Novel mitochondrial genes and gene reannotation: conservation of gene arrangements among available astigmatid mites

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Abstract

Background

Mitochondrial genomes (mitogenomes) of metazoans typically contain 37 genes, comprising 13 protein-coding genes, two rRNA genes, and 22 tRNA genes. To date, complete mitogenome sequences of 15 species of Astigmatina are available, and they present variation in a number of features, such as gene arrangements, tRNA unconventional secondary structures, and the number and internal structures of control regions. Furthermore, 11 astigmatid mites from six superfamilies share the same gene arrangement. Two available species from the genus *Histiostoma* reportedly have different mitochondrial (mt) tRNA gene arrangements.

Results

We sequenced the mitogenomes of *Lepidoglyphus destructor* and *Gohieria fusca*, both from the superfamily Glycyphagoidea (Astigmatina). In total, 37 mt genes were identified in the two Glycyphagoidea species. Based on AT content and stem-loop structures, we divided the largest non-coding regions (LNRs) in *L. destructor* and *G. fusca* into two domains, respectively. The novel feature of two domains for the LNR was also found in *Acalvilia sp.* (Astigmatina, Hemisarcoptoidae). Using MITOS 2, tRNAscan, ARWEN, and manual approaches, we reannotated the mitogenomes of *Histiostoma blomquisti*, *H. feroniarum*, and *Trouessartia rubecula*. We reannotated six tRNA genes in *H. blomquisti* and four tRNA genes in *H. feroniarum*. We were able to identify all of the mt tRNA genes that were reported as lost in *Tr. rubecula*. The phylogenetic relationships found in our study were fairly consistent with previous studies of astigmatid mites phylogeny. Within Astigmatina, Glycyphagoidea was recovered as a monophyletic group.

Conclusions

A novel feature of the LNR was found in *L. destructor*, *G. fusca* and *Acalvilia sp.* (Astigmatina, Hemisarcoptoidae). This feature was not found in other available Astigmatina mitochondrial sequences. In the current study, most available astigmatid mitochondrial genomes shared the same consistent gene arrangement that could be the potential ancestral pattern in Astigmatina.

Background

Astigmatina (Astigmata, astigmatid) mites are economically and medically important pests that cause damage to stored food products [1] and trigger human allergies [2]. Glycyphagoidea mites belong to Acari, Acariformes, Astigmatina, which also are found in mattresses, pillows, and upholstery [3, 4]. Being derived from endosymbiotic α-Proteobacteria, complete mitogenomes have the usual characteristics of an entire genome, such as gene rearrangements [5], RNA secondary structures [6], and structural conservation of the non-coding regions [7]. Mitogenomes also supply information for phylogenetics, phylogeography, and population genetics [8–11]. Complete mitogenome sequences of Astigmatina have been reported for 15 species (as of October 9, 2020) from the six superfamilies: Acaroidea, Analgoidea, Hemisarcoptoidae, Histiostomatoidae, Pterolichoidea, and Sarcoptoidae. No complete mitogenome sequences from Glycyphagoidea, however, have been published in GenBank. To determine the types of gene rearrangements and to search for unique features in Glycyphagoidea, we sequenced and annotated the complete mitogenomes of *Lepidoglyphus destructor* (Schrank) and *Gohieria fusca* (Oudemans) (Sarcoptiformes, Glycyphagoidea).

The great majority of mitochondrial tRNAs in metazoans (> 90%) are predicted to share the canonical cloverleaf structure of tRNA sequences [12]. The conserved structure of tRNAs containing the 7 bp acceptor stem, a 5 bp anticodon stem and a D- and a T-arm have been characterized by tRNAscan-SE [13], ARWEN [14], and MITOS [15] in numerous research studies related to the analysis of insect tRNAs. Nevertheless, tRNAs that have lost either a D- or a T-arm (truncated tRNAs) are exceedingly common in sarcoptiform mites, possibly leading to the prediction of an implausible secondary structure or to the tRNA being missed during prediction.

In a previous study, two species from the genus *Dermatophagoides* had different tRNA gene arrangements [16]. After Klimov and O’Connor’s reannotation of *Dermatophagoides pteronyssinus*, the two species were considered to share a consistent gene arrangement [17]. In *Histiostoma*, there are two available complete mitogenome sequences from the same genus, and the tRNA gene arrangements are different [18].

The presence of the full set of tRNA genes is universal, because tRNA genes are essential to the translation process [19]. In Sarcoptoidae, two tRNA genes (*tmA* and *tmV*) are absent from *Sarcoptes scabiei*. Xue et al. identified the two mt tRNA genes that were reported as lost in *Sarcoptes scabiei* through a manual approach [18]. In Acaroidea, three mt tRNA genes (*tmF*, *tmST*, and *tmQ*) were reported as lost in *Ty. longior* [20] and *Ty. Putrescentiae* [21]. Fang et al. de novo sequenced and analyzed the mitochondrial genome of *Ty. Putrescentiae*, and identified the three tRNA genes that were reported as lost in a previous study [22]. In Analgoidea, five tRNA genes (*tmA*, *tmE*, *tmL*, *tmY*, and *tmV*) were reported absent from *Trouessartia rubecula* [23]. Nevertheless, this tRNA genes loss lacks convincing evidence.

The mitogenome gene arrangement of known astigmatid mites is rearranged compared with the hypothetical ancestor of the arthropods and shows several major rearrangements. Astigmatid mites’ mitochondrial genomes usually exhibit conserved gene arrangements. Ten of the fifteen species share one gene arrangement that is supposed to reflect the possible common ancestor of astigmatid mites [11].

Thus, we reannotated the mitogenomes of *H. blomquisti*, *H. feroniarum*, and *Tr. rubecula*. Our goals were to (1) investigate whether tRNA gene arrangements really exist in the genus *Histiostoma*, (2) ascertain whether or not any tRNA genes really have been lost in *Tr. rubecula*, and (3) document mt gene
rearrangements in the astigmata investigated to date.

Results

Mitochondrial genomes of Lepidoglyphus destructor and Gohieria fusca

We determined the complete mitochondrial genomes of L. destructor (GenBank accession number: MT075728) and G. fusca (GenBank accession number: MN608156), with lengths of 14,663 bp and 14,741 bp, respectively. Both contained the typical set of 13 protein-coding genes (PCGs), 22 transfer RNAs (tRNAs), two ribosomal RNAs (rRNAs), and (Table 1). The overall base composition of the entire L. destructor mitogenome consisted of 25.0% A, 39.2% T, 15.2% C, and 20.6% G. In G. fusca, the overall base composition of the entire mitogenome consisted of 25.8% A, 38.4% T, 15.8% C, and 20.1% G. In both species, the nucleotide composition had the same AT bias (64.2%). The combined protein-coding sequence lengths in the two species were 10,785 bp and 10,827 bp, respectively. In the G. fusca mitogenome, 11 protein-coding genes used standard start codons known as ATN. In addition, several unorthodox initiation codons were used. ATP8 initiated with GTG, and nad3 started with codon TTG (Table 1). In L. destructor, 12 protein-coding genes used standard start codons, and nad3 starts with codon TTG (Table 1).

Table 1. Mitochondrial genome organization of Lepidoglyphus destructor and Gohieria fusca. Int = intergenic nucleotides; negative numbers indicate overlapping nucleotides between adjacent genes.

