Contents lists available at ScienceDirect



International Journal for Parasitology: Parasites and Wildlife

journal homepage: www.elsevier.com/locate/ijppaw



Comparative analyses of the fragmented mitochondrial genomes of wild pig louse *Haematopinus apri* from China and Japan



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ARTICLE INFO

Keywords: Anoplura Wild pig louse Mitochondrial genome Minichromosome

ABSTRACT

The wild pig louse *Haematopinus apri* is one of the commonest ectoparasites of wild pigs. In the present study, the entire mitochondrial (mt) genome of wild pig louse *H. apri* from China was sequenced and compared with previously characterized wild pig louse *H. apri* from Japan. We identified all of the 37 mt genes in the wild pig louse *H. apri* from China which are on nine circular minichromosomes. Each mt minichromosome is 2.9 kb–4.2 kb size and contains 2–8 genes and one non-coding region (1543 bp-2534 bp). The number of minichromosomes, gene content and gene order in the both mt genomes of wild pig louse *H. apri* from China and Japan are the same. The identity of the both mt genomes (except for non-coding regions) was 98.3% between wild pig louse *H. apri* from China is longer (3 bp) than that from Japan. For the 13 protein-coding genes, this comparison showed sequence differences in each gene at both the nucleotide (0.8%–2.4%) and amino acid (0.4%–3.5%) levels. The most conserved of these genes was the *nad*6, whereas the *nad*2 was least conserved at the nucleotide levels. This is the first comprehensive comparison of the mt genomes of a louse species from different geographic locations. This useful data provides additional genetic markers to study the phylogeny, systematics and population genetics of wild pig louse *H. apri*.

1. Background

The genus *Haematopinus* belongs to the family Haematopinidae of the suborder Anoplura, known as the blood-sucking lice (Barker, 1994; Durden and Musser, 1994; Meleney and Kim, 1974). This genus consists of 21 species, which mainly parasitize even-toed ungulates (Durden and Musser, 1994; Scofield et al., 2012). *Haematopinus* species are vectors of disease-causing microorganisms, such as African swine fever virus (Saegerman et al., 2021), swinepox virus (Thibault et al., 1998), classical swine fever virus (CSFV) (Wall R and Shearer D, 2008) and *Anaplasma* spp. (Da Silva et al., 2013). Meanwhile, wild pig louse *H. apri* is one of the most common ectoparasites of wild pigs.

Metazoan mitochondrial (mt) genomes are usually circular DNA organization (13–20 kb) that contain 36 to 37 genes: 12–13 proteincoding genes, two ribosomal RNA genes, and 22 transfer RNA genes (Boore, 1999; Lavrov, 2007; Wolstenholme, 1992). However, fragmented mt genomes have been reported from all sequenced blood-sucking lice (Dong et al., 2014a, b; Herd et al., 2015; Fu et al., 2020a, b; Jiang et al., 2013; Shao et al., 2012, 2015, 2017; Song et al., 2014) and some chewing lice (Song et al., 2019; Sweet et al., 2020, 2021). Although the suborder Anoplura contains 540 species of blood-sucking lice (Durden and Musser, 1994; Kim and Ludwig, 1978), to date, only mt genomes of 15 blood-sucking louse species have been sequenced and deposited in GenBank. In the genus Haematopinus, the entire fragmented mt genome was available for only three species (Jiang et al., 2013; Song et al., 2014). A recent study revealed highly divergent mt genomes in the booklouse (Liposcelis bostrychophila) from different geographical locations (Feng et al., 2018). This finding raised the possibility that parasitic lice from different geographical origins may have geographic differences in the number of minichromosomes, gene order, and nucleotide composition in their mt genomes. However, there is still no comprehensive comparison of the mt genomes of a louse species from different geographic locations. The mt genome sequences of wild pig louse H. apri from wild pig in Japan has been sequenced, but its non-coding regions have not been fully identified (Jiang et al., 2013). In addition, molecular data on wild pig louse H. apri from other countries and regions is totally lacking.

The objectives of the present study were: (i) to determine the number

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https://doi.org/10.1016/j.ijppaw.2022.03.013

Received 1 December 2021; Received in revised form 30 March 2022; Accepted 31 March 2022 Available online 2 April 2022

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of minichromosomes, gene order and nucleotide composition of the entire mt genome of wild pig louse *H. apri* from China, (ii) to compare this fragmented mt genome with that of wild pig louse *H. apri* from Japan, (iii) to test the hypothesis that wild pig louse *H. apri* from different geographic locations have geographic differences in the number of minichromosomes, gene order and nucleotide composition in their mt genomes.

