Mitogenomics and phylogenetics of twelve species of African Saturniidae (Lepidoptera)

Zwannda Nethavhani1, Rieze Straeuli1, Kayleigh Hiscock1, Ruan Veldtman2,3, Andrew Morton4, Rolf G. Oberprieler5 and Barbara van Asch1

1 Department of Genetics, University of Stellenbosch, Stellenbosch, Western Cape, South Africa
2 Department of Conservation Ecology and Entomology, University of Stellenbosch, Stellenbosch, Western Cape, South Africa
3 Kirstenbosch Research Centre, South African National Biodiversity Institute, Cape Town, Western Cape, South Africa
4 Unaffiliated, Cape Town, Western Cape, South Africa
5 Australian National Insect Collection, Commonwealth Scientific and Industrial Research Organisation, Canberra, Australia

ABSTRACT

African Saturniidae (Lepidoptera) include numerous species consumed at the caterpillar stage throughout the continent, and their importance to local communities as a source of nutrition and seasonal income cannot be overestimated. However, baseline genetic data with utility for the characterization of their diversity, phylogeography and phylogenetic relationships have remained scarce compared to their Asian counterparts. To bridge this gap, we sequenced the mitochondrial genomes of 12 species found in southern Africa for comparative mitogenomics and phylogenetic reconstruction of the family, including the first representatives of the tribes Eochroini and Micragonini. Mitochondrial gene content and organization were conserved across all Saturniidae included in the analyses. The phylogenetic positions of the 12 species were assessed in the context of publicly available mitogenomes using Bayesian inference and maximum likelihood (ML) methods. The monophyly of the tribes Saturniini, Attacini, Bunaeni and Micragonini, the sister relationship between Saturniini and Attacini, and the placement of Eochroa trimenii and Rhodinia fugax in the tribes Eochroini and Attacini, respectively, were strongly supported. These results contribute to significantly expanding genetic data available for African Saturniidae and allow for the development of new mitochondrial markers in future studies.

INTRODUCTION

The family Saturniidae (Lepidoptera) comprises approximately 3,400 species in 180 genera distributed in a diversity of terrestrial habitats worldwide (Kitching et al., 2018). The African fauna of the family comprises about 400 species in 50 genera, classified in six tribes (Oberprieler, Morton & Noort, 2021). The pupal cocoons of several Asian saturniids are used for silk production, e.g., Antheraea assamensis, Antheraea pernyi,
Eriogyna pyretorum, Saturnia japonica and Samia cynthia ricini (Jiang et al., 2009; Kim et al., 2012; Singh et al., 2017; Kim et al., 2018; Liu et al., 2021). Sericulture is not generally practiced in Africa and the potential of African silk-producing species such as Argema mimosae (Saturnini) and Epiphora bauhiniae (Attacini) is largely untapped, although traditional uses are documented (Peigler & Oberprieler, 2017). Nevertheless, the larvae of at least 50 saturniid species are consumed in southern, central and western Africa, where they represent important sources of nutrition and income to rural communities (Jongema, 2015; Kelemu et al., 2015; Malaisse, Mabossy-Mobouna & Latham, 2017; Hlongwane, Slotow & Munyai, 2020; Sekonya, McClure & Wynberg, 2020; Hlongwane, Slotow & Munyai, 2021; Kusia et al., 2021). However, large-scale harvesting of caterpillars from the wild is a threat to populations (McGeoch, 2002), exacerbated by lack of distribution data and monitoring.

In southern Africa, the most popular edible caterpillars are those of Gonimbrasia belina and Gynanisa maja, commonly known as mopane worms. Other edible Saturniidae recorded in southern Africa include species in the tribes Bunaeani (genera Athletes, Bunaea, Bunaeoides, Bunaeopsis, Cirina, Gonimbrasia, Gynanisa, Heniocha, Imbrasia, Lobobunaea, Melanocera, Nudaurelia, Pseudobunaea and Rohaniella), Micragonini (Goodia and Micragone) and Urotini (Pseudaphelia, Urota and Usta) (Silow, 1976; Nonaka, 1996; Thomas, 2013; Jongema, 2015; Malaisse, Mabossy-Mobouna & Latham, 2017; Hlongwane, Slotow & Munyai, 2021). However, knowledge regarding the exact number of species is limited due to taxonomic instability and inconsistencies in the family Saturniidae (Kitching et al., 2018; Siozios et al., 2020). Therefore, specimens belonging to different species are often assigned the same species name and vice versa (Siozios et al., 2020). As a result, the total number of Saturniidae species found and consumed in Africa may be either under- or over-reported, with implications for implementing long-term conservation measures. Despite their importance, no formally protected areas specifically designed for the conservation of these species exist in southern Africa, or for any other edible insect for that matter (Dzeresos, 2018).

Mitogenomes are widely used for phylogenetic reconstruction in insects, and deeper taxonomic coverage and recent methodological developments have allowed for improved recovery of relationships among taxa, even those affected by compositional heterogeneity and accelerated evolutionary rates (Song et al., 2016). Mitochondrial phylogeny estimates of Saturniidae have mostly been limited to Asian species (Jiang et al., 2009; Zhang, Zhao & Zhou, 2016; Kim et al., 2017a; Kim et al., 2017b; Li et al., 2017; Saikia, Nath & Devi, 2019; Kim et al., 2009; Liu et al., 2021), and only two African species in the tribe Bunaeani (G. belina and Gy. maja) were recently included (Langley et al., 2020), representing approximately 1% of the total species recorded in Africa and 4% of all saturniid mitogenomes publicly available. Thus, assessments of phylogenetic relationships, phylogeographic structure and genetic diversity of African Saturniidae have been hampered by limited baseline genetic information.

In this study, we generated mitogenome sequences for 12 African Saturniidae species in four tribes and assessed their phylogenetic position in the family: Eochroa trimenii (Eochroini), Bunaea alcinoe, Heniocha apollonia, Heniocha dyops, Gonimbrasia tyrrhea, Nudaurelia cytherea, Nudaurelia wahlbergii (Bunaeani), Epiphora bauhiniae (Attacini),
Ludia delegorguei, Holocerina smilax, Vegetia ducalis and Vegetia grimmia (Micragonini). This novel dataset covers 24% of the total number of species occurring in South Africa, Botswana and Namibia (R. Oberprieler, pers. obs., 2022) and is a significant improvement on the present paucity of mitogenomic data for sub-Saharan Saturniidae.

MATERIALS & METHODS
Specimen collection, morphological identification, DNA extraction
Adult and caterpillar specimens were collected at several locations in South Africa and Namibia, between January and March 2020 (Fig. 1, Table 1) under permits approved by the Ezemvelo KZN Permits Office (OP408/2020), Cape Nature (CN44-59-13497) and National Commission on Research Science and Technology (AN20190911). Specimens were euthanized by freezing within a few hours of collection, whenever field conditions allowed. Species identification of adults and caterpillars was made by R. Oberprieler based on photographs of reference specimens using current literature (Pinhey, 1972; Oberprieler, 1995; Staude et al., 2016; Staude et al., 2020). Legs from each adult and caterpillar were excised and stored in 100% ethanol at −20 °C until DNA extraction. Total DNA was extracted from one leg from each adult and caterpillar using a standard phenol-chloroform protocol (Sambrook & Russell, 2012).