Mitogenomes of L. destructor and G. fusca have been predicted to have the full set of tRNA genes. The putative mt tRNA genes were highly truncated in both L. destructor (46 to 61 bp) and G. fusca (48 to 61 bp). Only tmK showed the typical cloverleaf in these two species (Fig. S1, S2). Other tRNAs showed the reduction of tRNA-D- and/or T-arms. In L. destructor, we identified 21 tRNAs using automated prediction. We determined the tmV manually by alignment with homologous sequences in other astigmatid mites based on their anticodon and secondary structure. In G. fusca, we identified 19 tRNA genes using automated prediction. We identified the other three tRNA genes (tmN, tmV, and tmA) manually.

The novel feature of the largest non-coding region

Two large (>50 bp) non-coding regions (NCRs) could be identified in the L. destructor and G. fusca mitogenomes (Table 2). In L. destructor, the largest noncoding region (LNR, 813 bp) had an overall AT content of 68.4% and could be divided into two domains, domain I (463 bp) and domain II (350 bp), based on AT content and stem-loop structures (Fig. 1A). Domain I was on the J-strand at the 3’ end of tmF and had an AT content of 62.2%. A peculiar feature of this domain is the presence of a stem-loop structure with a length of 432 bp and short sequences (5’-GGGGGTAGGGG and CCCCTACCCCC-3’) (Fig. 1A I-box). Domain II was on the J-strand at the 5’ end of the tmS7 gene. The AT content was 78.9%, which was more AT-rich than domain I. Comparative sequence analysis of this domain identified conserved sequences, one microsatellite-like AT-repeat element, and several stem-loop structures (Fig. 1). In the mitochondrial genome of G. fusca, the LNR was 861 bp long and had an overall AT content of 70.7%. The LNR also could be divided into two domains. Domain I had a stem loop structure with a length of 435 bp (Fig. 1A). The short sequences (5’-GGGGGTAGGGG and CCCCTACCCCC-3’) were not found in domain I. The AT contents of domain I and domain II were 57.3% and 86.6%, respectively. The conserved sequences, one microsatellite-like AT-repeat element, and several stem-loop structures, were found in domain II. In total, we found the feature of two domains in both species from Glycyphagoidea.

Fig. 1. Conserved sequence blocks and secondary structures of the largest non-coding region in astigmatid mites. (A) Mapped on L. destructor, G. fusca, and Acalvolia sp. The domain I (striped box) and domain II (white box) are marked. The conserved sequences in domain I and the conserved palindromic sequences in domain II are shown in underlined letters. The A+T content of two domains is indicated. (B) Mapped on Aleuroglyphus ovatus, Caloglyphus berlesei, Rhizoglyphus robinii, Tyrophagusputrescentiae, Dermatophagoides pteronyssinus, D. farinae, Psoroptes cuniculi, Ardeacarus ardeae, Carpglyphus lactis, Sarcoptes scabiei, and Trouessartia rubecula (after revision). The conserved palindromic sequences are shown in underlined letters. Accession numbers of the analyzed sequences are reported in Table 6.

To investigate whether the feature of two domains existed in other astigmatid mites, we conducted a comparative analysis of non-coding regions from all available Astigmatid mitogenomes (Table 2). In astigmatid mites, most reported mitogenomes feature a compact structure that usually contains two conserved site-specific NCRs (except for Ty. longior, H. blo Mahmoudi, and H. feroniarum) and several nonconserved NCRs. These conserved site-specific NCRs are located between tmF-tmS1 and tmW-nad1 with high AT content (the average content is 84.8% A and 71.9%, respectively). The LNR is usually located in tmF-tmS1, and the length of this region averages 420 bp. The three longest conserved NCRs are in L. destructor, G. fusca, and Acalvolia sp. (813, 861, and 753 bp, respectively). The shortest NCRs located in tmF-tmS1 is only 76 bp in Sarcoptes scabiei.

Table 2. Distribution of NCRs in the astigmatid mites mitochondrial genomes.

Note: Non-coding regions marked with a star are assumed to be putative control regions. The data for Trouessartia rubecula, Histioctoma blo Mahmoudi, and H. feroniarum in the table are after our revision. Note that the several conserved sequences in domain II of L. destructor and G. fusca, including microsatellite-like (AT)n and several hypothetical stem-loop structures, were observed in the NCRs between tmF-tmS1 for most reported astigmatid mitogenomes (Fig. 1B). These conserved sequences were reported in the LNRs for Acaroidea species (Aleuroglyphus ovatus, Caloglyphus berlesei, Rhizoglyphus robinii, and Tyrophagusputrescentiae), Pyroglyphidae species (Dermatophagoides pteronyssinus and D. farinae), Hemisarcoptoidea species (Acalvolia sp., Carpglyphus lactis), and Sarcoptoidea (Psoroptes cuniculi) [22, 24-26]. In addition, we also found these conserved sequences in Pterolichoidea (Ardeacarus ardeae). Additionally, we observed microsatellite-like (AT)n sequence in the LNR in Sarcoptoidea (Sarcoptes scabiei) and Analogoeida (Trouessartia rubecula).
The other conservative non-coding region was located in \textit{tmW-nadI}, and the length of this region averaged 72 bp (Table 2). The longest reached 273 bp in \textit{Acalvolia sp.} The shortest was in \textit{Ardeacarus ardeae}, with only 29 bp. A similar sequence \([5'-(G)_nTA(G)_n-3']\) was found in the NCR of most available astigmatid mites (including the \textit{Tr. rubecula} and the \textit{H. feroniarum} after reannotation), except for \textit{Ardeacarus ardeae}. A similar sequence \([5'-(G)_nTA(G)_n-3']\) also was found in domain I of the LNR for \textit{L. destructor} and \textit{Acalvolia sp.}, except domain I of the LNR for \textit{G. fusca}. This sequence, \([5'-(G)_nTA(G)_n-3']\), seemed to be conserved in most available astigmatid mites.

Additionally, some astigmatid mites also exhibited other nonconservative NCRs (Table 2). Both \textit{H. blomquisti} and \textit{H. feroniarum} had one nonconservative NCR, and several stem-loop structures were found in these NCRs. The sequences and the stem-loop structures did not seem to be conserved.

Compared with the other described astigmatid mite NCRs, only \textit{Acalvolia sp.} (Astigmata, Hemisarcoptoidea) showed a similar feature of two domains in the LNR. In \textit{Acalvolia sp.}, the LNR also could be divided into two domains: a 469 bp fragment of domain I and a 284 bp fragment of domain II. The AT contents of domain I and domain II were 71.6% and 84.9%, respectively. In domain I, we found short sequences (5'-GGGGGTAGGGG and CCCCTACCCC-3') and a stem-loop structure. The conserved sequences among astigmatid mites were found in domain II (Fig. 1A).

As described earlier, the feature of two domains for the LNR was found only in \textit{L. destructor}, \textit{G. fusca}, and \textit{Acalvolia sp.} among all available astigmatid mites.

**Reannotation of four tRNA genes for \textit{Histiosoma blomquisti}**

In terms of previous study for \textit{H. blomquisti}, the smallest four tRNAs (\textit{trnA, trnS2, trnR,} and \textit{trnV}) could only be annotated manually, but the remaining tRNAs (\textit{trnC, trnF,} and the other 16 tRNAs) were identified using more than a manual approach [27]. To find more probable structures, we reannotated four tRNAs (\textit{trnC, trnA, trnF,} and \textit{trnS2}) based on the mitogenome from our analysis. We identified \textit{trnF} and \textit{trnS2} manually, and verified \textit{trnC} and \textit{trnA} based on the minimum free energy (MFE).

