TCAAAAATGAAATGGKGYTGAWCCTAT	3

Table 2. List of taxa used in this study

Superfamily	Family	Insect species	Accession number	References
Bombycoidea	Bombycidae	<i>R. menciana</i> ^a	KC881286	This study
		<i>R. menciana</i>	KJ647172	Kim et al. (2014)
		<i>B. mori</i> Xiafang	AY048187	Li et al. (2010b)
		<i>B. mori</i> C108	AB070264	Yukuhiro et al. (2002)
		<i>B. mandarina</i> Qingzhou	FJ384796	Hu et al. (2010)
		<i>B. mandarina</i> Japanese	GU966593	Li et al. (2010a)
		<i>A. pernyi</i>	AY242996	Liu et al. (2012b)
	Saturniidae	<i>A. yamamai</i>	EU726630	Kim et al. (2009)
		<i>Eriogyna pyretorum</i>	FJ685653	Jiang et al. (2009)
		<i>Samia cynthia ricini</i>	JN215366	Kim et al. (2012)
		<i>Ac. selene</i>	JX186589	Liu et al. (2012a)
		<i>Saturnia boisduvalii</i>	EF622227	Hong et al. (2008)
		<i>M. sexta</i>	NC_010266	Cameron and Whiting (2008)
		<i>Phthonandria atrilineata</i>	EU569764	Yang et al. (2009)
Geometridae	Geometridae	<i>Biston panterinaria</i>	JX406146	Yang et al. (2013)
		<i>Helicoverpa armigera</i>	NC_014668	Yin et al. (2010)
Noctuoidea	Noctuidae	<i>Spodoptera exigua</i>	JX316220	Wu et al. (2013)
		<i>Sesamia inferens</i>	JN039362	Chai and Du (2012)
		<i>Ochrogaster lunifer</i>	AM946601	Salvato et al. (2008)
		<i>Hyphantria cunea</i>	NC_014058	Liao et al. (2010)
		<i>Chilo suppressalis</i>	HQ860290	Yin et al. (2011)
Pyraloidea	Crambidae	<i>Diatraea saccharalis</i>	FJ240227	Li et al. (2011)
		<i>Ostrinia nubilalis</i>	NC_003367	Coates et al. (2005)
		<i>Cnaphalocrocis medinalis</i>	JQ305693	Yin et al. (2014)
Tortricodea	Tortricidae	<i>Adoxophyes honmai</i>	DQ073916	Lee et al. (2006)
		<i>Grapholita molesta</i>	HQ116416	Gong et al. (2012)
		<i>Spilonota lechriaspis</i>	HM204705	Zhao et al. (2011)
		<i>Choristoneura longicellana</i>	HQ452340	Unpublished
		<i>Acleris fimbriana</i>	HQ662522	Unpublished
		<i>D. melanogaster</i>	DMU35741	Clary et al. (1982)
Diptera	Drosophilidae	<i>D. melanogaster</i>	DMU35741	Clary et al. (1982)
Coleoptera	Tenebrionidae	<i>Tribolium castaneum</i>	AJ312413	Friedrich and Muqim (2003)

^a This study.

and the tandem repeats in the control region were predicted using the Tandem Repeats Finder program (Benson 1999).

Phylogenetic Analysis. The complete mitogenomes of 29 lepidopteran species (Table 2) were used to reconstruct the phylogenetic relationship. The mitogenomes of *Drosophila melanogaster* (Diptera: Drosophilidae) and *Tribolium castaneum* (Coleoptera: Tenebrionidae) were used as outgroups (Clary et al. 1982, Friedrich and Muqim 2003). The amino acid sequences of each of the 13 mitochondrial PCGs were aligned by Clustal X 1.83 using default settings (Thompson et al. 1997) and then backtranslated into nucleotide sequences after alignment. The concatenated set of nucleotide sequences were performed in

phylogenetic analysis, using ML method with the MEGA version 5.05 program.

Results

Genome Organization and Base Composition. In this study, the organization of *R. menciana* mitogenome was shown in Fig. 1. The complete mitogenome is a closed circular molecule of 15,301 bp in length, containing 13 PCGs (*coxI-III*, *nad1-6*, *nad4L*, *cyt B*, *atp6*, *atp8*), 22 tRNA genes, 2 rRNAs (*rrnL* and *rrnS*), and an A+T-rich region (Table 3). The order and orientation of *R. menciana*