2. Methods

2.1. Sample collection and DNA extraction

Adult specimens of wild pig louse *H. apri* were collected from a wild pig in Hunan province, China. These wild pig lice were washed five times in physiological saline solution, identified preliminarily to wild pig louse *H. apri* from China based on morphology and host information (Fig. 1) (Kim and Ludwig, 1978), and stored in 70% (v/v) ethanol at -40 °C. The total genomic DNA of wild pig louse *H. apri* from China was extracted from individual wild pig louse using a commercially available kit (Promega, Madison, USA) following the manufacturer's procedure. The identity of each wild pig louse specimen was further confirmed by PCR with primer reported previously (Fu et al., 2020a). The mt *cox*1 and *rrnS* gene sequences of the *H. apri* from China had 98.5% and 98.6% similarity with that of wild pig louse *H. apri* from wild pig in Japan (GenBank accession nos. KC814616 and KC814619), respectively.

2.2. Sequencing, assembling and annotation

The DNA concentration was evaluated using the Qubit system (Thermo Fisher Scientific, Waltham, MA, USA). Then, the entire genomic DNA was sequenced on Illumina Miseq Platform. The quality of raw data (300 bp each, paired-end reads) was assessed using FastQC 0.11.9 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc) and then were filtered for quality and trimmed using Skewer v0.2.2 (Jiang et al., 2013). Finally, 2 GB clean data was assembled using Geneious 11.1.5 (Kearse et al., 2012) based on obtained mt cox1 and rrnS gene sequences. We assembled all minichromosomes individually in full length as described previously (Fu et al., 2020a). The gene borders were predicted with MITOS web serves (http://mitos.bioinf.uni-leipzig. de/index.py) and manually curated. The location of protein-coding genes was further confirmed with BLAST searches of the NCBI database. tRNA genes were further confirmed using ARWEN (Laslett and Canbäck, 2008) and the program tRNAscan-SE (Lowe and Chan, 2016). The rRNA genes (rrnL and rrnS) were further confirmed by the boundary of the adjacent tRNA genes. The entire fragmented mt genome maps

were produced using Microsoft Powerpoint.

2.3. Verification of mt minichromosomes

The size and the circular organization of each mt minichromosome of wild pig louse *H. apri* from China were further verified by PCR using designed primers from the coding regions of each mt minichromosome (Table S1; Fig. S1). The two primers in each pair were next to each other with a small gap in between (10–90 bp). To obtain full-length sequences of each minichromosomes, the amplicons from nine full-length mt minichromosomes of wild pig louse *H. apri* from China were sequenced in a separate batch with high throughput sequencing as described above. The clean data of each minichromosomes was obtained and assembled as described above, respectively.

3. Results and discussion

3.1. Genome organization

By employing the sequencing from 6.099,122 \times 2 clean reads, we successfully assembled the entire mt genome of wild pig louse H. apri from China. We annotated 37 mt genes, including 13 protein-coding genes, two ribosomal RNA genes, and 22 transfer RNA genes (Fig. 2; Table 1). The nucleotide sequences of the mt minichromosomes of wild pig louse H. apri from China were deposited in the GenBank database under the accession numbers ON000914-ON000922. These genes were on nine circular minichromosomes, which was consist with those of previous studies (Jiang et al., 2013; Song et al., 2014). The mt minichromosomes of wild pig louse H. apri from China is 2.9 kb-4.2 kb in length, and each minichromosome has a coding region and a non-coding region (NCR) (Table 1). There are 2-8 genes in each coding region, varying in length from 787 bp to 2669 bp (Table 1). Each non-coding region varies in length from 1543 bp (nad2-trnI-cox1-trnL2 minichromosome) to 2534 bp (trnR-nad4L-nad6-trnM minichromosome) (Table 1). With the exception of nad1 gene, all mt genes are transcribed in the same direction.