DNA barcoding
The standard COI barcoding region was amplified using the primer pair LepF/LepR in a total reaction volume of 5 µL containing 2.5 µL of Qiagen Multiplex PCR Kit (QIAGEN), 0.5 µL of each primer, 0.5 µL of Milli-Q water and 1 µL of template DNA. PCR amplifications were performed with initial denaturation at 95 °C for 15 min; 35 cycles of 94 °C for 30 s, 50 °C for 90 s; followed by a final extension at 72 °C for 10 min. Sequencing reactions were performed using LepR with the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Waltham, MA, USA), at the Central Analytical Facilities of Stellenbosch University. The sequences were queried against the BOLD Systems (https://www.boldsystems.org/) and GenBank (https://blast.ncbi.nlm.nih.gov/) for confirmation of species identification on February 23, 2022.

Sequencing, assembly, and annotation of mitogenomes
Specimens were individually sequenced using the Ion Torrent™ S5™ platform (ThermoFisher Scientific, Waltham, MA, USA) available at the Central Analytical Facilities of Stellenbosch University, South Africa. Sequence libraries were prepared using the Ion Xpress™ Plus gDNA Fragment Library Kit (ThermoFisher Scientific, Waltham, MA, USA), according to the protocol MAN0009847, REV J.0. Libraries were pooled and sequenced using the Ion 540™ Chef Kit (ThermoFisher Scientific, Waltham, MA, USA). The Ion Torrent reads were quality trimmed with a 30-base sliding-window at an average threshold of Q16. The remaining sequencing reads were then filtered for read length, where any reads shorter than 25 bases were removed from the data. Reference-based assembly of the next-generation sequencing (NGS) reads was performed using mitogenomes of the closest Saturniidae relatives available on GenBank (Table S1). NGS
Figure 1  Representative specimens of 12 African Saturniidae species. Larva and adult specimens used in this study for new mitogenomic sequencing and phylogenetic analyses or collected from the same area. (A) *Nudaurelia wahlbergii* (Boisduval, 1847), (B) *Nudaurelia cytherea* (Fabricius, 1775), (C) *Bunaea alcinoe* (Stroll, 1780), (D) *Gonimbrasia tyrhrrea* (Cramer, 1776), (E) *Heniocha apollonia* (Cramer, 1779), (F) *Heniocha dyops* (Maassen, 1872), (G) *Vegetia ducalis* (Jordan, 1922), (H) *Vegetia grimmia* (Geyer, 1832), (C) *Holocerina smilax* (Westwood, 1849), (D) *Ludia delegorguei* (Boisduval, 1847), (I) *Eochroa trimenii* (C. & R. Felder, 1874), and (L) *Epiphora bauhiniae* (Guérin-Méneville, 1832).
| Species                  | Tribe        | Specimen | Life stage | Collection date | Country     | Region    | GPS       | Edible | GenBank |
|-------------------------|--------------|----------|------------|-----------------|-------------|-----------|-----------|--------|---------|
| *Bunaea alcinoe* (Stoll, 1780) | Bunaenini    | BA1.6    | Adult      | 2019/10/11      | South Africa| Gauteng    | −25.7479  | 28.2293| OL912807|
| *Eochroa trimenii* (C. & R. Felder, 1874) | Eochroini    | ET6      | Larva      | 2020/08/01      | South Africa| Western Cape| −34.0771  | 18.4951| Not reported |
| *Epiphora bauhiniae* (Guérin-Méneville, 1832) | Attacini     | GM5.2    | Larva      | 2020/02/26      | Namibia     | Okonjima   | −20.7579  | 16.7898| OL912809|
| *Gonimbrasia tyrrea* (Cramer, 1776) | Bunaenini    | GT1.1    | Adult      | 2020/05/25      | South Africa| Western Cape| −32.7681  | 18.1607| OL912810|
| *Heniocha apollonia* (Cramer, 1779) | Bunaenini    | HA1      | Adult      | 2020/10/04      | South Africa| Western Cape| −34.0367  | 23.0495| OL912811|
| *Heniocha dyops* (Maassen, 1872) | Bunaenini    | GM8.2    | Adult      | 2020/03/15      | Namibia     | Windhoek   | −22.8577  | 17.1089| OL912812|
| *Holocerina smilax* (Westwood, 1849) | Micragonini  | HS5      | Larva      | 2020/01/25      | South Africa| KwaZulu Natal| −26.15    | 27.9637| Not reported |
| *Ludia delegorguei* (Boisduval, 1847) | Micragonini  | LD1      | Larva      | 2020/12/12      | South Africa| KwaZulu Natal| −29.8455  | 30.9197| Not reported |
| *Nudaurelia cytherea* (Fabricius, 1775) | Bunaenini    | NC1.2    | Larva      | 2020/10/12      | South Africa| Western Cape| −34.3322  | 18.9878| OL912953|
| *Nudaurelia wahibergii* (Boisduval, 1847) | Bunaenini    | NW1      | Larva      | 2020/12/11      | South Africa| KwaZulu Natal| −29.8244  | 30.9263| OL912817|
| *Vegetia ducalis* (Jordan, 1922) | Micragonini  | VD1      | Adult      | 2020/05/22      | South Africa| Western Cape| −33.4201  | 18.4005| Not reported |
| *Vegetia grimmia* (Geyer, 1832) | Micragonini  | VG2      | Adult      | 2020/03/12      | South Africa| Western Cape| −34.3053  | 18.4619| Not reported |

reads were assembled and mapped to the reference sequences on Geneious Prime v2021.1.1 ([https://www.geneious.com/](https://www.geneious.com/)) using the “Map to reference” function, under the low to medium sensitivity option and fine-tuning up to five iterations. The open reading frames of the expected 13 mitochondrial protein-coding genes (PCGs) were identified with Geneious Prime using the invertebrate mitochondrial genetic code. Transfer RNA (tRNA) genes were located with ARWEN software ([http://130.235.244.92/ARWEN/](http://130.235.244.92/ARWEN/)) ([Laslett & Canbäck, 2008](http://130.235.244.92/ARWEN/)), under the default composite metazoan mitochondrial genetic code. The two ribosomal RNA (rRNA) genes and the non-coding AT-rich region were annotated by manual comparison to other Saturniidae mitogenomes available on GenBank. The raw sequence data were deposited on GenBank (PRJNA796275), as well as the 12 assembled and annotated mitogenomes (OL912807 to OL912817 and OL912953).

**Mitogenomic and phylogenetic analyses**

Mitogenome nucleotide composition and biases [(AT-skew = (A−T)/(A+T) and GC-skew = (G−C)/(G + C)] were calculated in Geneious Prime. Gene overlapping regions and intergenic spaces were counted manually. Non-synonymous (Ka) and synonymous (Ks) substitution rates in PCGs were calculated in DnaSP6 ([Rozas et al., 2017](http://130.235.244.92/ARWEN/)), and relative synonymous codon usage was calculated in MEGA X ([Kumar et al., 2018](http://130.235.244.92/ARWEN/)), with the invertebrate mitochondrial genetic code in both cases. The presence of tandem repeat
elements in the AT-rich region was searched using the Repeat Finder plug-in available on Geneious Prime.