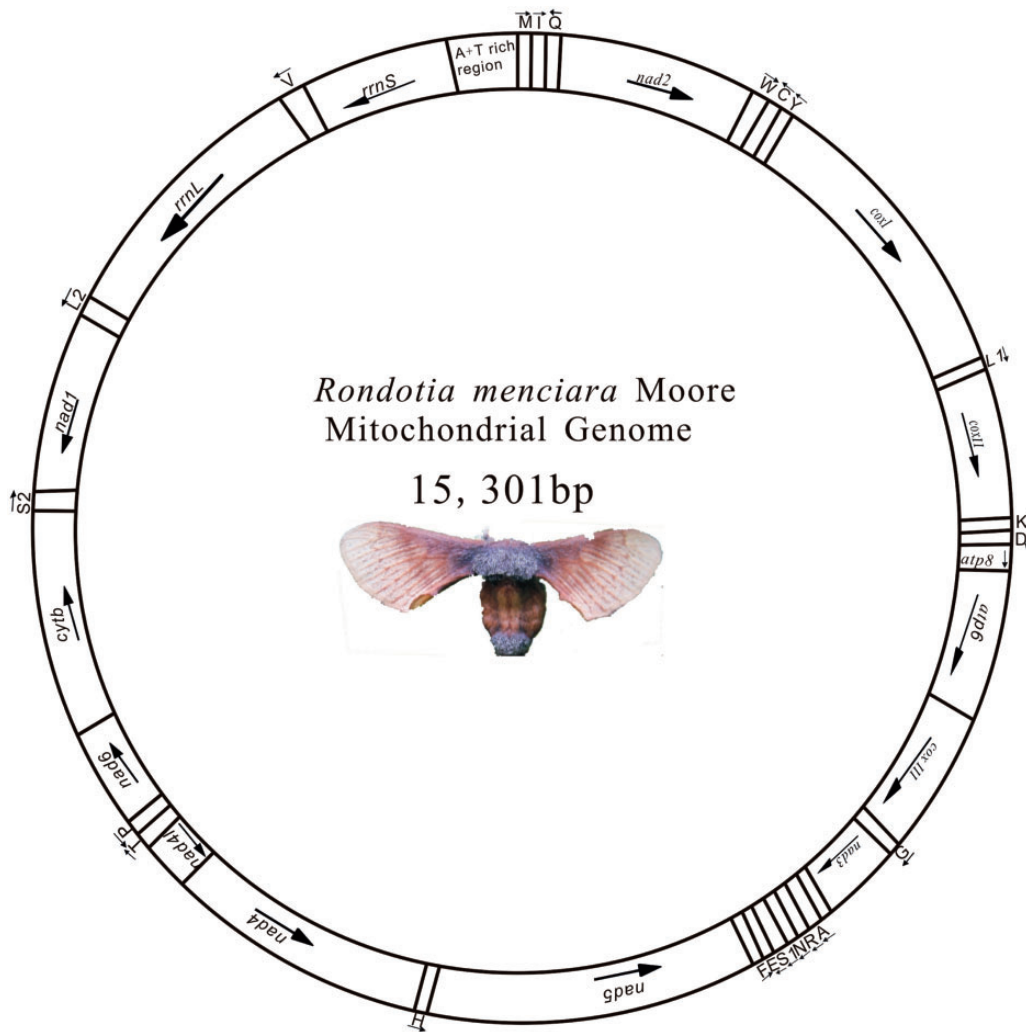


Fig. 1. Map of the mitogenome of *R. menciara*. The direction of all PCGs is designed by the underlined arrows. The transfer RNA genes are designated by single-letter amino acids codes. L1, L2, S1, and S2 denote *trnL(UUR)*, *trnL(CUN)*, *trnS(AGN)*, and *trnS(UCN)*, respectively.

mitochondrial genes are identical to those found in available lepidopteran mitogenomes (Cameron and Whiting 2008, Liu et al. 2013, Yang et al. 2013). The gene order *trnM*, *trnL*, and *trnQ* of the lepidopteran mitogenomes differs from the most common type *trnL*, *trnQ*, and *trnM*, which was found in a variety of insect orders and inferred to be ancestral for insects (Boore et al. 1998). The nucleotide composition of the *R. menciara* mitogenome was as follows: A (41.42%), T (37.45%), G (7.82%), and C (13.31%), and the A + T content (78.87%) was the lowest in the Bombycidae (Table 4). The AT skewness and GC skewness of *R. menciara* mitogenome was 0.050 and -0.26 , respectively, as observed in other Bombycidae, more biased toward A (the value of AT skewness is above zero) and C (Table 4).

PCGs and Codon Usage. The total length of 13 PCGs is 11,194 bp, and the A + T content of them is between 69.9% (*coxI*) and 89.51% (*atp8*) (Table 3). All PCGs in the *R. menciara* mitogenome were initiated by typical ATN codons: ATT for *nad2*, *coxI*, *atp8*, and *nad3* genes, ATA for *nad5*, *nad6*, and *cytb* genes, and ATG for the other six genes. Twelve of the PCGs were terminated with the canonical stop codons TAA or TAG, and *coxII* gene was terminated with a single T (Table 3). The presence of an incomplete stop codon seems a common phenomenon and had been found in several invertebrate mitochondrial genes (Jiang et al. 2009, Liu et al. 2013, Yang et al. 2013).

Codon usage of the PCGs exhibited a notable AT bias with an A + T composition of 77.05% (Table 4), which plays a major role in the A + T bias of the entire mitogenome. The six most frequently used codons in

the *R. menciara* mitogenome (TTA for Leu, ATT for Ile, TTT for Phe, ATA for Met, AAT for Asn, and TAT for Tyr) are composed of T or a combination of A and T, and the least frequent codons (CCG for Pro, TCG, AGG, and AGC for Ser, CGC for Arg, CTG for Leu, and CGG for Arg) have a high CG content (Table 5).

