Complete mitochondrial genomes of the slugs *Deroceras laeve* (Agriolimacidae) and *Ambigolimax valentianus* (Limacidae) provide insights into the phylogeny of Stylommatophora (Mollusca, Gastropoda)

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Abstract

In this study, we sequenced two complete mitogenomes from *Deroceras laeve* and *Ambigolimax valentianus*. The mitogenome of *Ambigolimax valentianus* represented the first such data from the family Limacidae. The lengths of the mitogenomes of *Deroceras laeve* and *Ambigolimax valentianus* were 14,773 bp and 15,195 bp, respectively. The entire set of 37 mitochondrial genes were identified for both mitogenomes. Compared with the mitogenome of *Achatina fulica*, the *trnP*-*trnA* tRNA cluster was rearranged in both *Deroceras laeve* and *Ambigolimax valentianus*. The secondary structures of tRNA and rRNA genes for the two species were predicted. Phylogenetic analyses based on amino acid sequences supported (1) monophyly of Stylommatophora, (2) division of Stylommatophora into the ‘achatinoid’ clade (i.e., the suborder Achatinina) and the ‘non-achatinoid’ clade (i.e., the suborder Helicina), (3) placement of the Orthurethra in the ‘non-achatinoid’ clade, and (4) monophyly of each of the superfamilies Helicoidea, Urocoptoidea, Succineoidea, Arionoidea, Pupilloidea and Limacoidea. The exemplars of Helicidae, Philomyoidae and Achatinellidae displayed many more mitochondrial gene rearrangements than other species of Stylommatophora.

Key words: Gene rearrangement, Limacoidea, mitogenome, next generation sequencing, phylogeny

Introduction

Typically, the metazoan mitochondrial genome (mitogenome) is a closed-circular and small (15–20 kb) genome encoding 13 protein-coding genes (PCGs), 22 transfer RNAs (tRNAs) and two ribosomal RNAs (rRNAs) (Boore 1999). Mitogenomes have been widely used to resolve phylogenetic relationships within molluscs (e.g., He et al. 2016; Minton et al. 2016; Xie et al. 2019; Guzmán et al. 2021). With the development of new sequencing technologies and the significantly decreased cost of next generation sequencing, the numbers of available mitogenomes has increased rapidly. Molluscs are the second largest...
phylum next to Arthropoda, with about 52,500 extant species (Ponder et al.
2020). However, relatively few mitogenomes are available for this group. Within
Mollusca, Stylommatophora includes the vast majority of terrestrial snails and
slugs. As of December 2022, only 58 stylommatophoran mitogenomes were
available in GenBank.

The Stylommatophora is the largest group within the pulmonate gastro-
pods, containing 20,000 ± 1500 species (Rosenberg et al. 2022). Based on the
structure of the excretory system Pilsbry (1900) divided the Stylommatophora
adapted this system by recognizing a fourth major group, viz. Mesurethra. Of
these four taxa, only Orthurethra is still widely accepted as a natural group.
Deep-level relationships within Stylommatophora have been controversial.
Nordsieck (1992) suggested dividing Stylommatophora into two subclades,
Orthurethra and Sigmurethra. The first comprehensive molecular study of sty-
ломматофор relationships was undertaken by Wade et al. (2001), who rec-
ognized an ‘achatinoid’ clade and a ‘non-achatinoid’ clade. This hypothesis was
subsequently supported by Wade et al. (2006). Bouchet et al. (2005) divided the
Stylommatophora into three clades, Elasmognatha, Orthurethra and Sigmure-
thra, based a combined analysis of morphological and molecular data. Elas-
mognatha contains Succineoidea and Athoracophoroidea. Orthurethra con-
tains Partuloidea, Achatinelloidea, Cochlicopoidea, Pupilloidea and Enoidea.
Sigmurethra was suggested to be an informal group, which contains Clausil-
ioidea, Orthalicoidea, Achatinoidea, Aillyoidea, Testacelloidea, Papilloderma-
toidea, Streptaxoidea, Rhytidoidea, Acavoidae, Plectopyloidea, Puncticidae
and Sagdoidae. Bouchet et al. (2017) recognized Stylommatophora as an or-
der and divided it into three suborders, Achatinina, Helicina and Scolodontina.
Achatinina comprises Achatinoidea and Streptaxoidea. Helicina includes Coo-
lociontoidea, Papillodermatoidea, Plectopyloidea, Puncticidae, Testacelloidea and
Urocoptoidea. Scolodontina contains the single family Scolodontidae. Sa-
di and Wade (2019) further refined this hypothesis by recognizing Scolodon-
tidae as sister to all other stylommatophoran groups comprising the ‘achatini-
od’ and ‘non-achatinoid’ clades. Scolodontidae corresponded to the suborder
Scolodontina proposed by Bouchet et al. (2017), while the ‘achatinoid’ clade
corresponded to the suborder Achatinina and the ‘non-achatinoid’ clade corre-
sponded to the suborder Helicina.