The phylogenetic positions of the 12 African Saturniidae species were inferred in the context of other sequences available on GenBank for the family as of October 2021 (Table S2). The final dataset \(n = 44\) included 35 species in 19 genera and five tribes, with Bombycidae (\(Bombyx\) mori, \(Bombyx\) mandarina and \(Rondotia\) menciana), Sphingidae (\(Manduca\) sexta and \(Sphinx\) morio) and non-Bombycoidea (\(Biston\) panterinaria, \(Phthonandria\) atrilineata, \(Protantigius\) superans and \(Spindasis\) takanonis) as outgroups. Phylogenetic analyses were performed using the 13 PCGs (all codon positions). Individual PCG sequences were extracted from each mitogenome and stop codons were removed manually. Translation alignments were performed separately for each PCG using the MAFFT algorithm in Geneious Prime and then concatenated in a single alignment for each specimen. Poorly aligned regions and alignment gaps were removed using GBlocks v0.91b (Castresana, 2000). Two Bayesian inference (BI) and one maximum likelihood (ML) methods were used for phylogenetic reconstruction. The best partitioning scheme and model for MrBayes v3.1.2 (Huelsenbeck & Ronquist, 2001) and ML were determined using the edge-linked greedy strategy (Lanfear et al., 2012) in PartitionFinder v2.1.1 (Lanfear et al., 2017) built-in Anaconda programme (https://repo.anaconda.com/), and GTR+I+G was considered the best model. BI in MrBayes was performed under the GTR+I+G nucleotide substitution model selected with PartitionFinder. Analyses were performed as follows: two independent runs of four heat chains, ten million generations run simultaneously, resampling every 1000 generations. The first 25% of trees were discarded as burn-in, and the decision criterion for the convergence of the two runs was set as an average split frequency of \(\leq 0.01\). BI analyses were also performed under the site-heterogeneous mixture model CAT+GTR in PhyloBayes MPI in XSEDE v1.8c (Lartillot et al., 2013) to minimize the effect of mitochondrial compositional heterogeneity on phylogenetic reconstructions (Lartillot, Brinkmann & Philippe, 2007; Cai et al., 2020). Constant sites were removed from the alignment, and the minimum number of cycles was set to 30,000 with the burn-in set to 1000. The “maxdiff” was set to 0.3, and the minimum effective size was set to 50. Nodal support in the MrBayes and PhyloBayes trees was estimated as Bayesian posterior probabilities (BPP). ML analyses were performed in IQ-TREE v1.6.12 (Nguyen et al., 2015) under the GTR+I+G substitution model selected by PartitionFinder. Branch supports were determined using 1000 replicates for both the ultrafast bootstrapping (UFBoot) and the SH-aLRT branch test (Guindon et al., 2010; Hoang et al., 2018). MrBayes, PhyloBayes and IQ-TREE analyses were run on the CIPRES Science Gateway Portal (Miller, Pfeiffer & Schwartz, 2010). The final trees were drawn using FigTree v1.4.4 (https://bio.tools/figtree).

RESULTS AND DISCUSSION

Mitochondrial phylogeny estimates of Saturniidae have been largely limited to Asian species, and the only mitogenomes publicly available for African species prior to this study were those of \(G.\) belina and \(Gy.\) maja, both in the tribe Bunaeini (Langley et al., 2020). This work adds the mitogenomes for 12 species of the tribes Bunaeini, Micragonini,
Eochroini and Attacini and significantly expands the genetic resources available for African Saturniidae, thus allowing further insights into the mitochondrial phylogeny of the family.

DNA barcoding

The sequence queries against BOLD Systems confirmed the morphological identification for all specimens used for mitogenome sequencing with sequence similarity between 99.44 and 100% (Table S3). The BLASTn queries against GenBank produced no matches with high identity (>98%) except for B. alcinoe and N. cytherea due to the absence of sequences for these species on the database.

Comparative mitogenomics of Saturniidae

The Ion Torrent runs generated between 11,321,643 (V. ducalis) and 21,284,325 (E. bauhiniae) reads, with an average size of 184 bp. Total mapped reads ranged from 16,637 (H. smilax) to 191,295 (H. dyops). Average sequence coverage varied between 197 × (H. smilax) and 1810 × (H. dyops) (Table 2). The total length of the new mitogenomes ranged from 15,218 bp (V. ducalis) to 15,363 bp (N. cytherea), in line with those of other Saturniidae (Kim et al., 2017a; Kim et al., 2017b; Langley et al., 2020; Liu et al., 2021; Chen et al., 2021). All mitogenomes generated in this study are identical in gene content and organization and include the 13 PCGs, two rRNA genes and 22 tRNA genes typical of Metazoa. Nine PCGs (ATP6, ATP8, CYTB, COI, COII, COIII, ND2, ND3 and ND6) and 14 tRNAs are located on the major (J) strand, and four PCGs (ND1, ND4, ND4L and ND5), eight tRNAs and the two rRNAs are located on the minor (N) strand (Table S4). Gene order is conserved across all species and identical to that of other Saturniidae, with the reciprocal translocation of tRNA_{Met} and tRNA_{Gln} relative to tRNA_{Ile} (M-I-Q) as the only difference to the hypothetical ancestral arrangement for the non-Ditrysia lineage Hepialoidea of Lepidoptera (I-Q-M) (Cao et al., 2012) (Fig. 2).

All tRNAs have the typical cloverleaf structure except for tRNA_{Ser1} in B. alcinoe, E. bauhiniae, G. belina, G. tyrrhea, G. maja, H. apollonia, H. dyops, H. smilax, L. delegorguei, V. ducalis and V. grimmia, in that the dihydrouridine (DHU) arm is absent, as occurs in many Metazoa (Cameron, 2014) (Fig. 3). In contrast, the DHU arm of tRNA_{Ser1} is present in E. trimenii, N. cytherea and N. wahlbergii. The length of the tRNAs ranges from 57 bp (tRNA_{Phe}) in N. wahlbergii to 74 bp (tRNA_{Trp}) in H. apollonia. The location and average size of 16S rRNA (1,357 bp; between tRNA_{Leu1} and tRNA_{Val}) and 12S rRNA (760 bp; between tRNA_{Val} and AT-rich region) across the 12 species is in line with the average size and position of the two genes in other Saturniidae. No tandem repeat elements were detected in the AT-rich regions of the 12 species.