The analysis of the base composition at each codon position of the 13 PCGs of *R. menciara* mitogenome shows that the third codon position (87.4%) is considerably higher in A + T content than the first (72.8%) and the second (70.3%) ones (Table 6). Further analysis of the base composition at third codon position among Bombycidae insects shows that the highest and lowest A + T content appeared in the *R. menciara* (Kim et al. 2014) and this article, respectively, whereas the C + T content of each codon position was similar.

Overlapping and Intergenic Spacer Regions. Eleven overlaps, a total of 34 bp, were observed in the *R. menciara* mitogenome (Table 3). The largest overlap, 8 bp, occurred between the *trnW* and *trnC* genes, as reported in other Bombycidae species, such as *B. mandarina* (Li et al. 2010a) and *B. mori* Dazao (Liu et al. 2013). As in other lepidopteran species, a 7 bp overlap occurred between the *atp8* and *atp6* genes, as well. The *R. menciara* mitogenome harbors 17 short non-coding regions of 1–52 bp, for a total of 149 bp, as in the other insects from Bombycidae, the longest spacer is located between the *trnQ* and *nad2* genes (Table 3).

Transfer and rRNA Genes. As in other lepidopteran insects, all 22 tRNA genes with characteristic cloverleaf secondary structure in the

Table 3. Summary of the mitogenome of *R. menciiana*

Genes	Location	Size (bp)	Intergenic nucleotides	Direction	Anticodon	Start codon/stop codon	A + T(%)
<i>trnM(CAU)</i>	1–68	68		F	cat		77.94
<i>trnI(GAU)</i>	69–132	64	0	F	gat		78.13
<i>trnQ(UUG)</i>	130–198	69	–3	R	ttg		84.06
<i>nad2</i>	251–1,264	1,014	52	F		att/taa	84.12
<i>trnW(UCA)</i>	1,276–1,345	70	11	F	tca		82.86
<i>trnC(GCA)</i>	1,338–1,404	67	–8	R	gca		76.12
<i>trnY(GUA)</i>	1,405–1,473	69	0	R	gta		78.26
<i>coxI</i>	1,471–3,015	1,545	–3	F		att/taa	69.9
<i>trnL(UUR)</i>	3,011–3,078	68	–5	F	taa		73.53
<i>coxII</i>	3,079–3,760	682	0	F		atg/t	75.07
<i>trnK(CUU)</i>	3,761–3,831	71	0	F	ctt		74.65
<i>trnD(GUC)</i>	3,831–3,897	67	–1	F	gtc		88.06
<i>atp8</i>	3,898–4,059	162	0	F		att/taa	89.51
<i>atp6</i>	4,053–4,730	678	–7	F		atg/taa	76.84
<i>coxIII</i>	4,736–5,524	789	5	F		atg/taa	71.99
<i>trnG(UCC)</i>	5,527–5,592	66	2	F	tcc		86.36
<i>nad3</i>	5,590–5,946	357	–3	F		att/tag	79.55
<i>trnA(UGC)</i>	5,969–6,034	66	22	F	tgc		80.30
<i>trnR(UCG)</i>	6,037–6,101	65	2	F	tcg		78.46
<i>trnN(GUU)</i>	6,103–6,167	65	1	F	gtt		80.00
<i>trnS(AGN)</i>	6,170–6,238	69	2	F	gct		81.16
<i>trnE(UUC)</i>	6,240–6,306	67	1	F	ttc		92.54
<i>trnF(GAA)</i>	6,306–6,372	67	–1	R	gaa		85.07
<i>nad5</i>	6,372–8,102	1,731	–1	R		ata/taa	80.59
<i>trnH(GUG)</i>	8,107–8,173	67	4	R	gtg		83.58
<i>nad4</i>	8,186–9,526	1,341	12	R		atg/taa	78.23
<i>nad4L</i>	9,526–9,816	291	–1	R		atg/taa	83.51
<i>trnT(UGU)</i>	9,821–9,888	68	4	F	tgt		80.88
<i>trnP(UGG)</i>	9,889–9,953	65	0	F	tgg		78.46
<i>nad6</i>	9,956–10,468	513	2	F		ata/taa	81.68
<i>cytb</i>	10,471–11,622	1,152	2	F		ata/taa	74.39
<i>trnS(UCN)</i>	11,632–11,699	68	9	F	tga		80.88
<i>nad1</i>	11,717–12,655	939	17	R		atg/tag	75.08
<i>trnL(CUN)</i>	12,657–12,726	70	1	R	tag		78.57
<i>rrnL</i>	12,726–14,090	1,365	–1	R			83.37
<i>trnV(UAC)</i>	14,091–14,159	69	0	R	TAC		82.61
<i>rrnS</i>	14,160–14,941	782	0	R			84.4
A + T rich region	14,942–15,301	360	0				91.11