Our \textit{trnF} identified by manual sequence alignment was more conservative, showing fewer mismatches in the acceptor stem. This tRNA lies on the J-strand at the 3’ end of \textit{nad5}. Without predictions from tRNA search programs, we manually retrieved a tentative \textit{trnS2} from the sequence of previous study [27]. Similarly, we manually inferred the putative \textit{trnS2} to be a D-loop with fewer mismatches on stems. As a common phenomenon in Astigmatina, this D-loop was extremely truncated (43 bp). Compared with \textit{H. feroniarum} and other astigmatid species, \textit{trnS2} in our study was more conserved (Table 3).

**Table 3.** The alignment of nucleotide sequences of four mitochondrial tRNA genes

(\textit{trnS2, tmW, trnF,} and \textit{tmI}) in five reported astigmatid species in four different superfamilies.

Whenever contradictory predictions occurred, we calculated the minimum free energy (MFE) as a proxy. According to the MFE, we determined \textit{trnC} and \textit{trnA} in our study. These were more probable as having fewer mismatches on stems and arms (Fig. 2). After being amended, \textit{trnC} apparently shared 54bp nucleotides with the adjacent \textit{tmP}; these two genes overlapped and were on opposite strands (Table S1).

**Fig. 2** Secondary structures of \textit{trnC} and \textit{trnA} for \textit{Histiosoma blomquisti} and \textit{trnC, trnV,} and \textit{trnQ} for \textit{Histiosoma feroniarum} in our study.

*Note:* Bars indicate Watson-Crick base pairings; dots and circles represent GU pairs and mismatched pairs, respectively. *Hb* and *Hf* are obtained in this study. MFE indicated the minimum free energy value.

Because of the \textit{trnF} occupancy, the boundaries of \textit{nad5} in \textit{H. blomquisti} changed. Due to the tRNA reannotation, the NCRs also were changed. In a previous study, the NCRs were located in \textit{msT-trnV, trnF-trnA}. After the revision, one conserved site-specific NCR was identified between \textit{trnF-trnS1}, with high AT content. The other NCR was still located in \textit{msT-trnV}, with the length increased to 1598 bp (Table 2).

**Reannotation of six tRNA genes for \textit{Histiosoma feroniarum}**

In previous work, 16 of 22 tRNA genes (\textit{trnC, trnV, trnQ, trnF, trnW, tmI}, and 10 other tRNAs) were predicted using tRNA programs; the other six were identified manually [18].

In this study, we used manual annotation to reannotate \textit{tmF, tmW,} and \textit{tmI} of \textit{H. feroniarum}. Our \textit{tmF} for \textit{H. feroniarum} was identified by manual alignment and was conserved with the reannotated \textit{tmF} in \textit{H. blomquisti} (Table 3). In our analysis, \textit{tmW} was reannotated by sequence alignment. This sequence also had fewer mismatches (Table 3). In the previous study, the D-loop \textit{tmI} with the less common anticodon sequence AAU was reported in \textit{H. feroniarum} [18], but D-loop \textit{tmI} had never been reported in other Astigmatina. Our \textit{tmI} was reannotated manually based on anticodon sequences and secondary structures. In our study, we also observed the less common anticodon sequence AAU. We inferred this reannotated \textit{tmI} to be a TV-loop structure. This sequence was aligned to \textit{H. blomquisti} but was the only other remaining member of Histiosomatoidae. The reannotated \textit{tmI} presented considerable similarity (Table 3).

To select the most probable tRNAs, minimum free energy (MFE) was calculated for three tRNA genes (\textit{trnC, trnV, trnQ}). We found smaller MFE values than in previous annotations. In our study, \textit{trnC} was retrieved between \textit{tmS2} and \textit{tmP}, and \textit{tmV} was retrieved between 125S and 165S whereas the positions were coincidentally the opposite from those predicted by Xue et al. [18]. Our \textit{trnC} had common anticodon sequences GCA among astigmatid mites, whereas the previously described \textit{trnC} used ACA instead of GCA in anticodon sequences. When we calculated MFE, both \textit{trnC} and \textit{trnV} in our study were more thermodynamically stable (Fig. 2). After reannotation, the \textit{tmV} had 52 bp overlapping with the contiguous \textit{msS} on the same chain. Notably, the previously described \textit{tmQ} was annotated on the J-strand at the 3’ end of \textit{nad5} [18], but many mismatches on stems and arms were found in this position. In fact, most nucleotides of \textit{tmQ} among Astigmatina were highly conserved, and when located on the N-strand downstream of \textit{tmS1}, the reannotated \textit{tmQ} had a smaller MFE.
After amendment of the tRNA genes, the boundaries of \textit{nad2} of \textit{H. feroniarum} changed because of \textit{tml} occupancy. We also observed changes to the \textit{ml} size of \textit{H. feroniarum} on account of \textit{tmw} occupancy (Table S1). With alignment by Clustal W 2.0, the \textit{ml} boundaries of the two \textit{Histiostoma} species were more conserved. In addition, we changed one position of the NCR because of the reannotation of \textit{tmw} and \textit{tml}. After the revision, we could identify one conserved site-specific NCR with high AT content between \textit{tmw} and \textit{nad1}.

Notably, the reannotations of \textit{H. blomquisti} and \textit{H. feroniarum} mitogenomes led to the mtDNA gene order for both \textit{Histiostoma} species sharing the gene arrangement of a possible common ancestor of astigmatid mites, excluding the different locations of non-coding regions (Fig. 3).

\textbf{Fig. 3.} Mitochondrial gene arrangements in the sarcoptiform mites. The arrow pointing to the right represents the (+)-strand, and the arrow to the left the (−)-strand. Translocated or inverted genes are color-coded (green: inversion and translocation; pink: translocation; yellow: inversion). tRNA genes are in blue. The possible common ancestor among astigmatid mites is found in different genera and families. Species marked with a star: tRNAs reannotated in our study. Species marked with a triangle: the gene order in previous studies.

\textbf{Retrieving the “lost” mt tRNA genes in \textit{Trouessartia rubecula}}

In \textit{Trouessartia rubecula}, Esteban \textit{et al.} predicted tRNAs based solely on the MITOS WebServer, and five tRNA genes (\textit{trnA}, \textit{trnE}, \textit{tml}, \textit{tmY}, and \textit{tmV}) were not identified [23]. In protein annotation, the common start codon is ATN, but the \textit{cox1} gene start codon of \textit{T. rubecula} is the uncommon TCT. \textit{T. Rubecula} was reported to contain three D-loop sequences, which are lengths of these non-coding regions are normally short, and the location of these region is different from the common non-coding regions.

In the current study, we used trNAScan-SE, ARWEN, and MITOS 2 to predict the missing \textit{tmE}, \textit{tml}, and \textit{tmY}, and we annotated \textit{tnA} and \textit{tmV} using manual comparison. The secondary structure with the smallest constrained MFE was considered to be the most likely one (Fig. 4). We verified the tRNA genes of the two sarcoptiform mites by comparison of secondary structures and conserved nucleotide sequences with several Astigmata species (Table 4). Nucleotides that pair at the arms (acceptor arm, D arm, anticodon arm, and T arm) are underlined in the figure. We analyzed the codon usage in the mt protein-coding genes of the 17 species of astigmatid mites to see whether and how the corresponding codons of the “lost” tRNA genes were used. Overall, the codon use was similar between \textit{T. rubecula} and other astigmatid mites (Fig. 5).