The new sequences have the high AT content characteristic of insect mitogenomes, ranging from 78.80% in H. apollonia to 81.00% in E. trimenii (Table S5). The AT content of the total PCGs follows the same trend and varies from 77.40% in H. apollonia to 79.90% in E. trimenii. Average AT content across the 12 species is highest in ATP8 (92.70%) and lowest in COI (69.30%) (Fig. 4), in line with previous reports on Saturniidae (Jiang et al., 2009; Singh et al., 2017; Chen et al., 2021). AT-skew is negative for PCGs on the J-strand across all species and varies from −0.22 in ND3 to −0.02 in ATP8. All four PCGs on the...
Table 2  Results of next-generation sequencing for the assembly and annotation of the complete mitochondrial genomes of 12 African Saturniidae (Lepidoptera) species.

| Specimen | Number of reads | Average read length (bp) | Number of rmapped reads | Average coverage | Sequence length (bp) |
|----------|-----------------|--------------------------|-------------------------|-----------------|---------------------|
| Bunaea alcinoe BA1.6 | 15,634,521 | 187 | 49,550 | 582 × | 15,305 |
| Eochroa trimenii ET6 | 16,515,662 | 182 | 62,423 | 718 × | 15,254 |
| Epiphora bauhiniae GM5.2 | 21,284,25 | 173 | 108,486 | 1147 × | 15,259 |
| Gonimbrasia tyrrea GT1.1 | 17,135,076 | 177 | 132,321 | 1412 × | 15,296 |
| Heniocha apollonia HA1 | 16,037,132 | 186 | 137,494 | 1397 × | 15,270 |
| Heniocha dyops GM8.2 | 15,407,906 | 173 | 191,295 | 1810 × | 15,306 |
| Holocerina smilax HS5 | 19,181,464 | 192 | 16,637 | 197 × | 15,220 |
| Ludia delegorguei LD1 | 14,679,181 | 196 | 26,826 | 320 × | 15,238 |
| Nudaurelia cytherea GM5.2 | 19,034,516 | 174 | 33,793 | 349 × | 15,363 |
| Nudaurelia wahlbergi NW1 | 13,837,894 | 188 | 30,529 | 333 × | 15,287 |
| Vegetia ducalis VD1 | 11,321,643 | 195 | 42,382 | 509 × | 15,217 |
| Vegetia grimmia VG2 | 14 153 225 | 193 | 88 880 | 1048 × | 15,251 |

Figure 2  Mitochondrial gene organization. Linear map of the complete mitochondrial genomes of (A) African Saturniidae and (B) hypothetical ancestral of the non-Ditrysian lineage Hepialoidea of Lepidoptera.

N-strand have a positive AT-skew, ranging from 0.14 in ND5 to 0.24 in ND1 and ND4L. Additionally, most PCGs have a slightly negative GC-skew, ranging from −0.69 (ATP8) to −0.02 (COI), except for COIII = 0.00 in L. delegorguei.

All 13 PCGs initiate with standard ATN start codons except COI and COII, which start with the alternative codons CGA and GTG, respectively. CGA as the start codon for COI has been reported to be highly conserved in Lepidoptera (Margam et al., 2011), and GTG was reported as the start codon for COII in A. assamensis, E. pyretorum, S. boisduvalii and S. ricini (Jiang et al., 2009; Kim et al., 2012; Singh et al., 2017; Kim et al., 2017a; Kim et al., 2017b). PCGs terminate with complete TAA or TAG codons or with incomplete TA- or T-, which are presumed to be completed by posttranscriptional modifications such as polyadenylation (Salinas-Giegé, Giegé & Giegé, 2015) (Fig. 5, Table S6). The most frequent amino-acid codons are Leu, Ile, Phe and Ser across all species, whereas Cys, Asp, Arg, Glu and Gln are rare (Fig. 6A). The most frequently used codons are AT-rich, and this feature is reflected in the total mitogenome nucleotide bias towards A and T. ATP8 has the largest
Figure 3  Structure of tRNA-Ser in 12 African Saturniidae species. Predicted structure of tRNA<sub>Ser</sub> in the complete mitochondrial genomes of 12 new and two previously sequenced African Saturniidae species. Inferred canonical Watson-Crick bonds are represented by lines, and non-canonical bonds are represented by dots.

number of different codons compared to all other genes across the 12 species (Fig. 6B). Relative synonymous codon usage (RSCU) is higher than 1.0 for all codons and highest for Leu1 (Fig. 7). Average Ka/Ks was found to be less than 1.0 in all PCGs across all species,
indicating purifying or stabilizing selection, and to be highest for ATP8 (Ka/Ks = 0.31) (Fig. 8).

**Phylogenetic position of African Saturniidae in the family**

The phylogeny of the family Saturniidae has been reconstructed using morphological, nuclear and mitochondrial sequence data. Regier et al. (2008) analysed four protein-coding nuclear genes using parsimony and ML methods and included five tribes present in Africa (Attacini, Bunaenini, Micragonini, Saturniini and Urotini). The recovered tribal structure strongly supported Attacini as sister group of Saturniini, which together formed the sister group of Urotini, Bunaenini and Micragonini, although with Urotini paraphyletic in regard to Bunaenini and Micragonini. More recently, the phylogeny of Saturniidae was reconstructed using a phylomorphology approach based on anchored-hybrid-enriched (AHE) loci followed by geometric morphometrics of hindwings (Rubin et al., 2018). The
Figure 5  Usage of start and stop codons in Saturniidae mitogenomes. Usage of start and stop codons of 13 protein-coding genes in 14 mitogenomes of African Saturniidae species.

Figure 6  Codon frequencies in Saturniidae mitogenomes. (A) Mean codon frequencies and (B) standard deviation in the mitochondrial protein-coding genes of 14 African Saturniidae. Vertical axis: Protein-coding genes; Horizontal axis: Single-letter amino acid codons.

tree also supported a sister relationship between Attacini and Saturniini, the African tribes Bunaeini, Urotini and Micragonini together forming a monophylum and the sister group of Attacini+Saturniini, with Urotini again paraphyletic in regard to Bunaeini. In addition to phylogeny estimates based on nuclear genes and phylomorphology, the relationships among Saturniidae have been reconstructed using mitogenomic data (Sima et al., 2013; Kim et al., 2018; He, Wang & Chen, 2017; Zhong et al., 2017; Langley et al., 2020; Chen et al., 2021; Liu et al., 2020; Liu et al., 2021; Zhao et al., 2021). Although these studies focused on Asian species in the tribes Attacini and Saturniini, reciprocal monophyly and sister relationships were largely consistent, except for the positions of Cricula trifenestrata and Neoris haraldi, which were not consistently recovered, likely due to low sequence quality. The addition of the African tribe Bunaeini to the mitochondrial phylogeny did not challenge the sister relationship between Attacini and Saturniini, which together formed the sister group of Bunaeini (Langley et al., 2020; Chen et al., 2021), in agreement with
the non-mitochondrial phylogeny estimates (Regier et al., 2008; Rubin et al., 2018). Thus, mitochondrial phylogeny estimates of Saturniidae have included only the tribes Attacini, Bunaeeini and Saturniini, and African species were severely under-represented compared to Asian species (Langley et al., 2020; Chen et al., 2021). To contribute to the filling of this gap, we assessed the phylogenetic position of 12 African species of Saturniidae including the first representatives of the tribes Eochroini and Micragonini. Due to the suspected low quality of the mitogenomes of *C. trifenestrata* and *N. haraldi*, we performed phylogeny estimates with and without these sequences.