Table 4. Comparison of the nucleotides composition and skewness of Bombycoidea insects

Insect species	Whole genome			PCGs codon ^a		<i>rrnL</i>		<i>rrnS</i>		A + T rich	
	Size(bp)	A + T%	AT skewness/GC skewness	Size	A + T%	Size	A + T%	Size	A + T%	Size	A + T%
<i>R. menciiana</i> ^b	15,301	78.87	0.050/–0.260	11,157	77.05	1,365	83.37	782	84.4	360	91.11
<i>R. menciiana</i>	15,364	82.14	0.021/–0.195	11,178	80.96	1,398	85.77	775	85.03	360	91.11
<i>B. mori</i> Xiafang	15,664	81.35	0.058/–0.215	11,142	79.51	1,376	84.38	783	85.44	498	95.38
<i>B. mori</i> C108	15,656	81.36	0.059/–0.216	11,160	79.53	1,378	84.4	783	85.57	494	94.55
<i>B. mandarina</i> Qingzhou	15,717	81.42	0.057/–0.211	11,142	79.5	1,380	84.64	788	85.66	495	95.56
<i>B. mandarina</i> Japanese	15,928	81.73	0.054/–0.212	11,166	79.64	1,377	84.68	783	85.95	747	95.18
<i>A. pernyi</i>	15,566	80.16	–0.021/–0.216	11,181	78.46	1,369	83.86	775	84.13	552	90.4
<i>A. yamamai</i>	15,338	80.3	–0.022/–0.22	11,187	78.89	1,380	83.99	776	84.41	334	89.52
<i>Eriogyna pyretorum</i>	15,327	80.82	–0.031/–0.204	11,193	79.35	1,338	84.6	778	84.45	358	92.18
<i>Samia cynthia ricini</i>	15,384	79.78	–0.006/–0.228	11,196	78.26	1,358	84.02	779	83.83	361	90.86
<i>Ac. selene</i>	15,236	78.91	–0.023/–0.236	11,184	77.3	1,364	83.58	762	83.99	339	87.91
<i>Saturnia boisduvalii</i>	15,360	80.62	–0.024/–0.217	11,199	79.11	1,391	84.76	774	84.11	330	91.52
<i>M. sexta</i>	15,516	81.78	–0.005/–0.181	11,157	80.24	1,391	85.26	777	85.71	324	95.37

^a Termination codons excluded.^b This study.

R. menciiana mitogenome were predicted using the tRNAscan-SE Search Server. The length of tRNA genes ranged from 64 bp (*trnI*) to 71 bp (*trnK*) and the A + T content ranged from 73.53% (*trnL(UUR)*) to 92.54% (*trnE*) (Table 3). A total of six mismatches were found in five tRNA genes, two in amino acid acceptor stems, three in anticodon stems, and one in pseudouridine (TΨC) stems (Fig. 2). The mismatched bases show significant nucleotide bias, four U-U, one A-G, and one U-G.

Two rRNA genes were identified on the N-strand in the *R. menciiana* mitogenome: the *rrnL* gene, located between *trnL(CUN)* and *trnV* genes, and the *rrnS* gene, between the *trnV* gene and the A + T-rich region (Table 3). The length of *rrnL* and *rrnS* genes was 1,365 bp and 782 bp, and their A + T content was 83.37% and 84.4%, respectively.

A+T-Rich Region. The A + T-rich region of *R. menciiana* mitogenome was exactly same as that of Kim et al. (2014). The A + T-rich region was 360-bp long and located between the *rrnS* and *trnM* genes.

Table 5. Codon usage of PCGs in *R. menciiana* mitogenome

Codon	No. of codons	RSCU ^a	Codon	No. of codons	RSCU ^a
AAA(Lys)	92	1.69	TAA ^b	10	1.67
AAG(Lys)	17	0.31	TAG ^b	2	0.33
AAC(Asn)	39	0.31	TAC(Tyr)	27	0.28
AAT(Asn)	211	1.69	TAT(Tyr)	167	1.72
ACA(Thr)	78	1.88	TGA(Trp)	83	1.77
ACG(Thr)	5	0.12	TGG(Trp)	11	0.23
ACC(Thr)	22	0.53	TGC(Cys)	6	0.38
ACT(Thr)	61	1.47	TGT(Cys)	26	1.63
AGA(Ser)	86	2.21	TCA(Ser)	76	1.95
AGG(Ser)	2	0.05	TCG(Ser)	2	0.05
AGC(Ser)	3	0.08	TCC(Ser)	23	0.59
AGT(Ser)	31	0.8	TCT(Ser)	88	2.26
ATA(Met)	248	1.78	TTC(Phe)	37	0.2
ATG(Met)	30	0.22	TTT(Phe)	338	1.8
ATC(Ile)	43	0.2	TTA(Leu)	442	4.74
ATT(Ile)	394	1.8	TTG(Leu)	32	0.34
GTA(Val)	51	1.32	CTA(Leu)	51	0.55
GTG(Val)	10	0.26	CTG(Leu)	2	0.02
GTC(Val)	8	0.21	CTC(Leu)	6	0.06
GTT(Val)	86	2.22	CTT(Leu)	27	0.29
GAA(Glu)	53	1.49	CAA(Gln)	54	1.71
GAG(Glu)	18	0.51	CAG(Gln)	9	0.29
GAC(Asp)	7	0.21	CAC(His)	17	0.49
GAT(Asp)	60	1.79	CAT(His)	53	1.51
GCA(Ala)	38	1.32	CCA(Pro)	51	1.62
GCG(Ala)	4	0.14	CCG(Pro)	1	0.03
GCC(Ala)	14	0.49	CCC(Pro)	24	0.76
GCT(Ala)	59	2.05	CCT(Pro)	50	1.59
GGA(Gly)	86	1.78	CGA(Arg)	31	2.34
GGG(Gly)	41	0.85	CGG(Arg)	3	0.23
GGC(Gly)	7	0.15	CGC(Arg)	2	0.15
GGT(Gly)	59	1.22	CGT(Arg)	17	1.28