\textit{Note:} Bars indicate Watson Crick base pairings; dots and circles represent GU pairs and mismatched pairs, respectively. MFE indicated the minimum free energy value.

\textbf{Table 4} The alignment of nucleotide sequences of two mitochondrial tRNA genes (\textit{tma} and \textit{tmV}) in six species. Conserved nucleotides are shaded in gray across these five mites.

\textit{Note:} Nucleotides that pair at the arms (acceptor arm, D arm, anticodon arm, and T arm) are underlined. Sequences of anticodons are in boldface.

\textbf{Aleuroglyphus ovatus} (Aa), \textit{Caloglyphus berlesei} (Cb), \textit{Rhizoglyphus robini} (Rr), \textit{Dermatophagoides farinae} (Df), \textit{Ardeacarus ardeae} (AA), and \textit{Trouessartia rubecula} (TR) in our study.

\textbf{Fig. 5} Relative synonymous codon usage (RSCU) and codon numbers of 22 amino acids (a, c, d, e, f, g, h, i, k, l, m, n, p, q, r, s1, s2, t, v, w, and y) in the mitochondrial (mt) genomes of astigmatid mites. The X-axis indicates the astigmatid mites; the Y-axis indicates the RSCU or total number of codons. The blue column indicates the codons that are complimentary to the anticodons of their corresponding mt tRNA. The red, green and purple columns indicate the imperfect, synonymous codons to the anticodons of their corresponding mt tRNA genes.

After revision, we reannotated the position of the \textit{cox1} gene. The reannotated \textit{cox1} gene had a length of 1536 bp and had ATA as the initiation codon rather than the uncommon start codon TCT identified in a previous study. The result from BLASTn supported our boundaries for \textit{cox1}. The LNR (301 bp in length) was arranged in the same way as most LNRs in the available sequences for astigmatid mites.

\textbf{New insight into astigmatid mitochondrial gene arrangement}

Compared with \textit{Limulus polyphemus} as the hypothetical ancestor of arthropods, gene orders were rearranged in the mitogenomes of 17 astigmatid mites (Fig. 3). We found that the mitogenomes of 13 astigmatid mites from six superfamilies (Glycyphagoidea, Acatroidea, Analgoidea, Sarcoptoida, Pterolichoidea, and Hemisarcoptoida) shared a consistent gene order, indicating that this gene arrangement was from the possible common ancestor of astigmatid mites. The mitogenomes of \textit{Histiostoma blomquisti}, \textit{Histiostoma feroniarum}, and \textit{Acalvolia sp.} represented three types of gene arrangements resulting from the different locations of the non-coding regions from the gene arrangement of the possible common ancestor of astigmatid mites (Fig 3). The mt gene order of \textit{Tyrophagus longior} was rearranged compared with other astigmatid mites because of tRNA loss.

\textbf{Phylogenetic analysis}

We constructed a phylogenetic tree based on a nucleotide data set from 13 mt PCGs of 17 astigmatid mites. The ML and BI analyses showed fully consistent topologies, and we provided the percentage of the bootstrap support at each node. The phylogenetic tree showed that the monophyly of Glycyphagoidea recovered with strong support (BPP = 1 and BSP = 100%), as well as the monophyly of Acatroidea, Hemisarcoptoida, and Histiostomatoidea (Fig. 6). The monophyly of two superfamilies, Analgoidea and Sarcoptoida, was always rejected. The logical next steps would be the integration of more astigmatina mitogenomes, including species from each of the six superfamilies.

\textbf{Fig. 6} Phylogenetic tree inferred from mitochondrial genome sequences using maximum likelihood and Bayesian methods. Branch lengths presented here reflect the Bayesian methods analysis. Node numbers indicate Bayesian posterior probabilities (BPP) and maximum likelihood bootstrap proportions (BSP). *
indicates clades with BSP <70%.

Discussion

For the first time, we determined the complete L. destructor and G. fusca mitogenomes in the superfamily Glycyphagoidea. Like most astigmatid mites, we identified 37 mitochondrial genes and one CR in the two Glycyphagoidea species. For comparison, the 15 other species of Astigmata with complete mitochondrial genomes available in GenBank exhibited an average genome size of 14,227 bp with high AT content. These values closely agreed with those we observed for both Glycyphagoidea species. The length variation of mitogenomes was usually due to length variation in the LNRs [28].

The LNRs in the mitochondrial genomes were less conserved than the coding genes, and they have not been well studied compared with other mitochondrial genes [29]. Astigmatid mites have two conservative non-coding regions and several nonconservative non-coding regions. The LNRs were usually located in tmF-tmS1, and several conserved sequences were found in the regions, including one microsatellite-like (AT)n element and several stem-loop structures. The conserved sequences were common and present in 12 astigmatid mites, including L. destructor and G. fusca, and these sequences were involved in the regulation of mitogenome replication [30].

In addition to abovementioned features, in L. destructor and G. fusca, based on AT content and hypothetical stem-loop structures, we divided the LNR into two domains and identified several conserved features in both species. Domain II was more AT-rich than domain I. Similar LNR features have not been reported for other Acari. A comparison of the LNR with available Astigmata mitochondrial sequences showed that similar unusual features of the LNR could be found only in Acalvolia sp. (Hemisarcoptoidea). Whether this LNR feature is general needs to be investigated by sequencing of additional Astigmata mitogenomes.

Truncated tRNAs have been found in all published mitogenomes of Acariformes [11, 12]. In addition, mismatches in the stems occur regularly in astigmatid mites. Both such tRNAs pose challenges for tRNA prediction programs [31]. In previous studies with two species of the genus Histioptera, tRNA gene arrangements were different from each other. In Tr. rubecula, five tRNAs were reported to be lost. In the current study, we reannotated the mitogenomes of H. blomquisti, H. feroniarum, and Tr. rubecula. In Histioptera, in addition to prediction programs, we used manual annotation by sequence comparison when alternative anticodons were considered [18]. We reannotated tmC, tmA, tmF, and tmS2 in H. blomquisti, and tmC, tmV, tmQ, tmF, tmW, and tmI in H. feroniarum. The reannotation of tmF (H. blomquisti) was based on a comparison among astigmatid mites. The tmS2 (H. blomquisti) and the tmF and tmW (H. feroniarum) were identified by alignment with the tRNA of the only other species in the same genus. The tmI was reannotated to have a TV-loop, which was a more conserved secondary structure than the D-loop of tmI identified previously. The D-loop of tmI was extremely rare in Astigmata, only known in H. feroniarum [18]. We used the MFE to verify tmC and tmA (H. blomquisti) and tmC, tmV, and tmQ (H. feroniarum). Compared with the tmC from a previous study, our tmC (H. feroniarum) had the common anticodon among astigmatid mites. In conclusion, the tRNAs after our reannotation contained more conserved sequences among homologous species of Histioptera or Astigmatina, and had more stable secondary structures than previously identified. Among these manually retrieved tRNAs, the extreme situations were found in tmC of H. blomquisti and tmV of H. feroniarum. The genes overlapped between tmC-tmP on the opposite strand; thus, the transcripts did not overlap [32]. The overlap between tmV-trnS in H. feroniarum was on the same chain. Sequence overlaps universally existed in astigmatid species and may be corrected later through post-transcriptional editing, alternate treatment, and relaxed restrictions on tRNA structure [33, 34]. Whether the consistent gene arrangement for Histioptera is general needs to be investigated by sequencing additional Histioptera mitogenomes.