**Phylogenetic trees excluding *C. trifenestrata* and *N. haraldi***

The ML and MrBayes trees excluding *C. trifenestrata* and *N. haraldi* (Fig. 9) displayed the same phylogenetic structure, recovering Bunaeeini as a basal lineage, a sister relationship between Micragonini and Eochroini and the sister relationship between Saturniini and Attacini as found in previous analyses based on nuclear genes (Regier et al., 2008), phylomorphology (Rubin et al., 2018) and mitogenome data (Langley et al., 2020; Chen et al., 2021) (Fig. 9). In contrast, the PhyloBayes tree (Fig. 10) recovered a topology different from those of ML and MrBayes and of previous phylogenetic hypotheses in that Saturniini appear non-monophyletic, and several nodes have lower statistical support. Therefore, our discussion will focus on the topology recovered by the ML and MrBayes methods.

*Eochroa trimenii*, traditionally classified in Bunaeeini (e.g., Bouvier, 1936; Pinhey, 1972; Staude et al., 2016; Staude et al., 2020), was not recovered in this tribe but as sister group of Micragonini, albeit with full nodal support only in the MrBayes tree. Its exclusion
from Bunaeini had earlier been proposed by Oberprieler (1997) and was formalized by Cooper (2002), who placed it in a separate tribe, Eochroini. This result cannot be compared with previous mitochondrial and non-mitochondrial phylogeny estimates, because Eochroa is here included in a phylogenetic analysis for the first time. Although our result does not discount the possible inclusion of Eochroa in Micragonini, such a placement is repudiated by the fact that Eochroa does not possess the synapomorphic characters of Micragonini, as outlined by Oberprieler & Nässig (1994) and Oberprieler (1997). Furthermore, its relationship to the genus Usta, which appears in a similar position as sister group of Micragonini in the analyses of Regier et al. (2008) and Rubin et al. (2018), needs to be investigated. The ML and MrBayes trees recovered the monophyly of the tribes Bunaeini and Micragonini with full nodal support, and the results were broadly in agreement with previous phylogeny estimates (Rubin et al., 2018; Chen et al., 2021).

ML and MrBayes outperformed PhyloBayes in that the broad tribal structure of the first two methods is congruent with those of non-mitochondrial phylogeny estimates (Regier et al., 2008; Barber et al., 2015; Rubin et al., 2018), except that the tribes Micragonini and Eochroini are not recovered as sister group of Bunaeini but of Attacini+Saturniini instead. Different phylogenetic reconstruction methods (i.e., Bayesian inference vs ML) may impact tree topology, especially with regards to the order of deeper nodes. For example, a study of the high-level phylogeny of the Coleoptera inferred with mitogenome sequences showed that different inference models (ML and Bayesian inference) can yield inconsistent
topologies for the same data (Yuan et al., 2016). Additionally, our analyses did not include all African Saturniidae lineages (missing in particular the various genera currently classified in Urotini), and this incomplete sampling may also not be sufficient for a well-supported recovery of the tribal structure of the family. For example, mitogenomic phylogeny estimates of Lepidoptera (Timmermans, Lees & Simonsen, 2014) and Hymenoptera (Mao, Gibson & Dowton, 2015) have failed to resolve some of the relationships with confidence, likely due to limited taxon sampling. The inclusion of representatives of Urotini will allow further insights into the mitochondrial phylogenetic reconstruction of Saturniidae, as this tribe was non-monophyletic in the phylogeny estimates based on nuclear genes (Regier et al., 2008) and phylomorphology (Rubin et al., 2018).

**Phylogenetic trees including C. trifenestrata and N. haraldi**

The phylogenetic analyses including *C. trifenestrata* and *N. haraldi* (Figs. S1–S2) resulted in lower support for the monophyly of Saturniini, and *N. haraldi* was recovered in a basal position relative to all other Saturniidae. The genus *Neoris* was not represented in the phylogeny estimates derived from nuclear genes (Regier et al., 2008) or phylomorphology (Rubin et al., 2018); therefore, it is not possible to establish comparisons. The monophyly of Attacini had low support in the ML tree with *Rhodinia fugax*, traditionally classified...
in Saturniini but morphologically similar to Attacini, as the basal-most member of Attacini, but the Bayesian trees more strongly and consistently supported the monophyly of this tribe including *R. fugax*. *Cricula* was recovered as sister group of *Antheraea* in non-mitochondrial phylogeny estimates (Regier et al., 2008; Rubin et al., 2018), but the position of *C. trifemestra* was recovered inconsistently in our trees as well as in a previous work, where it formed a group diverged from the other Saturniini along with *N. haraldi* (Langley et al., 2020). In the present study, we detected several issues in the mitogenomes of *C. trifemestra* and *N. haraldi*, including cases of shifts in the reading frame caused by artefactual single nucleotide indels that we corrected, but it is possible that other sequencing errors evaded our curation. Therefore, the positions of *N. haraldi* and *C. trifemestra* seem be the result of substandard sequencing quality of these publicly available mitogenomes.

**CONCLUSIONS**

Prior to this study, baseline mitogenomic data for African Saturniidae with utility for assessments of the genetic diversity, phylogeography and phylogenetic relationships in the family were scarce compared with Asian counterparts. We sequenced the mitogenomes of 12 African Saturniidae species, including the first representatives of the tribes Eochroini and Micragonini, and significantly expanded the available genetic information for this group,
which contains numerous species of economic, nutritional and cultural importance in sub-Saharan Africa. Our results support the monophyly of the tribes Bunaenini, Micragonini, Saturniini and Attacini, but with \textit{E. trimenii} excluded from Bunaenini and placed in a separate tribe, Eochroini, and as sister group of Micargonini, and the sister relationship between Saturniini and Attacini, with the latter tribe including \textit{R. fugax} as its basal-most member. However, a sister group relationship between Micragonini+Eochroini and Bunaenini was not consistently found, recovered only in the PhyloBayes reconstruction but not in the ML and MrBayes ones. These present findings contribute towards a more comprehensive understanding of the diversity and relationships among African Saturniidae.

\section*{ACKNOWLEDGEMENTS}

The authors are grateful to Bronwyn Egan (University of Limpopo), Gail Morland (Namibia University of Science and Technology), William Versfeld (Ongava Research Center), Chrissie Fourie and Margriet Brink for assistance with specimen collection.

\section*{ADDITIONAL INFORMATION AND DECLARATIONS}

\subsection*{Funding}

This study was funded by the National Research Foundation of South Africa (Grant No. 121293). Ruan Veldtman was funded by The South African Department of Forestry, Fisheries and the Environment (DFFtE). Zwannda Nethavhani received a PhD bursary from the Foundational Biodiversity Information Programme (Grant No. 128333). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

\subsection*{Grant Disclosures}

The following grant information was disclosed by the authors:
The National Research Foundation of South Africa: 121293.
The South African Department of Forestry, Fisheries and the Environment (DFFtE).
The Foundational Biodiversity Information Programme: 128333.

\subsection*{Competing Interests}

The authors declare there are no competing interests.

\subsection*{Author Contributions}

- Zwannda Nethavhani, Rieze Straeuli and Kayleigh Hiscock performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Ruan Veldtman conceived and designed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Andrew Morton conceived and designed the experiments, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Rolf G. Oberprieler and Barbara van Asch conceived and designed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
Field Study Permissions
The following information was supplied relating to field study approvals (i.e., approving body and any reference numbers):

Specimen collections were approved by Ezemvelo KZN Permits Office (OP408/2020), Cape Nature (CN44-59-13497) and National Commission on Research Science and Technology (AN20190911).