^a Relative synonymous codon usage.^b Stop codon.

The A + T content of the region was 91.11%, lower than the other Bombycidae insects (94.42–95.55%) (Yukuhiro et al. 2002; Hu et al. 2010; Li et al. 2010b; Liu et al. 2013). Several structures conserved in other Bombycidae mitogenomes were also observed in the *R. menciiana* A + T-rich region (Fig. 3). The conserved “ATAGA + poly T” motif with 17 consecutive Ts was located 24 bp downstream of the *rrnS* gene. A microsatellite (ATAT)_n element and a 12-bp poly-A region, commonly observed in other lepidopteran mitogenomes, were also found immediately upstream of the *trnM* gene. We also identified 2.6 tandem repeats elements of 37 bp in the *R. menciiana* A + T-rich region (Fig. 3).

Phylogenetic Analysis. In this study, the concatenated nucleotides sequences of 13 PCGs of 29 mitogenomes were used to reconstruct the phylogenetic relationships by ML method (Fig. 4). These 29 mitogenomes represent five superfamilies within the lepidopteran suborder: Bombycoidea, Geometroidea, Noctuoidea, Pyraloidea, and Tortricidea. The results show that the phylogenetic relationships among these five superfamilies are Tortricidea + (Pyraloidea + (Noctuoidea + (Geometroidea + Bombycoidea))), the relationship between Geometridae and Bombycoidea was close. The phylogenetic relationships inside Bombycoidea and Bombycidae were Bombycidae + (Sphingidae + Saturniidae) and *R. menciiana* + (*B. mori* + *B. mandarina*), respectively.

Discussions

The complete mitogenome of *R. menciiana* with a circular molecule of 15,301 bp was determined using the PCR method, which is the shortest in known complete mitogenomes of Bombycidae. The gene organization and order of *R. menciiana* mitogenome are identical to the studied lepidopteran mitogenomes (Cameron and Whiting 2008, Liu et al. 2013, Yang et al. 2013). The order of the *trnM* tRNA gene in the lepidopteran mitogenomes is A + T-rich region, *trnM*, *trnI*, *trnQ*, and *nad2*, whereas the deduced ancestral gene order is A + T-rich region,

Table 6. Summary of base composition at each codon position of PCGs in the Bombycidae mitogenome

Insect species	First codon position				Second codon position				Third codon position			
	T-1	C-1	A-1	G-1	T-2	C-2	A-2	G-2	T-3	C-3	A-3	G-3
<i>R. menciiana</i> ^a	36.4	10.8	36.4	16.4	48.2	16.3	22.1	13.4	46.2	7.7	41.1	5.0
<i>R. menciiana</i>	37.5	9.2	37.9	15.4	49.1	15.4	22.4	13.1	50.0	3.1	45.2	1.6
<i>B. mori</i> C108	37.3	9.7	37.0	16.0	48.4	16.2	22.0	13.4	49.0	4.4	43.9	2.7
<i>B. mori</i> Xiafang	37.3	9.6	37.1	15.9	48.4	16.2	22.0	13.3	48.9	4.6	43.8	2.7
<i>B. mandarina</i> Qingzhou	37.3	9.6	37.2	15.9	48.4	16.2	22.0	13.4	49.0	4.5	43.7	2.8
<i>B. mandarina</i> Japanese	37.4	9.6	37.2	15.8	48.4	16.2	22.2	13.3	49.2	4.3	43.8	2.8