After the revision, the Tr. rubecula had a full set of tRNAs. Codons for all 22 amino acids were present in the protein-coding genes in all astigmatid mites. No evidence has yet indicated that nuclear tRNA genes can be imported into mitochondria in mites or other animals [19]. If an mt tRNA gene is indeed lost and there is no nuclear replacement, then its corresponding codons in the mt protein-coding genes will not be translated. Thus, the "loss" of mt tRNAs may not exist in astigmatid mites. Rearranged gene orders in the mitogenomes not only can provide phylogenetic information but also are useful in resolving the phylogeny of insects [35, 36]. Therefore, the correct annotation of genes is important for phylogenetic and evolutionary studies. Our study had four types of gene arrangements in Astigmatina. The mitogenomes of 13 astigmatid mites from six superfamilies shared one type of gene arrangement. This gene arrangement likely was the most conserved gene arrangement among the astigmatid mites. In addition, these four types of gene arrangements for 37 genes (13 PCGs, 22 tRNAs, and 2 rRNAs) were consistent. The gene arrangement of Ty. longior represented a unique type because of missing tRNAs [20]. Two species had complete mitogenomes reported in the genus Tyrophagus. The gene arrangement of Ty. putrescentiae was consistent with the possible common ancestor of astigmatid mites [22]. Murillo et al. used COI to analyze molecular phylogenetics and suspected that a misidentification existed in the Ty. longior [20] sequence, as it clustered with other sequences of Ty. putrescentiae [36]. Hence, our results did not support the loss of mt tRNA genes in these astigmatid mites. Our results did support the hypothesis that the conserved gene arrangement in most available species was likely the arrangement of the possible common ancestor of astigmatid mites [11].

Within the Astigmata, we observed the monophyly of Glycyphagoidea, Hemisarcoptoidea, Acaroida, and Histioptera. Consistent with a previous study, the monophyly of two superfamilies, Sarcotoidea and Analgoidea, was rejected [11, 23, 31]. Our finding supported the suggestion that the phylogeny of Astigmata is far from resolved at the superfamily level [11, 38, 39].

Conclusion

In this study, we determined the complete L. destructor and G. fusca mitogenomes. A novel feature of the LNR was found in L. destructor, G. fusca, and Acalvolia sp., the feature that was not found in other available Astigmata mitochondrial sequences. We reannotated the mitogenomes of Tr. rubecula, H. blomquisti, and H. feroniarum. The gene order of 13 astigmatid mites from six superfamilies shared the gene order that was identical to the possible common ancestor of astigmatid mites. This information from our study has important ramifications for our understanding of mitogenome evolution in astigmatid mites.
**Methods**

**Mite collection and DNA extraction**

We collected samples of *Gohieria fusca* from a flour factory in Wuhu, southeast China, in July 2016. We collected samples of *L. destructor* from a flour shop in Wuhu in spring 2018. *L. destructor* and *G. fusca* were reared at 25°C and 85% relative humidity in the dark using the following medium: wheat germ, corn flour, and baker's yeast at a ratio of 10:10:1 by weight [40]. Mites were stored in 100% ethanol at -20°C until use. We identified specimens based on their morphological characteristics [41]. The cox1 sequences of *L. destructor* and *Gohieria fusca* were obtained with the primers LWCO1U and LWCO1L, as suggested by Webster et al. [42]. After performing BLASTn searches of the nucleotide collection (nr/nt) database of the NCBI, the results showed that the cox1 sequence from two species were very similar to several sequences from *L. destructor* and *Gohieria fusca*. In *L. destructor*, the identities were 98.91% (accession number: EU078972) and 98.64% (accession number: AY525569). In *G. fusca*, the identities were 98.64% (accession number: MG279724) and 98.38% (accession number: MG279719). Therefore, the two specimens collected in the current study were identified as *L. destructor* and *G. fusca*. We extracted whole-genomic DNA by standard phenol-chloroform extraction [43].

**Sequence assembly, annotation, and analysis**

We obtained the mitogenomes of *L. destructor* and *G. fusca* by the Shanghai Majorbio Bio-pharm Biotechnology Company (Shanghai, China) and Shanghai BIOZERON Company (Shanghai, China), respectively. Both were sequenced with an Illumina HiSeq sequencer [44]. We annotated the assembled genomes using the MITOS WebServer [15] using the mitochondrial invertebrate genetic code for invertebrates. We calculated relative synonymous codon usage and amino acid frequencies using MEGA X [45]. The PCGs boundaries were confirmed manually in MEGA X software and BLASTp searches conducted at the NCBI website [46]. We identified secondary structures of the LNR using the Mfold web server [47]. The two rRNA genes, rml and rmS, were curated by BLAST searches based on sequence similarity. The boundaries of tRNAs were assumed to be immediately after their upstream genes and before their downstream genes [18]. To predict tRNAs, we used tRNAscan-SE [13], ARWEN [14], MITOS, and MITOS2 [15]. The tRNAs that could not be predicted by programs were determined manually by alignment with their homologues in other astigmatid mites [11]. We also used the manual comparative approach to determine the results of tRNA search programs.

**The largest non-coding region of mitogenomes amplification and sequencing**

To verify the number of variable length AT-repeats in the LNR, we extracted DNA from 10 individual mite offspring from a single female using the DNeasy Blood and Tissue Kit (QIAGEN, Hilden, Germany). According to the obtained sequences, we designed specific polymerase chain reaction (PCR) primers for each species (Table 5) to amplify the remaining genome by long PCR as a single fragment using the manufacturer's rapid PCR protocol.

We amplified the LNRs for *G. fusca* and *L. destructor* using four flanking primers. The primer specifications and the length of each amplified fragment are given in Table 5. We used prime STAR GXl DNA polymerase (Takara) for PCR under the following cycling conditions: 30 cycles of 98°C for 10 s, 52°C for 16 s, and 68°C for 1 min 40 s. We conducted PCR amplification reactions in 12.5 μl volumes containing 2.5 μl 5 × buffer, 1 μl dNTP, 0.25 μl GXl DNA Polymerase, 0.25 μl of each primer, 0.5 μl of template DNA, and double-distilled water to a final volume of 12.5 μl. Finally, PCR products were separated by electrophoresis on a 1% agarose gel. PCR products were purified with a QIAquick Spin PCR Purification Kit (Qiagen). PCR products were cloned into a Blunting Kit (BKL) (Takara). After heat-shock transformation of *Escherichia coli* (DH5a cells), we aligned their similarity against the sequence by using Clustal W 2.0 [48].

**Table 5** The primers used for the amplification of *Gohieria fusca* and *Lepidoglyphus destructor* mitochondrial genomes, with the size of each fragment that was amplified in bp.

**The reannotations of *Histostoma blomquisti*, *H. feroniarum*, and *Trouessartia Rubecula* mitogenomes**

The assembled genome was annotated using the MITOS WebServer [15]. The boundaries of PCGs were confirmed manually by MEGA X version 10.1.5 [45] software and BLASTp searches conducted in the NCBI database [46]. Based on highly conserved sequence motifs, rml and rmS were identified by BLAST searches of the NCBI database. To predict tRNAs, we used tRNAscan-SE [13], ARWEN [14], and MITOS 2. The tRNA genes that could not be identified by the tRNAscan-SE, ARWEN, and MITOS 2 were determined manually by alignment with their homologues in related species in the Astigmatid mites based on their anticodon and secondary structures [11]. If a tRNA can not be determined by automated prediction or manual comparative approach, such as predictions of the different tools were in some cases contradictory, the Mfold Server [47] and RNAeval (ViennaRNA package v.2.3.3) [49] were used for calculating the minimum free energy (MFE) to select the most probable among alternative structures (constrained analysis). The secondary structure with the smallest constrained MFE was considered as the most stable one [6]. We also used the manual comparative approach to determine the results of tRNA prediction programs.