Data Availability
The following information was supplied regarding data availability:

The raw Next-Generation Sequencing data is available at GenBank: PRJNA796275. The mitogenomes reported are available at GenBank: OL912807 to OL912817 and OL912953.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.13275#supplemental-information.

REFERENCES

Barber J, Leavell BC, Keener AL, Breinholt JW, Chadwell BA, McClure CJW, Hill GM, Kawahara A. 2015. Moth tails divert bat attack: evolution of acoustic deflection. Proceedings of the National Academy of Sciences of the United States of America 112:2812–2816 DOI 10.1073/pnas.1421926112.

Bouvier E. 1936. Étude des Saturnioides normaux: famille des Saturniidés. Mémoires du Muséum National d'Histoire Naturelle (Nouvelle Série) 3:1–354.

Cai C, Tihelka E, Pisani D, Donoghue PCJ. 2020. Data curation and modelling of compositional heterogeneity in insect phylo-genomics: a case study of the phylogeny of Dytiscoidea (Coleoptera: Aderhaga). Molecular Biology and Evolution 147:1–7 DOI 10.1016/j.ympev.2020.106782.

Cameron SL. 2014. Insect mitochondrial genomics: implications for evolution and phylogeny. Annual Reviews of Entomology 59:95–117 DOI 10.1146/annurev-ento-011613-162007.

Cao Yong-Qiang Cao Y-Q, Chuan M, Chen J-Y, Yang D-R. 2012. The complete mitochondrial genomes of two ghost moths, Thitarodes renzhiensis and Thitarodes yunnanensis: the ancestral gene arrangement in Lepidoptera. BMC Genomics 13:276 DOI 10.1186/1471-2164-13-276.

Castresana J. 2000. Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. Molecular Biology and Evolution 17:540–552 DOI 10.1093/oxfordjournals.molbev.a026334.

Chen D-B, Zhang RS, Jin XD, Yang J, Li P, Liu YQ. 2021. First complete mitochondrial genome of Rhodinia species (Lepidoptera: Saturniidae): genome description and phylogenetic implication. Bulletin of Entomological Research 3:1–10 DOI 10.1017/S0007485321000808.

Cooper MR. 2002. Note(s) on classification. In: Cooper MR, Cooper MD, eds. The emperor moths of KwaZulu-Natal. 1st edition. New Germany: Peroniceras Press, 103.
Dzerefos CM. 2018. Conservation of edible insects in sub-Saharan Africa. In: Halloran A, Flore AR, Vantomme P, Roos N, eds. Edible insects in sustainable food systems. Cham: Springer, 181–197 DOI 10.1007/978-3-319-74011-9_12.

Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O. 2010. New algorithms and methods to estimate maximum-likelihood phylogenies: Assessing the performance of PhyML 3.0. Systematic Biology 59:307–321 DOI 10.1093/sysbio/syq010.

He YY, Wang X, Chen LS. 2017. The complete mitochondrial genome of Neoris haraldi Schawerda (Lepidoptera: Saturniidae). Mitochondrial DNA Part B Resources 8:625–626 DOI 10.1080/23802359.2017.1372721.

Hlongwane ZT, Slotow R, Munyai TC. 2020. Nutritional composition of edible insects consumed in Africa: a systematic review. Nutrients 12:1–28 DOI 10.3390/nu12092786.

Hlongwane ZT, Slotow R, Munyai TC. 2021. Indigenous knowledge about consumption of edible insects in South Africa. Insects 12:1–19 DOI 10.3390/insects12010022.

Hoang DT, Chernomor O, Von Haeseler A, Minh BQ, Vinh LS. 2018. UFBoot2: improving the ultrafast bootstrap approximation. Molecular Biology and Evolution 35:518–522 DOI 10.1093/molbev/msx281.

Huelsenbeck JP, Ronquist F. 2001. MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics 17:754–755 DOI 10.1093/bioinformatics/17.8.754.

Jiang ST, Hong GY, Yu M, Li N, Yang Y, Liu YQ, Wei ZJ. 2009. Characterization of the complete mitochondrial genome of the giant silkworm moth, Eriogyna pyretorum (Lepidoptera: Saturniidae). International Journal of Biological Sciences 5:351–365 DOI 10.7150/ijbs.5.351.

Jongema Y. 2015. List of edible insects of the world. Wageningen: Wageningen University & Research 1–75. Available at https://www.wur.nl/en/Research-Results/Chair-groups/Plant-Sciences/Laboratory-of-Entomology/Edible-insects/Worldwide-species-list.htm.

Kelemu S, Niassy S, Torto B, Fiaboe K, Affognon H, Tonnang H, Maniania NK, Ekesi S. 2015. African edible insects for food and feed: inventory, diversity, commonalities and contribution to food security. Journal of Insects As Food Feed 1:103–119 DOI 10.3920/JIFF2014.0016.

Kim JS, Kim MJ, Jeong JS, Kim I. 2018. Complete mitochondrial genome of Saturnia jonasi (Lepidoptera: Saturniidae): Genomic comparisons and phylogenetic inference among Bombycoidea. Genomics 110:274–282 DOI 10.1016/j.ygeno.2017.11.004.

Kim JS, Park JS, Kim MJ, Kang PD, Kim SG, Jin BR, Han YS, Kim I. 2012. Complete nucleotide sequence and organization of the mitochondrial genome of eri-silkworm, Samia cynthia ricini (Lepidoptera: Saturniidae). Journal of Asia-Pacific Entomology 15:162–173 DOI 10.1016/j.aspen.2011.10.002.

Kim SR, Kim JS, Kim KY, Kim MJ, Jeong JS, Kim I. 2017a. Complete mitochondrial genome of the wild silkmoth, Saturnia boisduvalii (Lepidoptera: Saturniidae). Entomological Research 47:344–351 DOI 10.1111/1748-5967.12256.
Kim SR, Kim KY, Jeong JS, Kim MJ, Kim K, Choi KH. 2017b. Population genetic characterization of the Japanese oak silk-moth, *Antheraea yamamai* (Lepidoptera: Saturniidae), using novel microsatellite markers and mitochondrial DNA gene sequences. *Genetic and Molecular Research* 16:gmr16029608 DOI 10.4238/gmr16029608.

Kim SR, Kim M-I, Hong MY, Kim KY, Kang PD, Hwang JS, Han YS, Jin BR, Kim I. 2009. The complete mitogenome sequence of the Japanese oak silkmoth, *Antheraea yamamai* (Lepidoptera: Saturniidae). *Molecular Biology Reports* 36:1871–1880 DOI 10.1007/s11033-008-9393-2.

Kitching IJ, Rougerie R, Zwick A, Hamilton CA, St Laurent RA, Naumann S, Mejia LB, Kawahara AY. 2018. A global checklist of the Bombycoidea (Insecta: Lepidoptera). *Biodiversity Data Journal* 6:e22236 DOI 10.3897/BDJ.6.e22236.

Kumar S, Stecher G, Li M, Knyaz C, Tamura K. 2018. MEGA X: molecular evolutionary genetics analysis across computing platforms. *Molecular Biology and Evolution* 35:1547–1549 DOI 10.1093/molbev/msy096.