Limacoidea is a superfamily of Stylommatophora that is subdivided into
four families: Agriolimacidae, Limacidae, Boettgerillidae and Vitrienidae (Haus-
dorf 1998; Bouchet et al. 2017). The slug *Ambigolimax valentianus* is an inva-
sive species in North and South America, Africa and Asia (Robinson 1999).
Several studies have proved the usefulness of mitogenome data to resolve
phylogenetic relationships in Stylommatophora (Minton et al. 2016; Xie et al.
2019; Guzmán et al. 2021). In this study, we applied next generation sequenc-
ing to obtain the complete mitogenomes of *Deroceras laeve* (O. F. Müller,
1774) and *Ambigolimax valentianus* (A. Férussac, 1821). The mitogenome
of *Deroceras laeve* represented the second for Agriolimacidae and that of
*Ambigolimax valentianus* was the first for Limacidae. This contribution aims
at characterizing these two new mitogenomes and using them for a phyloge-
netic analysis of Stylommatophora.
Material and methods

Specimens and DNA extraction

Specimens of Deroceras laeve and Ambigolimax valentianus were collected from Zunyi, Guizhou Province, China, in July, 2020. They were identified by checking their adult morphological characters and blasting the mitochondrial cox1 gene sequences in the BOLD system. The voucher specimens were deposited at the Henan Agricultural University, Zhengzhou, China, under the accession numbers MT-Zy20200701 and MT-Zy20200702. The specimens were preserved in absolute ethanol, and stored at -80 °C until DNA extraction. Total genomic DNA of the individual specimen was extracted with the TIANamp Genomic DNA Kit (TIANGEN BIOTECH CO., LTD), following the manufacturer's protocol.

Genome sequencing, assembly and annotation

Genome sequencing was performed on an Illumina HiSeq2500 platform, with a strategy of 150 paired-end sequencing. Library generation for the Illumina Hiseq sequencing was carried out using the Illumina TruSeqTM DNA Sample Prep Kit (Illumina, San Diego, CA, USA), with 350 bp insert size. NGS QC Toolkit v.2.3.3 (Patel and Jain 2012) was used to check the quality of the data. Adapters, poly-N, and low-quality reads were removed from raw data. About 3 Gb clean data obtained by NGS for each species were used to assemble the mitochondrial scaffold.

GetOrganelle v.1.7.5.2 (Jin et al. 2020) was used for mitogenome assembly. The GetOrganelle animal database (-F animal_mt) was applied to identify, filter, and assemble target-associated reads. The new mitogenomes were annotated with the MITOS webserver (Bernt et al. 2013) (http://mitos2.bioinf.uni-leipzig.de/index.py). The gene boundaries of protein-coding genes were refined by alignment against mitochondrial gene sequences of closely related species. tRNA genes were identified using MITOS (Bernt et al. 2013) and ARWEN (Laslett and Canbäck 2008), and the secondary structures were redrawn in Adobe Illustrator CC 2019. The secondary structures of rRNA genes were predicted with reference to Omalonyx unguis (Guzmán et al. 2021). The mitogenome structure images were generated using mtviz (http://pacosy.informatik.uni-leipzig.de/mtviz). The annotated mitogenome sequences were submitted to GenBank under the accession numbers of OQ198714 (Deroceras laeve) and OQ198715 (Ambigolimax valentianus).

Characterization of the new mitogenomes

Pairwise comparisons of gene order with the gene order of Achatina fulica (He et al. 2016; Yang et al. 2016; Xie et al. 2019) and assessment of rearrangement events were performed using CREx (http://pacosy.informatik.uni-leipzig.de/crex/form) (Bernt et al. 2007). The nucleotide compositions of the mitogenome sequences were calculated with MEGA 11 (Kumar et al. 2018). AT and GC-skew values were obtained using the following formulas: AT-skew = (A-T)/(A+T) and GC-skew = (G-C)/(G+C) (Perna and Kocher 1995).
Sequence alignment

Protein-coding genes were aligned individually using MUSCLE as implemented in MEGA 11 (with default settings) (Kumar et al. 2018). Protein-coding genes were translated into amino acid sequences using the invertebrate mitochondrial genetic code. The alignments of genes were concatenated with FASconCAT-G_v.1.04 (Kück and Longo 2014) to create the amino acid dataset PCG_aa.