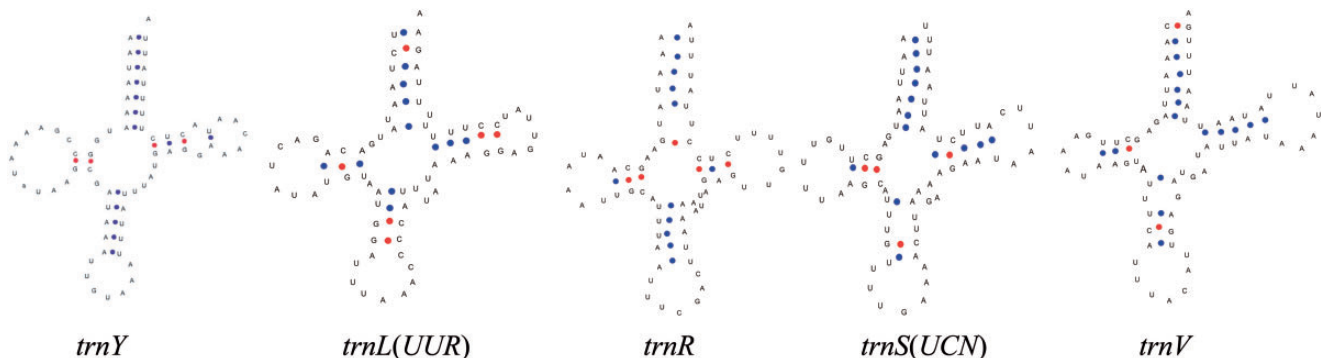
^a This study.

Fig. 2. Putative secondary cloverleaf structures for the tRNA genes of the *R. menciiana* mitogenome with mismatch bases. The blue point and red point indicate Watson–Crick base pairing A-U and G-C, respectively, and the blank indicate the mismatch bases. Six mismatches (four U-U, one A-G, and one U-G) lies in five tRNA genes (two in amino acid acceptor stems, three in anticodon stems and one in pseudouridine) (TΨC).

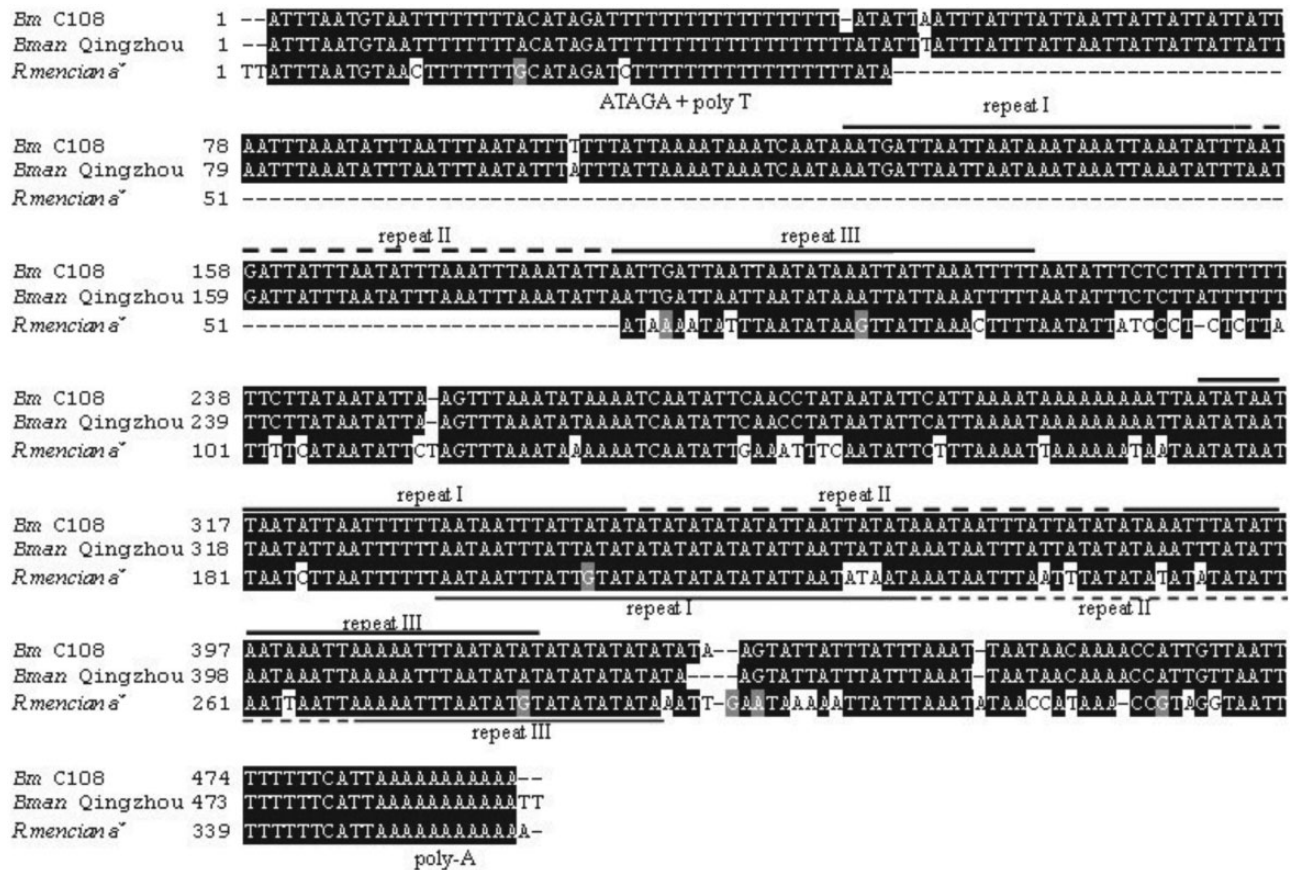


Fig. 3. Alignments of the A+T-rich region in Bombycidae. The thread underlined and thick overlined indicate the tandem repeat in *R. menciiana*, and *B. mori* and *B. mandarina*, respectively.

trnI, *trnQ*, *trnM*, *nad2* (Boore et al. 1998). This observation suggests that lepidopteran insects may have acquired the typical gene orientation and order independently after diverging from the ancestral insect.