3.2. Annotation

The wild pig louse *H. apri* from China mt genome contains 13 protein-coding genes, which had five initiation codons (ATA, ATG, TTG ATT, and GTG). Among them, four protein-coding genes use ATA (*cox3*, *nad5*, *nad6*, *atp6*), three genes use ATG (*nad1*, *nad2*, *nad4*L), three genes use TTG (*cox1*, *nad3*, *atp8*), two genes use ATT (*cox2*, *cytb*) and one gene use GTG (*nad4*). This mt genome has three termination codons (TAA,



Fig. 1. Female and male Haematopinus apri from China.





Fig. 2. The complete mitochondrial genome of wild pig louse *Haematopinus apri* form China. Each minichromosome has a coding region and a non-coding region (NCR, in black). The names and transcript orientation of genes are indicated in the coding region and the minichromosomes are placed in alphabetical order of protein-coding genes and rRNA genes. *Abbreviations: atp6* and *atp8*, ATP synthase F0 subunits 6 and 8; *cytb*, cytochrome *b*; *cox1-3*, cytochrome *c* oxidase subunits 1–6; *nad1-6* and *AL*; *rrnS* and *rrnL*, small and large subunits of ribosomal RNA. tRNA genes are indicated with their single-letter abbreviations of the corresponding amino acids.

Table 1

Mitochondrial minichromosomes of the wild pig louse *Haematopinus apri* from China, identified by Illumina sequencing.

Minichromosome	Size (bp)	Size of coding region (bp)	Size of non- coding region (bp)	Intergenic region (bp)
nad2-trnI-cox1-trnL ₂	4212	2669	1543	0
trnD-trnY-cox2-trnS ₁ -	3832	1840	1980	12
trnE-cytb-trnV	2780	1217	1561	2
trnQ-nad1-trnT-trnG- nad3-trnW	3603	1509	2087	7
trnK-nad4-atp8-atp6- trnN	3873	2297	1568	8
trnR-nad4L-nad6-trnM	3463	872	2534	57
trnH-nad5-trnF	3346	1777	1563	6
rrnS-trnC	3107	787	2319	1
trnL1-rrnL	3378	1226	2152	0
Total	31594	14194	17307	93

TAG and T). Among them, five genes use T (*cox1*, *cox2*, *cox3*, *nad2*, *nad3*), five genes use TAA (*cytb*, *nad4*, *nad5*, *nad6*), three genes use TAG (*nad4L*, *atp6*, *atp8*). The use of the start codon and stop codon of these protein-coding genes in wild pig louse *H. apri* from China is consistent with that in wild pig louse *H. apri* from Japan, except for *nad4* gene (use ATG for Japan). Incomplete termination codons (TA and T) are also present in other sucking lice, including horse louse *H. asini* (Song et al., 2014), rat louse *Hoplopleura kitti* (Dong et al., 2014b), rat louse *Polipleura kitti* (Dong et al., 2014b), rat louse *Polipleura kitti* (Herd et al., 2015), guanaco louse *Microthoracius praelongiceps* (Shao et al., 2017) and pubic louse *Pthirus pubis* (Shao et al., 2012). In the mt genome of wild pig louse *H. apri* from China, the sizes of the *rrnL* and *rrnS* genes were 718 bp and 1160 bp,

respectively. Sequence identity in the *rrn*L and *rrn*S genes is 98.2% and 99.2% between wild pig louse *H. apri* from China and Japan, respectively. The 22 tRNA genes identified in the mt genome of wild pig louse *H. apri* from China ranged from 63 to 75 bp in length. The secondary structure predictions in *H. apri* from China (not shown) were similar to those of horse louse *H. apri* from Japan and domestic pig louse *H. suis* (Jiang et al., 2013).

3.3. Non-coding regions

We obtained the entire non-coding region sequences of all of the nine mt minichromosomes of the wild pig louse *H. apri* from China, which range from 1543 bp (*nad2-trnI-cox1-trnL*₂ minichromosome) to 2534 bp (*trnR-nad4L-nad6-trnM* minichromosome) (Table 1). The non-coding regions contain tandem repetitive sequences (137 bp) and copy number (2–3 repeats). The non-coding regions of *H. apri* from China have 52.4%–96.7% pairwise identity to each other. There are sequence motifs in the non-coding regions that are highly conserved in many bloodsucking lice, such as *H. suis*, *H. apri* (Jiang et al., 2013), *Pedicinus obtusus*, *P. badii* (Fu et al., 2020b) and *M. praelongiceps* (Shao et al., 2017). As in previous studies (Shao et al., 2012; Jiang et al., 2013), a GC-rich motif (66 bp, 62.1% G and C) is downstream of the 3'-end of the coding region and an AT-rich motif (121 bp, 73.6% A and T) is upstream of the 5'-end. Interestingly, this study found another AT-rich motif (72 bp, 79.2% A and T) upstream of the GC-rich motif.