Kusia ES, Borgemeister C, Khamis FM, Copeland RS, Tanga CM, Ombura FL, Subramanian S. 2021. Diversity, host plants and potential distribution of edible saturniid caterpillars in Kenya. *Insects* 12:1–21 DOI 10.3390/insects12070600.

Lanfear R, Calcott B, Ho SYW, Guindon S. 2012. PartitionFinder: combined selection of partitioning schemes and substitution models for phylogenetic analyses. *Molecular Biology and Evolution* 29:1695–1170 DOI 10.1093/molbev/mss020.

Larson JD, Calcott B, Ho SYW, Guindon S. 2017. Partitionfinder 2: new methods for selecting partitioned models of evolution for molecular and morphological phylogenetic analyses. *Molecular Biology and Evolution* 34:772–773 DOI 10.1093/molbev/msw260.

Langley J, Van der Westhuizen S, Morland G, Van Asch B. 2020. Mitochondrial genomes and polymorphic regions of *Gonimbrasia belina* and *Gynanisa maja* (Lepidoptera: Saturniidae), two important edible caterpillars of Southern Africa. *International Journal of Biological Macromolecules* 144:632–642 DOI 10.1016/j.ijbiomac.2019.12.055.

Lartillot N, Brinkmann H, Philippe H. 2007. Suppression of long-branch attraction artefacts in the animal phylogeny using a site-heterogeneous model. *BMC Evolutionary Biology* 7:1–14 DOI 10.1186/1471-2148-7-S1-S4.

Lartillot N, Rodrigue N, Stubbs D, Richer J. 2013. PhyloBayes-MPI A Bayesian software for phylogenetic reconstruction using mixture models MPI version. *Systematic Biology* 62:611–615.

Laslett D, Canbäck B. 2008. ARWEN: A program to detect tRNA genes in metazoan mitochondrial nucleotide sequences. *Bioinformatics* 24:172–175 DOI 10.1093/bioinformatics/btm573.

Li W, Zhang Z, Lin L, Terenius O. 2017. *Antheraea pernyi* (Lepidoptera: Saturniidae) and its importance in sericulture, food consumption, and traditional Chinese medicine. *Journal of Economic Entomology* 110:1404–1411 DOI 10.1093/jee/tox140.

Liu J, Dai J, Jia J, Zong Y, Sun Y, Peng Y, Wang L, Qian C, Zhu B, Wei G. 2021. Characterization and phylogenetic analysis of the complete mitochondrial
genome of *Saturnia japonica*. Biochemical Genetics online ahead of print DOI 10.1007/s10528-021-10129-9.

Liu YC, Li XY, Chen D-B, Liu YQ, Li XS. 2020. The complete mitochondrial genome of *Antheraea pernyi* strain qing_6 (Lepidoptera: Saturniidae). *Mitochondrial DNA Part B Resources* 5:3397–3398 DOI 10.1080/23802359.2020.1823903.

Malaisse F, Mabossy-Mobouna G, Latham P. 2017. Un Atlas des chenilles et chrysalides consommées en Afrique par l’homme. *Geo-E* 41:55–66.

Mao M, Gibson T, Dowton M. 2015. Higher-level phylogeny of the Hymenoptera inferred from mitochondrial genomes. *Molecular Phylogenetics and Evolution* 84:34–43 DOI 10.1016/j.ympev.2014.12.009.

Margam VM, Coates BS, Hellmich RL, Agunbiade T, Seufferheld MJ, Sun W, Ba MN, Sanon A, Binso-Dabire CL, Baoua I, Ishiyaku MF, Covas FG, Srinivasan R, Armstrong J, Murdock LL, Pittendrigh BR. 2011. Mitochondrial genome sequence and expression profiling for the legume pod borer *Maruca vitrata* (Lepidoptera: Crambidae). *PLOS ONE* 6:e16444 DOI 10.1371/journal.pone.0016444.

McGeoch MA. 2002. Insect conservation in South Africa: an overview. *African Entomology* 10:1–10.

Miller MA, Pfeiffer W, Schwartz T. 2010. The CIPRES science gateway: a community resource for phylogenetic analyses. In: *Proceedings of the 2010 gateway computing environments workshop (GCE)*, New Orleans, LA, USA. 1–8.

Nguyen LT, Schmidt HA, Haeseler AVon, Minh BQ. 2015. IQ-TREE: a fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. *Molecular Biology and Evolution* 32:268–274 DOI 10.1093/molbev/msu300.

Nonaka K. 1996. Ethnoentomology of the central Kalahari San. *African Study Monographs* 22:29–46.

Oberprieler RG. 1995. *The Emperor Moths of Namibia*. Hartbeespoort, South Africa: Ekogilde.

Oberprieler RG. 1997. Classification of the African Saturniidae (Lepidoptera) the quest for natural groups and relationships. *Metamorphosis. Occasional Supplement* 3:142–155.

Oberprieler RG, Morton AS, vam Noort Noort S. 2021. The life history of *Vegetia grimmia* (Geyer, 1832) (Saturniidae: Bunaenae: Micragonini), with an account of its discovery, distribution and taxonomic distinction. *Metamorphosis* 32:74–92.

Oberprieler RG, Nässig WA. 1994. Tarn- oder Warntrachten —ein Vergleich larvaler und imaginaler Strategien bei Saturniiinen (Lepidoptera: Saturniidae). *Nachrichten des Entomologischen Vereins Apollo, Neue Folge* 15:267–304.

Peigler RS, Oberprieler RG. 2017. Ethnographic description of cocoons and silk of the moth families Saturniidae, Lasiocampidae and Psychidae. *Nachrichten des Entomologischen Vereins Apollo* 38:113–120.

Pinhey ECG. 1972. Emperor Moths of south and south central Africa. Cape Town: C. Struk (Pty) Ltd.

Regier JC, Grant MC, Mitter C, Cook CP, Peigler RS, Rougerie R. 2008. Phylogenetic relationships of wild silkmoths (Lepidoptera: Saturniidae) inferred
from four protein-coding nuclear genes. *Systematic Entomology* 33:219–228 DOI 10.1111/j.1365-3113.2007.00416.x.

Rozas J, Ferrer-Mata A, Sanchez-DelBarrio JC, Guirao-Rico S, Librado P, Ramos-Onsins SE, Sanchez-Gracia A. 2017. DnaSP 6: DNA sequence polymorphism analysis of large data sets. *Molecular Biology and Evolution* 34:3299–3302 DOI 10.1093/molbev/msx248.

Rubin JJ, Hamilton CA, McClure CJW, Chadwell BA, Kawahara AY, Barber JR. 2018. The evolution of anti-bat sensory illusions in moths. *Science Advances* 4:eaar7428 DOI 10.1126/sciadv.aar7428.

Saikia M, Nath R, Devi D. 2019. Genetic diversity and phylogeny analysis of *Antheraea assamensis* Helfer (Lepidoptera: Saturniidae) based on mitochondrial DNA sequences. *Journal of Genetics* 98:1–12 DOI 10.1007/s12041-019-1072-7.

Salinas-Giegé T, Giegé R, Giegé P. 2015. TRNA biology in mitochondria. *International Journal of Molecular Sciences* 16:4518–4559 DOI 10.3390/ijms16034518.