The A + T content (78.87%) of the *R. menciiana* mitogenome is lower than the other Bombycidae insects (Yukuhiro et al. 2002; Cameron and Whiting 2008; Hong et al. 2008; Jiang et al. 2009; Kim et al. 2009; Hu et al. 2010; Li et al. 2010a; Liu et al. 2012b, 2013; Kim et al. 2014) (Table 4). The AT skewness of the *R. menciiana* mitogenome was 0.050 (Table 4), higher than the *R. menciiana* mitogenome (0.021) of Kim et al. (2014), and lower than that (0.054~0.059) of the *B. mandarina* and *B. mori*, indicating a higher occurrence of A compared with T nucleotides in Bombycidae. Different from the Bombycidae, there were higher occurrence of T compared with A nucleotides in the two families of Sphingidae and Saturniidae (Table 4). The GC skewness of the *R. menciiana* mitogenome was -0.260, indicating a heavy bias toward C versus G nucleotides and much more negative than observed for other Bombycoidea mitogenome (<0.236).

There was an incomplete stop codon of a single T in *R. menciiana coxII* gene. The incomplete stop codon had been found in several invertebrate mitochondrial genes, which seems a common phenomenon of mitochondrial genes (Jiang et al. 2009; Liu et al. 2013; Yang et al. 2013). The relative synonymous codon usage exhibits extensive similarity with other lepidopteran mitogenomes in previous study (Salvato et al. 2008). The most frequent codons in *R. menciiana* are composed of T or a combination of A and T, especially the third codon position. The observed differences in nucleotide composition may caused by the constraints on A + T content in the third codon position, which is more relaxed than those in the first and second codon positions due to degenerated genetic code (Taanman 1999). The C + T content in each of the codon positions were similar, which is agree with the point

that the high A + T content in insect mitogenome were caused by the mutation of C to T (The Honeybee Genome Sequencing Consortium 2006).

The length of the intergenic spacer regions (149 bp in 17 regions) is longer than that of *Manduca sexta* (115 bp in 13 regions) (Cameron and Whiting 2008) and *Ac. selene* (137bp in 13 regions) (Liu et al. 2012a) and is shorter than that of *C. boisduvalii* (194 bp in 16 regions) (Hong et al. 2008) and Bombycidae insects (about 338 bp) (Yukuhiro et al. 2002; Hu et al. 2010; Li et al. 2010a; Liu et al. 2013), which has been suggested to be constitutively synapomorphic and restricted to Ditrysia mitogenomes (Cameron and Whiting 2008, Hao et al. 2012). Similar to the mitogenome of the other insects, the tRNA genes have typical clover-leaf structure. However, the anticodon stem of *trnS(UCN)* could not form a stable stem-loop structure for the two U-U mismatches. The phenomenon of two U-G mismatches occurred also in the anticodon stem of *trnL(CUN)* of *E. pyretorum* (Jiang et al. 2009), *B. mori Dazao* (Liu et al. 2013), and *Ac. selene* (Liu et al. 2012a).

The exactly same A + T-rich region occurred between the *R. menciiana* mitogenome of this article and Kim et al. (2014). The length of 360 bp was shorter than that of Bombycidae insects (494 bp~747 bp) (Yukuhiro et al. 2002; Hu et al. 2010; Li et al. 2010a; Liu et al. 2013) and longer slightly than that of *A. yamamai* (334 bp) (Kim et al. 2009) and *C. boisduvalii* (330 bp) (Hong et al. 2008). The A + T content of the region was 91.11%, higher than that of *Ac. selene* (87.91%) (Liu et al. 2012a), *A. pernyi* (90.4%) (Li et al. 2011), *A. yamamai* (90.40%) (Kim et al. 2009), and lower than those from Bombycidae insects (94.42~95.55%). There are some common structures in the A + T-rich region of *R. menciiana* mitogenome, such as ATAGA + poly-T, and tandem repeats elements. The conserved motif of "ATAGA + poly T"