**Phylogenetic analysis**

To infer the phylogenetic positions of *L. destructor* and *G. fusca* within the Astigmata, we generated a dataset of 19 mite taxa (17 astigmatid mites and two orbibatid mites) (Table 6) that included only those species with a complete set of 13 PCGs. First, the nucleotide sequences of each PCG was translated under the invertebrate mt genetic code. Then the amino acid sequences of 13 PCGs were aligned individually by MAFFT v7 [50]. Additionally, large gaps and ambiguous sites were deleted by Gblocks v.0.91b [51]. Finally, we conducted phylogenetic analyses using maximum likelihood (ML) and Bayesian inference (BI) methods.

**Table 6. Mitochondrial genomes employed in this study.**

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We analyzed the dataset of amino acid sequences of 13 PCGs as two types of matrices: combined algorithm and partitioned algorithm. Dataset partitioning was performed by PartitionFinder v.2.1.1 [52] based on an initial total of 13 data blocks (amino acid sequences, 13 protein-coding genes). We predicted the models by PartitionFinder using the Akaike information criterion (AICc). PartitionFinder used unlinked branch lengths, and the greedy search algorithm was used for amino acid sequences and the MrBayes model. For the dataset of partitioned amino acid sequences based on genes, two substitution models [GTR+H+G (COI, COII, ATP6, NAD3, NAD1, NAD2, NAD4, NAD4L, NAD5, NAD6, CYTB), GTR+G (ATP8, COIII)] were chosen by PartitionFinder. For the dataset of combined amino acid sequences, the substitution model GTR+H+G was chosen by PartitionFinder.

For the dataset of amino acid sequences, we performed BI analyses with MrBayes v.3.2.2 [53]. For MrBayes v.3.2.2, we used separate data partitions plus mixed models and conducted two independent runs each with four Markov chain Monte Carlo runs (one cold chain and three heated chains). The two datasets were run for 2 million generations, with trees sampled every 1000 generations. We then applied a conservative burn-in of 25%. We evaluated the convergence of the parameter estimates with TRACER v.1.6. All estimated parameters showed ESS values above 200. We edited the consensus tree with FigTree v.1.4.0. Nodes supported by Bayesian posterior probabilities (BPP) ≥95% were considered strongly supported [54]. We performed ML analyses on the combined and partitioned datasets using the GTRGAMMAI model in the RAxML-7.0.3-WIN [55]. We assessed clade support using a nonparametric bootstrap with 1000 replicates. We considered nodes supported by bootstrap values (BSP) ≥70% to be strongly supported [56].

**Abbreviations**

*atp6* and *atp8*: Genes for ATP synthase subunits 6 and 8; bp: Base pair; cob: Gene for cytochrome b; *cox1*, *cox2*, and *cox3*: Genes for cytochrome oxidase subunits 1, 2, and 3; CR: Control region; DNA: Deoxyribonucleic acid; mitogenome: Mitochondrial genomes; Int: Intergenic nucleotides; LNRs: Largest non-coding regions; PCGs: Protein-coding genes; PCR: Polymerase chain reaction; rRNA: Ribosomal RNA; *rrnS* and *rrnL*: Genes for small and large subunits of ribosomal RNA; RSCU: Relative Synonymous Codon Usage; *tmA*: tRNA gene for alanine; *tmC*: tRNA gene for cysteine; *tmD*: tRNA gene for aspartic acid; *tmE*: tRNA gene for glutamic acid; *tmF*: tRNA gene for phenylalanine; *tmG*: tRNA gene for glycine; *tmH*: tRNA gene for histidine; *tmI*: tRNA gene for isoleucine; *tmK*: tRNA gene for lysine; *tmL1*: tRNA gene for leucine; *tmL2*: tRNA gene for leucine; *tmM*: tRNA gene for methionine; *tmN*: tRNA gene for asparagine; *tmP*: tRNA gene for proline; *tmQ*: tRNA gene for glutamic acid; *tmR*: tRNA gene for arginine; *tmS1*: tRNA gene for serine; *tmS2*: tRNA gene for serine; *tmT*: tRNA gene for threonine; *tmV*: tRNA gene for valine; *tmW*: tRNA gene for tryptophan; *tmY*: tRNA gene for tyrosine; μl: Microliter.

**Declarations**

**Ethics approval and consent to participate**

No specific permits were required for the mites collected for this study in China. The mite specimens were collected from mushrooms, and the field studies did not involve endangered or protected species. The species in our study are common mites and are not included in the “List of Protected Animals in China”.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Availability of data and materials**

The datasets generated and/or analysed during the current study are not publicly available due [the mt genomes of *Lepidoglyphus destructor* and *Gohieria fusca* submitted on GenBank under the accession numbers MT075728 and MN608156, and not released yet] but are available from the corresponding author on reasonable request.

**Authors' contributions**

SET, XXQ, ZSL, and FYu designed the study. FYu, XJY, ZXB performed the laboratory work. FYu, XJY, SX, CLM, FR, and FYing analyzed the data. FYu, XJY, ZXB, FWX, DFY, and ZZT contributed to draft writing and literature search. SET, ZSL, XXQ, and LF reviewed and revised the manuscript. All authors read and approved the final manuscript.

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Tyrophagus putrescentiae

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Additional Files

Additional file 1: Figure S1 Predicted secondary structures of the 22 mt tRNAs of Lepidoglyphus destructor. Bars indicate Watson-Crick base pairings; dots and circles represent GU pairs and mismatched pairs, respectively.

Additional file 2: Figure S2 Predicted secondary structures of the 22 mt tRNAs of Gohieria fusca. Bars indicate Watson-Crick base pairings; dots and circles represent GU pairs and mismatched pairs, respectively.

Additional file 3: Table S1 Mitochondrial genome organization of Histiostoma blomquisti and Histiostoma feroniarum.