Phylogenetic analysis

A total of 68 mollusk mitogenome sequences were used in the phylogenetic analyses, of which 59 species were included as ingroup to represent Stylommatophora. Four species from Systellommatophora, two species from Ellobiida and three species from Hygrophila were selected as outgroups (Suppl. material 1). Phylogenetic analyses were performed based on the amino acid dataset mentioned above, under maximum likelihood (ML) and Bayesian inference (BI) criteria. ML analysis was performed with IQ-TREE v.1.6.10 (Nguyen et al. 2015). The data was partitioned by gene types. The best-fitting substitution models for partitions were chosen using ModelFinder (Kalyaanamoorthy et al. 2017) implemented in IQ-TREE. Branch support (BS) values were calculated using ultrafast bootstrap with 10,000 replicates. BI analysis was conducted using MrBayes v.3.2.7 (Ronquist et al. 2012). Two runs with four chains each were performed. The initial number of generations for each run was set to 10 million. Sampling was done every 1000 generations. The Average Standard Deviation of Split Frequencies (ASDSF) were monitored using Tracer v.1.7 (Rambaut et al. 2018). After reaching convergence (ASDSF < 0.01), the tree and branch length information were summarized using the \texttt{sumt} command, discarding the first 25% as burn-in. The consensus tree was yielded, and posterior probability (PP) values were used to assess branch support.

Results

Characteristics of the new mitogenomes

The entire mitochondrial genomes of Deroceras laeve and Ambigolimax valentianus were 14,773 base pairs (bp) and 15,195 bp long, respectively. They contained the entire set of 37 genes usually present in the animal mitogenomes and had identical gene orders (Fig. 1). Compared with the gene order of Achatina fulica (He et al. 2016; Yang et al. 2016; Xie et al. 2019), the trnP_trnA cluster was rearranged in both Deroceras laeve and Ambigolimax valentianus. The CREx analysis showed that the rearranged gene order of Deroceras laeve and Ambigolimax valentianus has evolved as the result of transposition (Fig. 2). The nucleotide compositions of both mitogenomes were heavily biased towards A and T. The overall A+T content of Deroceras laeve was 73.0%, while the A+T content of Ambigolimax valentianus was 71.4%. Both mitogenomes had the negative AT-skew values (-0.115 for Deroceras laeve and -0.096 for Ambigolimax valentianus) and the positive GC-skew values (0.181 for Deroceras laeve and 0.113 for Ambigolimax valentianus) in the major strand. This indicated the occurrence of more T than A and more G than C.
For the protein-coding genes, ATG (five for *Deroceras laeve* and four for *Ambigolimax valentianus*), ATT (six for *Deroceras laeve* and three for *Ambigolimax valentianus*) and ATA (one for *Ambigolimax valentianus*) were used as the start codons. For the *cox2* gene and *atp8* gene of *Ambigolimax valentianus*, GTG was the start codon. For the *cox1* gene and *cob* gene of both *Ambigolimax valentianus* and *Deroceras laeve*, TTG was the start codon. All protein-coding genes terminated with the stop codon TAA or TAG, except for *nad3*.

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**Figure 1.** Gene order and gene content of the mitogenome of *Deroceras laeve*. The abbreviation of genes follows MITOS. Starting from *trnI* and ending at *cox3*, count clockwise. The outer numbers indicate the positions of each section.

**Figure 2.** The rearrangement event assessed from the CREx analysis for the mitogenome of *Deroceras laeve*.
and nad4L of *Ambigolimax valentianus*, which had the incomplete stop codon T. For both species, Leu, Ile, Phe and Val were the most frequently used amino acids. Relative synonymous codon usage (RSCU) for 13 protein-coding genes of *Deroceras laeve* is shown in Fig. 3. *Ambigolimax valentianus* (Suppl. material 2) had similar RSCU values to *Deroceras laeve*.