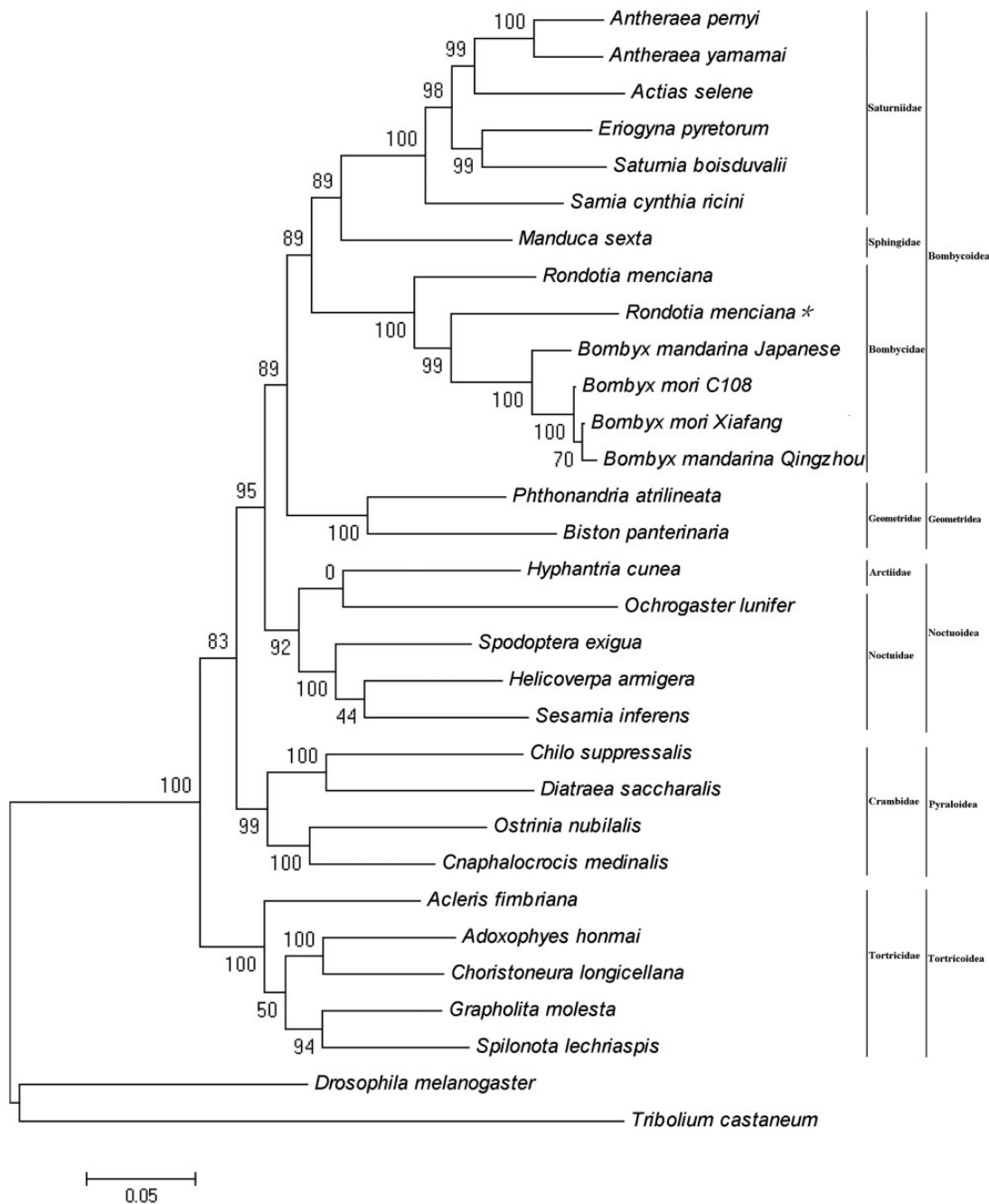


Fig.4. Phylogenetic analysis inferred from the concatenated nucleotides sequences of 13 PCGs of mitogenome using Mega 5.05 software and ML method. *D. melanogaster* and *T. castaneum* were used as outgroups. The numbers above the branches specify bootstrap percentages (1,000 replicates). *This study.

stretch at the 24 bp downstream of the *rrnS* gene is thought to be the origin of the DNA replication (Taanman 1999, Jiang et al. 2009). There are 2.6 tandem repeats elements of 37 bp in the *R. menciana* A + T-rich region, whereas three 32-bp and three 37-bp tandem repeats elements in *B. mori* C108 and *B. mandarina* Qingzhou. In *B. mori* Dazao, the A + T-rich region harbors two 31-bp repeat elements and three 36-bp repeat elements (Liu et al. 2013). The sequence and location of tandem repeats elements among Bombycidae mitogenomes are nonconserved. The existence of tandem repeats elements maybe caused mainly by the replication slippage.

Mitogenomes are effective markers for deep-level phylogenetic studies in the Lepidoptera. In our analysis, Bombycidae (*B. mori*, *B. mandarina*, and *R. menciana*), Sphingidae (*M. sexta*), and Saturniidae (*Ac. selene*, *A. pemyi*, *A. yamamai*, *S. cynthia ricini*, *E.*

pyretorum, and *C. boisduvalii*) were clustered in one branch of the phylogenetic tree, and the relationship of the three family was Bombycidae + (Sphingidae + Saturniidae), which is consistent with the morphological data and some previous studies (Zwick 2008, Zwick et al. 2011, Liu et al. 2013). The relationship of Geometridae is closer with Bombycoidea in our analyses, which is similar to the study by Yang et al. (2013).

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