Tables

Table 1 Mitochondrial genome organization of Lepidoglyphus destructor and Gohieria fusca. Int = intergenic nucleotides; negative numbers indicate overlapping nucleotides between adjacent genes.
| Strand | Gene   | *Lepidoglyphus destructor* |                     | *Gohieria fusca* |                     |
|--------|--------|----------------------------|-------------------|-----------------|-------------------|
|        |        | Position                  | Size   | int | AA  | start | stop  | anti | Position                  | Size   | int | AA  | start | stop  |
| J      | cox1   | 1-1548                    | 1548   | 2   | 515 | ATT   | TAG   | 1-1548 | 1548   | 7   | 515 | ATT   | TAG   |
| J      | cox2   | 1551-2306                 | 756    | 2   | 251 | ATA   | TAA   | 1556-2311 | 756    | 6   | 251 | ATG   | TAA   |
| J      | tmD    | 2309-2362                 | 54     | -1  |     |       |       | GTC   | 2318-2371 | 54    | -1  |     |       |
| J      | Atp8   | 2362-2517                 | 156    | 5   | 51  | ATC   | TAG   | 2371-2526 | 156    | 5   | 51  | GTG   | TAG   |
| J      | Atp6   | 2523-3194                 | 672    | 10  | 223 | ATG   | TAG   | 2532-3200 | 669    | 7   | 222 | ATG   | TAA   |
| J      | Cox3   | 3205-3987                 | 783    | 6   | 260 | ATG   | TAG   | 3208-3990 | 783    | 11  | 260 | ATG   | TAA   |
| J      | tmG    | 3994-4049                 | 56     | -1  |     |       |       | TCC   | 4002-4058 | 57    | -2  |     |       |
| J      | Nad3   | 4049-4393                 | 345    | 8   | 114 | TTG   | TAA   | 4057-4401 | 345    | -1  | 114 | TTG   | TAA   |
| J      | tmR    | 4402-4447                 | 46     | 13  |     |       |       | TCG   | 4401-4448 | 48    | 9   |     |       |
| J      | tmM    | 4461-4512                 | 52     | 3   |     | CAT   |       | 4458-4509 | 52    | 7   |     | CAT   |
| J      | tmS2   | 4516-4565                 | 50     | -2  |     | TGA   | 4517-4571 | 55    | -2  |     | TGA   |
| N      | tmC    | 4564-4615                 | 52     | 7   |     | GCA   | 4570-4621 | 52    | 12  |     | GCA   |
| J      | tmP    | 4623-4677                 | 55     | 3   |     | TGG   | 4634-4688 | 55    | 2   |     | TGG   |
| J      | tmY    | 4681-4734                 | 54     | -6  |     | GTA   | 4691-4745 | 55    | -4  |     | GTA   |
| J      | tmK    | 4729-4789                 | 61     | 3   |     | TTT   | 4742-4802 | 61    | 2   |     | TTT   |
| J      | tmN    | 4793-4846                 | 54     | -1  |     | GTT   | 4804-4860 | 57    | 0   |     | GTT   |
| J      | mS     | 4846-5504                 | 659    | -1  |     |       | 4861-5527 | 667   | -1  |     |       |
| J      | tmV    | 5504-5553                 | 50     | -1  |     | TAC   | 5527-5576 | 50    | 0   |     | TAC   |
| J      | mL     | 5553-6594                 | 1042   | 0   |     |      | 5577-6602 | 1026  | 1   |     |      |
| J      | tmW    | 6595-6650                 | 56     | 69  |     | TCA   | 6604-6660 | 57    | 92  |     | TCA   |
| N      | nad1   | 6720-7610                 | 891    | 37  | 296 | ATG   | TAA   | 6753-7688 | 936   | 0   | 311 | ATG   | TAA   |
| N      | nad6   | 7648-8082                 | 435    | -1  | 144 | ATG   | TAA   | 7689-8123 | 435   | -1  | 144 | ATG   | TAA   |
| N      | tmT    | 8082-8137                 | 56     | 9   |     | TGT   | 8123-8176 | 54    | 8   |     | TGT   |
| J      | nad4   | 8147-8401                 | 255    | 1   | 84  | ATG   | TAA   | 8185-8439 | 255   | 1   | 84  | ATG   | TAA   |
| J      | nad5   | 8403-9687                 | 1285   | 0   | 428 | ATG   | T     | 8441-9725 | 1285  | 0   | 428 | ATG   | T     |
| J      | tmH    | 9688-9744                 | 57     | -1  |     | GTG   | 9726-9781 | 56    | -1  |     | GTG   |
| J      | nad9   | 9744-11373                | 1630   | 0   | 543 | ATT   | T     | 9781-11416 | 1636  | -1  | 545 | ATT   | T     |
| J      | tmF    | 11374-11429               | 56     | 0   |     | GAA   | 11416-11471 | 56    | 0   |     | GAA   |
| J      | D-loop | 11430-12242               | 813    | 0   |     |   | 11472-12332 | 861   | 0   |     |   |
| N      | tmS1   | 12243-12292               | 50     | 15  |     | TCT   | 12333-12382 | 50    | 0   |     | TCT   |
| N      | tmQ    | 12308-12360               | 53     | -2  |     | TGG   | 12383-12436 | 54    | 2   |     | TGG   |
| N      | tmL    | 12359-12413               | 55     | -1  |     | GAT   | 12439-12493 | 55    | -1  |     | GAT   |
| N      | nad2   | 12413-13337               | 925    | -1  | 308 | ATA   | T     | 12493-13417 | 925   | -2  | 308 | ATT   | T     |
| N      | tmE    | 13337-13389               | 53     | 0   |     | TTC   | 13416-13473 | 58    | -1  |     | TTC   |
| N      | cob    | 13390-14493               | 1104   | 2   | 367 | ATG   | TAA   | 13473-14574 | 1102  | 2   | 367 | ATG   | T     |
| N      | tmL1   | 14496-14550               | 55     | 9   |     | TAG   | 14577-14631 | 55    | 8   |     | TAG   |
| J      | tmA    | 14560-14607               | 48     | -2  |     | TGC   | 14640-14690 | 51    | -4  |     | TGC   |
| J      | tmL2   | 14606-14663,1             | 59     | -1  |     | TAA   | 14687-14741 | 55    | -1  |     | TAA   |

Table 2. Distribution of NCRs in the astigmatid mites mitochondrial genomes.
| Species                   | Conservative non-coding region | Nonconservative noncoding region |
|---------------------------|-------------------------------|----------------------------------|
|                           | length | position | AT(%) | length | position | AT(%) |
| *Lepidoglyphus destructor*| 813    | 69       | 68.4% | 69     | tmWnad1  | 55.1% |
| *Gohieria fusca*          | 861    | 92       | 70.7% | 92     | tmWnad1  | 63.0% |
| *Acalvolia sp*            | 753    | 273      | 76.6% | 273    | tmWnad1  | 78.8% |
| *Carpoglyphus lactis*     | 303    | 66       | 88.4% | 66     | tmWnad1  | 72.7% |
| *Aleuroglyphus ovatus*    | 378    | 57       | 82.5% | 57     | tmWnad1  | 75.4% |
| *Caloglyphus berlesei*    | 341    | 53       | 89.4% | 53     | tmWnad1  | 71.7% |
| *Rhizoglyphus robini*     | 319    | 53       | 87.1% | 53     | tmWnad1  | 71.7% |
| *Tyrophagus putrescentiae*| 245    | 53       | 93.1% | 53     | tmWnad1  | 67.9% |
| *Dermatophagoides farinae*| 410    | 46       | 90.5% | 46     | tmWnad1  | 65.2% |
| *Dermatophagoides pteronyssinus* | 286   | 44       | 91.7% | 44     | tmWnad1  | 65.9% |
| *Ardeacarus ardeae*       | 214    | 29       | 91.6% | 29     | tmWnad1  | 89.7% |
| *Psoroptes cuniculi*      | 382    | 62       | 89.8% | 62     | tmWnad1  | 72.6% |
| *Sarcoptes scabiei*       | 76     | 62       | 98.7% | 62     | tmWnad1  | 79.0% |
| *Trouessartia rubecula*   | 301    | 56       | 84.7% | 56     | tmWnad1  | 73.5% |
| *Tyrophagus longior*      |        | 49       |       |       | tmWnad1  | 73.5% |
| *Histiosoma blomquisti*   | 624    | 50       | 68.6% | 50     | mSrmL    | 90.0% |
| *Histiosoma feroniarum*   | 146    | 143      | 73.9% | 143    | Nad4-trnH | 81.1% |

*Note:* Non-coding regions marked with a star are assumed to be putative control regions. The data for *Trouessartia rubecula, Histiosoma blomquisti,* and *H. feroniarum* in the table are after our revision.