<table>
<thead>
<tr>
<th>Gene</th>
<th>RSCU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gin</td>
<td>1.78</td>
</tr>
<tr>
<td>His</td>
<td>1.81</td>
</tr>
<tr>
<td>Asn</td>
<td>3.62</td>
</tr>
<tr>
<td>Pro</td>
<td>3.7</td>
</tr>
<tr>
<td>Thr</td>
<td>4.26</td>
</tr>
<tr>
<td>Leu1</td>
<td>2.98</td>
</tr>
<tr>
<td>Glu</td>
<td>2.09</td>
</tr>
<tr>
<td>Met</td>
<td>6.38</td>
</tr>
<tr>
<td>Arg</td>
<td>1.48</td>
</tr>
<tr>
<td>Tyr</td>
<td>4.04</td>
</tr>
<tr>
<td>Asp</td>
<td>1.75</td>
</tr>
<tr>
<td>Lys</td>
<td>2.31</td>
</tr>
<tr>
<td>Ala</td>
<td>4.9</td>
</tr>
<tr>
<td>Ile</td>
<td>5.01</td>
</tr>
<tr>
<td>Ser1</td>
<td>5.13</td>
</tr>
<tr>
<td>Ser2</td>
<td>13.23</td>
</tr>
<tr>
<td>Leu2</td>
<td>1.17</td>
</tr>
<tr>
<td>Cys</td>
<td>2.79</td>
</tr>
<tr>
<td>Trp</td>
<td>7.83</td>
</tr>
<tr>
<td>Val</td>
<td>6.82</td>
</tr>
<tr>
<td>Gly</td>
<td>8.38</td>
</tr>
<tr>
<td>Phe</td>
<td>4.9</td>
</tr>
</tbody>
</table>

Figure 3. Codon usage of the 13 mitochondrial protein-coding genes of *Deroceras laeve*. RSCU: relative synonymous codon usage.

All 22 tRNA genes were identified by both MITOS and ARWEN, and their lengths ranged from 61 to 68 bp. Most tRNA genes of both species can be folded into the classic clover-leaf structure (Fig. 4 and Suppl. material 3). The trnT gene of *Deroceras laeve* had an unusual TΨC loop. The trnK gene and trnS1 gene of *Ambigolimax valentianus* had an incomplete DHU arm. The position of the rrlL gene was located between trnV and trnL1, while the rrmS gene was found between trnE and trnM. In *Deroceras laeve*, rrlL had a length of 1052 bp with an A + T content of 74.4%, whereas rrmS had a length of 697 bp with an A + T content of 71.5%. In *Ambigolimax valentianus*, rrlL was 1091 bp long with an A + T content of 75.9%, while rrmS was 711 bp long with an A + T content of 72.7%. The secondary structures for rrlL and rrmS of *Deroceras laeve* and *Ambigolimax valentianus* are presented in Figs 5, 6 and Suppl. materials 4, 5, respectively. The rrlL molecule of both species contained six domains (labeled I–VI) comprising 43 helices. The rrmS molecule consisted of three domains (labeled I–III) and 28 helices.

**Phylogenetic inference**

ML and BI produced similar tree topologies. The monophyly of Stylommatophora was supported under both analyses (Figs 7, 8). Achatinoidea represented by *Achatina fulica* was consistently resolved as the sister group of all other Stylommatophora (BS = 100, PP = 1.0). This lineage corresponded to the suborder Achatinina. The remaining stylommatophorans formed the ‘non-achatinoid’ clade which corresponded to the suborder Helicina. In Helicina, all superfamilies with more than two representatives were well supported, with the exception of Punctoidea. All families with multiple representatives were supported as monophyletic. The newly sequenced *Deroceras laeve* was sister to *Deroceras reticulatum*, and both together formed the
sister group of *Ambigolimax valentinanus*. The three species form a monophyletic Limacoidea clade.

The major differences between the ML and BI analyses were the positions of Xanthonychidae and the clade Orthalicoidea + Punctoidea. The ML analysis placed Xanthonychidae as sister to all other Helicoidea. Whereas, the BI analysis recovered Xanthonychidae as the sister group of a clade Geomitridae + Helicidae. The ML analysis placed Orthalicoidea + Punctoidea between a clade comprising (Clausilioidea + Pupilloidea) and Arionoidea comprising (Philomycidae + Arionidae). But this arrangement received no statistical support. The BI analysis recovered Orthalicoidea + Punctoidea between the clade (Clausilioidea + Pupilloidea) and a clade including Limacoidea, Helicarioidea, Trochomphoidea and Punctidae.
Figure 5. The secondary structure of rml inferred for Deroceras laeve.
Figure 6. The secondary structure of rrnS inferred for Deroceras laeve.
Figure 7. ML phylogenetic tree inferred with IQ-TREE using amino acid sequences of 13 PCGs. Numbers at the nodes are ultrafast bootstrap values (BS > 70). Blue indicates the newly sequenced species. The branch of *Cepaea nemoralis* is depicted as half of its original branch length. Scale bar represents substitutions/site.
Discussion