Sambrook J, Russell DW. 2012. In: Michael R, Sambrook J, eds. *Molecular cloning: a laboratory manual*. 4th edition. New York: Cold Spring Harbor Laboratory Press, 2028.

Sekonya JG, McClure NJ, Wynberg RP. 2020. New pressures, old foodways: Governance and access to edible mopane cater-pillars, *Imbrasia (=Gonimbrasia) belina*, in the context of commercialization and environmental change in South Africa. *International Journal of the Commons* 14:139–153 DOI 10.5334/ijc.978.

Silow CA. 1976. Edible and other insects of mid-western Zambia: studies in ethnoentomology II. In occasional Papers of the Institutionen för och Janforonde Etnografivia Universitet, Almqvist & Wiksell, Uppsala. 223.

Sima YH, Chen M, Yao R, Li YP, Liu T, Jin X, Wang L-P, Su JF, Li XS, Liu YQ. 2013. The complete mitochondrial genome of the Ailanthus silkmoth, *Samia cynthia cynthia* (Lepidoptera: Saturniidae). *Gene* 526:309–317 DOI 10.1016/j.gene.2013.05.048.

Singh D, Kabiraj D, Sharma P, Chetia H, Mosahari PV, Neog K, Bora U. 2017. The mitochondrial genome of Muga silkworm (*Antheraea assamensis*) and its comparative analysis with other lepidopteran insects. *PLOS ONE* 12:e0188077 DOI 10.1371/journal.pone.0188077.

Siozios S, Massa A, Parr CL, Verspoor RL, Hurst GDD. 2020. DNA barcoding reveals incorrect labelling of insects sold as food in the UK. *PeerJ* 8:e8496 DOI 10.7717/peerj.8496.

Song F, Li H, Jiang P, Zhou X, Liu J, Sun C, Vogler AP, Cai W. 2016. Capturing the phylogeny of holometabola with mitochondrial genome data and Bayesian site-heterogeneous mixture models. *Genome Biology and Evolution* 8:1411–1426 DOI 10.1093/gbe/evw086.

Staude HS, Maclean M, Mecenero S, Pretorius RJ, Oberprieler RG, Van Noort S, Sharp A, Sharp I, Balona J, Bradley S, Brink M, Morton AS, Bodha MJ, Collins SC, Grobler Q, Edge DA, Williams MG, Sihvonen P. 2020. An overview of Lepidoptera-host-parasitoid associations for southern Africa, including an illustrated report...
on 2370 African Lepidoptera-host and 119 parasitoid-Lepidoptera associations. 

Staude HS, Mecenero S, Oberprieler RG, Sharp A, Sharp I, Williams MC, Maclean M. 2016. An illustrated report on the larvae and adults of 962 African Lepidoptera species. Results of the Caterpillar Rearing Group: a novel, collaborative method of rearing and recording lepidopteran life-histories. Metamorphosis 31:1–380.

Thomas B. 2013. Sustainable harvesting and trading of mopane worms (Imbrasia belina) in Northern Namibia: an experience from the Uukwaluudhi area. International Journal of Environmental Studies 70:494–502 DOI 10.1080/00207233.2013.829324.

Timmermans MJTN, Lees DC, Simonsen TJ. 2014. Towards a mitogenomic phylogeny of Lepidoptera. Molecular Phylogenetics and Evolution 79:169–178 DOI 10.1016/j.ympev.2014.05.031.

Yuan ML, Zhang QL, Zhang L, Guo ZL, Liu YJ, Shen YY, Shao R. 2016. High-level phylogeny of the Coleoptera inferred with mitochondrial genome sequences. Molecular Phylogenetics and Evolution 104:99–111 DOI 10.1016/j.ympev.2016.08.002.

Zhang YL, Zhao JH, Zhou QM. 2016. The complete mitochondrial genome of Antheraea pernyi strain Yu6 (Lepidoptera: Saturniidae). Mitochondrial DNA Part A 27:4228–4229 DOI 10.3109/19401736.2015.1022753.

Zhao F, Su T, He B, Jiang K, Zuo C, Lin G, Huang Z. 2021. Complete mitochondrial genome of Actias dubernardi (Lepidoptera: Saturniidae). Mitochondrial DNA Part B Resources 6:627–628 DOI 10.1080/23802359.2021.1875929.

Zhong J, Liu Z-H, Yang W-K, Zhu F, Zhan-Peng D. 2017. Sequencing and analysis of the complete mitochondrial genome of the Antheraea assama (Lepidoptera: Saturniidae). Acta Entomologica Sinica 60:936–949 DOI 10.16380/j.kcxb.2017.08.011.

FURTHER READING

Chen MM, Li Y, Chen M, Wang H, Li Q, Xia RX, Zeng CY, Li YP, Liu YQ, Qin L. 2014. Complete mitochondrial genome of the atlas moth, Attacus atlas (Lepidoptera: Saturniidae) and the phylogenetic relationship of Saturniidae species. Gene 545:95–101 DOI 10.1016/j.gene.2014.05.002.

Hong MY, Lee EM, Jo YH, Park HC, Kim SR, Hwang JS, Jin BR, Kang PD, Kim KG, Han YS, Kim I. 2008. Complete nucleotide sequence and organization of the mitogenome of the silk moth Caligula boisduvalii (Lepidoptera: Saturniidae) and comparison with other lepidopteran insects. Gene 413:49–57 DOI 10.1016/j.gene.2008.01.019.

Liu Y, Pan MLY, Dai F, Zhu X, Lu C, Xiang Z. 2008. The complete mitochondrial genome of the Chinese oak silkmoth, Antheraea pernyi (Lepidoptera: Saturniidae). Acta Biochimica et Biophysica Sinica 40:693–703 DOI 10.111/j.1745-7270.2008.00449.x.

Liu QN, Zhu BJ, Dai LS, Wei GQ, Liu CL. 2012. The complete mitochondrial genome of the wild silkmoth, Actias selene. Gene 505:291–299 DOI 10.1016/j.gene.2012.06.003.
Park JS, Kim MJ, Kim I. 2016. The complete mitochondrial genome of the moon moth, Actias aliena (Lepidoptera: Saturniidae). Mitochondrial DNA Part A 27:149–150 DOI 10.3109/19401736.2013.878918.

Shantibala T, Devi KM, Lokeshwari RK, Anju S, Luikham R. 2018. Complete mitochondrial genome of a latent wild oak tasar silkworm, Antheraea frithi (Lepidoptera: Saturniidae). Mitochondrial DNA Part B Resources 3:15–16 DOI 10.1080/23802359.2017.1413290.

Shantibala T, Victor T, Luikham R, Arunkumar KP, Debaraj SH, Lokeshwari RK, Kim I. 2016. Complete mitochondrial genome of the wild eri silkworm, Samia canningi (Lepidoptera: Saturniidae). Mitochondrial DNA Part A 27:844–845 DOI 10.3109/19401736.2014.919469.

Yang J, Zhang RS, Chen D-B, Chen MM, Li YP, Liu YQ. 2019. The complete mitochondrial genome of Antheraea proylei strain In981 (Lepidoptera: Saturniidae). Mitochondrial DNA Part B Resources 4:2264–2265 DOI 10.1080/23802359.2019.1627944.