**Table 3.** The alignment of nucleotide sequences of four mitochondrial tRNA genes (*trnS2, trnW, trnF,* and *trnI*) in five reported astigmatid species in four different superfamilies.
Table 4. The alignment of nucleotide sequences of two mitochondrial tRNA genes (tmA and tmV) in six species: Conserved nucleotides are shaded in gray across these five mites.

| Gene | Species | AA-arm | D-arm | AC-arm | V-loop | T-arm | AA-arm |
|------|---------|--------|-------|--------|--------|-------|--------|
| tmA  | Ao      | TGGATGG| AGAGAA | CTTTTTTGTGAGAAAG | TTTTT | TTAGAA | CAAGAGTT |
|      |Cb      | TGGATGG| AAGATA | CTTTTTTGTGAGAAAG | TTTTT | TTAGAA | CAAGAGTT |
|      |Rr      | TGGATGG| AAGATA | CTTTTTTGTGAGAAAG | TTTTT | TTAGAA | CAAGAGTT |
|      |Df      | TGGGCA| AAGAGGTTG | ACTTTTTTGAGAAAG | TTTTT | TTAGAA | CAAGAGTT |
|      |AA      | TGGGCA| AGAGAA | GTTTTTTGTGAGAAAG | TTTTT | TTAGAA | CAAGAGTT |
|      |TR      | TGGGCA| AGAGAA | GTTTTTTGTGAGAAAG | TTTTT | TTAGAA | CAAGAGTT |
| tmV  |Ao      | TGGAGCT| AGAGAA | TTTTGTGTACAAAAA | ATTTT | TTAGTA | TAAGTT |
|      |Cb      | TGGAGCT| AGAGAA | TTTTGTGTACAAAAA | ATTTT | TTAGTA | TAAGTT |
|      |Rr      | TGGAGCT| AGAGAA | TTTTGTGTACAAAAA | ATTTT | TTAGTA | TAAGTT |
|      |Df      | TGGAGCT| AGAGAA | TTTTGTGTACAAAAA | ATTTT | TTAGTA | TAAGTT |
|      |AA      | TGGAGCT| AGAGAA | TTTTGTGTACAAAAA | ATTTT | TTAGTA | TAAGTT |
|      |TR      | TGGAGCT| AGAGAA | TTTTGTGTACAAAAA | ATTTT | TTAGTA | TAAGTT |

Note: Nucleotides that pair at the arms (acceptor arm, D arm, anticodon arm, and T arm) are underlined. Sequences of anticodons are in boldface.

*Alleuroglyphus ovatus* (Ao), *Caloglyphus berlesei* (Cb), *Rhizoglyphus robinii* (Rr), *Dermatophagoides farinae* (Df), *Ardeacarus ardeae* (AA), and *Trouessartia rubecula* (TR) in our study.
Table 5 The primers used for the amplification of *Gohieria fusca* and *Lepidoglyphus destructor* mitochondrial genomes, with the size of each fragment that was amplified in bp.

| Specimen     | Primer name | Original sequence | Size fragment (bp) |
|--------------|-------------|-------------------|--------------------|
| *G. fusca*   | Gf-CR-F     | GCTTTGAAGGGCTAAAGGT | 852                |
|              | Gf-CR-R     | TCTAGGTGCAACACAAAG |                    |
| *L. destructor* | Ld-CR-F    | TGAAGGGGCTAAAGGTGCC | 962                |
|              | Ld-CR-R     | ACCGGATAGGGTACACCTTG |                |

Table 6 Mitochondrial genomes employed in this study.

| Cohort                  | Superfamily     | Family        | Species                      | GenBank ID  | Size(bp) | Reference                     |
|-------------------------|-----------------|---------------|------------------------------|-------------|----------|-------------------------------|
| Mixonomatides           | Phthiracoidea   | Phthiracaridae| *Steganacarus magnus*        | EU935607    | 13818    | Domes *et al.* (2008) [57]    |
| Brachypylina             | Oripodoidea     | Scheloribatidae| *Paraleius leontonychus*     | LT984407    | 14186    | Schäffer *et al.* (2018) [31]|
| Astigmatina             | Acaroidea       | Acaridae      | *Aleuroglyphus ovatus*       | KC700022    | 14328    | Sun *et al.* (2014b) [25]     |
|                         |                 |               | *Caloglyphus berlesei*       | KF499016    | 14273    | Sun *et al.* (2014a) [24]     |
|                         |                 |               | *Rhizoglyphus robini*        | MF596168    | 14244    | Xue *et al.* (2018) [18]      |
|                         |                 |               | *Tyrophagus longior*         | KR869095    | 13271    | Yang & Li (2016) [20]         |
|                         |                 |               | *Tyrophagus putrescentiae*   | MK_393792   | 14156    | Fang *et al.* (2020) [22]     |
| Astigmatina             | Acaroidea       | Acaridae      | *Aleuroglyphus ovatus*       | KC700022    | 14328    | Sun *et al.* (2014b) [25]     |
|                         |                 |               | *Caloglyphus berlesei*       | KF499016    | 14273    | Sun *et al.* (2014a) [24]     |
|                         |                 |               | *Caloglyphus berlesei*       | KF499016    | 14273    | Sun *et al.* (2014a) [24]     |
|                         |                 |               | *Rhizoglyphus robini*        | MF596168    | 14244    | Xue *et al.* (2018) [18]      |
|                         |                 |               | *Tyrophagus longior*         | KR869095    | 13271    | Yang & Li (2016) [20]         |
| Astigmatina             | Acaroidea       | Acaridae      | *Aleuroglyphus ovatus*       | KC700022    | 14328    | Sun *et al.* (2014b) [25]     |
| Analgoidea              | Pyroglyphidae   |               | *Dermatophagoides farinae*   | NC_013184   | 14266    | Klimov & O'Connor (2009) [17]|
|                        | *Dermatophagoides pteronyssinus* |     | EU884425                        | 14203        |          | Demauw *et al.* (2009) [16]    |
| Trouessartiidae         | Trouessartia rubedula |              | MH208456                       | 14125        |          | Esteban *et al.* (2018) [23]  |
| Hemisarcoptoidea        | Winterschmidtii | Acalvolia sp.  | MH921997                       | 14711        |          | Li & Xue (2019) [11]          |
| Carpoglyphidae          | Carpoglyphus lactis |             | MN073839                       | 14060        |          | Zhao & Li (2020) [58]         |
| Histiostomatoidea       | Histiostomatidae| *Histiostoma blomquisti* | KX452726                       | 15892        |          | Lee & Wang (2016) [27]        |
|                        |                 | *Histiostoma feroniarum* | MF596167                       | 13896        |          | Xue *et al.* (2018) [18]      |
| Pterolichoidea          | Pterolichidae   | Ardeacarus ardea | KY352304                       | 14069        |          | Han & Min (2017) [59]         |
| Sarcoptoidea            | Psoroptidae     | Psoroptes cuniculi | KJ957822                       | 14247        |          | Gu *et al.* (2014) [26]       |
|                        | Sarcoptidae     | Sarcoptes scabiei | CM003133                       | 13667        |          | Mofiz *et al.* (2016) [60]    |
| Glycyphagoidea          | Glycyphagidae   | Lepidoglyphus destructor | MT075728                       | 14663        |          | This study                    |
|                         | *Gohieria fusca*|                | MN608156                       | 14741        |          | This study                    |
|                         |                 |                |                               |             |          |                               |