3.4. Comparative mt genomic analyses of H. apri from China and Japan

A comparison of the nucleotide and the amino acid sequences of mt genes for *H. apri* from China and Japan are given in Table 2. The nucleotide sequence variation across the entire mt genome (except for non-coding regions) between the wild pig louse *H. apri* from China and

Table 2

Nucleotide (nt) and/or predicted amino acid (aa) sequence differences in mitochondrial genes between the wild pig louse *H. apri* from China (HaC) and from Japan (HaJ) upon pairwise comparison.

Gene/ region	Nt sequence length	Nt difference (%)	Number of aa		aa difference (%)	
	HaC	HaJ	HaC/ HaJ	HaC	HaJ	HaC/ HaJ
atp6	666	666	1.1	221	221	0.5
atp8	174	174	1.7	57	57	1.8
nad1	900	900	1.6	299	299	1.3
nad2	997	997	2.4	332	332	1.8
nad3	346	346	1.7	115	115	3.5
nad4	1317	1316	1.7	438	438	1.1
nad4L	282	282	1.8	93	93	2.2
nad5	1638	1638	2	545	545	1.7
nad6	453	453	0.9	150	150	0.7
cox1	1528	1528	1.5	509	509	1
cox2	658	658	1.4	219	219	0.5
cox3	784	784	2	261	261	0.4
cytb	1083	1083	1.5	360	360	1.1
rrnS	718	718	0.8			
<i>rrn</i> L	1160	1159	1.8			
tRNA	1490	1488	1.7			

Japan was 1.7%. Difference across both nucleotide and amino acid sequences of the 13 protein-coding genes between the wild pig louse H. apri from China and Japan was 1.7% and 1.4%, respectively. The magnitude of nucleotide sequence difference in each gene between the wild pig louse H. apri from China and Japan ranged from 0.8% to 2.4%. The greatest difference between the wild pig louse H. apri from China and Japan was nad2 gene, whereas the least difference was detected in the rrnS gene (Table 2). Amino acid sequences inferred from individual mt protein-coding genes of wild pig louse H. apri from China and Japan were also compared. The amino acid sequence differences ranged from 0.4% to 3.5%, with cox3 gene being the most conserved protein-coding gene and nad3 gene the least conserved. Previous studies of other lice have also detected low intra-specific sequence variation in mt sequences. For example, the intra-specific sequence variation in P. pubis was 0-1.5% for pcox1 (775 bp) (Amanzougaghene et al., 2020) and 1.0% for pcytb (696 bp) (Light and Reed, 2009; Shao et al., 2012). The intra-specific sequence variation in P. badii was less than 0.3% for pcox1 (346 bp) (Light et al., 2010). Taken together, the molecular evidence presented here indicated the wild pig louse *H. apri* from China and Japan represent the same species.

4. Conclusions

The present study sequenced and compared the entire fragmented mt genomes of *H. apri* from China and Japan, and demonstrated that wild pig louse *H. apri* from different geographic locations is highly conserved in mt genomic composition and structure, refuted our hypothesis. This is the first comprehensive comparison of the mt genomes of a louse species from different geographic locations. The useful data provide useful genetic markers for studying the population genetics, molecular systematics and phylogenetics of blood-sucking lice.

Ethics approval and consent to participate

All procedures involving animals in the present study were approved and this study was approved by the Animal Ethics Committee of Hunan Agricultural University (No. 43321503).

Authors' contributions

GHL and YN conceived and designed the study, and critically revised the manuscript. YN and RL performed the experiments. YTF WW and YN analyzed the data. YN YTF and GHL drafted the manuscript. WQT helped in study design, study implementation, and manuscript preparation. All authors read and approved the final manuscript.

Funding

This study was supported in part, by the National Natural Science Foundation of China (Grant No. 32172884) and the Training Programme for Excellent Young Innovators of Changsha (Grant No. KQ2106044).

Consent for publication

Not applicable.

Declaration of competing interest

The authors declare that they have no competing interests.

Acknowledgements

Not applicable.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ijppaw.2022.03.013.

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