Mitochondrial gene rearrangement

Large-scale changes in genomes are considered to be rare events (Rokas and Holland 2000). The gene set remains constant across bilaterian animals, mitochondrial gene rearrangements appear to be unusual, and gene order is selectively neutral (Boore et al. 1995; Dowton et al. 2002; Dowton et al. 2009; Cameron 2014). In a certain animal group, for example the hexapods, most species share an identical mitogenome organization (Dowton et al. 2002; Cameron 2014). However, Stylommatophora mitogenomes have experienced many more mitochondrial gene rearrangements than other groups (Minton et al. 2016; Xie et al. 2019; Guzmán et al. 2021). In this study, two new mitogenomes from *Deroceras laeve* and *Ambigolimax valentianus* had a tRNA gene rearrangement associated with the *trnP_trnA* tRNA cluster. We compared the gene order for all included mitogenomes and found that all the Helicina species have gene rearrangements compared to *Achatina fulica*. The exemplars of Helicidae, Philomyidae and Achatinellidae displayed many more mitochondrial gene rearrangements than others (Fig. 8).

Phylogeny of Stylommatophora

Within Stylommatophora, the division of the order into Achatinina and Helicina is well accepted (Wade et al. 2001; Wade et al. 2006). Recently, some authors added Scolodontidae to the phylogenetic analysis (Ramírez et al. 2012; Bouchet et al. 2017; Saadi and Wade 2019). Currently, no mitogenomes of Scolodontidae have been published, so the phylogenetic position of Scolodontidae could not be assessed in this analysis. Our analyses consistently supported the division of Stylommatophora into two principal clades, Achatinina and Helicina. This result is consistent with the previous mitogenome analyses (Xie et al. 2019; Guzmán et al. 2021).

The present mitogenome data recovered Clausilioidea as a sister group of the orthurethran clade. This result contrasted with Wade et al. (2006), who recovered Orthurethra to be close to a clade comprising Arionoidea and Limacoidea. In a prior mitogenome analysis, a sister group relationship between Succineoidea and Arionoidea was supported (Xie et al. 2019). In this study, this pattern was supported by the BI analysis (PP = 0.96) based on expanded taxon sampling of mitogenomes. In our analyses, only one species of *Haplotrema minimum* (Ancey, 1888) representing Haplotrematoidea was included due to mitogenome data availability. Both the ML and BI analyses placed Haplortematoidea as sister to Helicoidea. However, this relationship had low support values. In future researches, larger taxon samples are needed to identify the phylogenetic placement of Haplotrematoidea.

The superfamily Punctoidea included the families Punctidae, Charopidae, Cystopeltidae, Discidae (“Endodontidae”), Helicodiscidae, Oopeltidae and Oreohelicidae (Bouchet et al. 2017). In this study, we included three species of Punctoidea in the phylogenetic analyses, which respectively represented Discidae [*Discus perspectivus* (Megerle von Mühlfeld, 1816)], Oreohelicidae [*Oreohelix idahoensis* (Newcomb, 1866)] and Punctidae [*Punctum randolphii* (Dall, 1895)]. *Discus perspectivus* and *O. idahoensis* were significantly supported to
Figure 8. Bayesian phylogenetic tree inferred in MrBayes using amino acid sequences of 13 PCGs (left), and gene order comparisons for the stylommatophoran mitogenomes (right). Numbers at the nodes in the tree are Bayesian posterior probabilities (PP > 0.9). Blue indicates the newly sequenced species. The branch of *Cepaea nemoralis* is depicted as half of its original branch length. Scale bar represents substitutions/site. In the mitogenome structure maps, gene rearrangements are highlighted by red.
be a sister group. *Punctum randolphi*i was placed separately, and clustered with a clade including Trochomorphoidea, Helicarinoidea and Limacoidea. Taxon sampling of Punctoidea was very limited in our analysis. The monophyly of Punctoidea needs to be further tested by additional species in future studies.

Previous analyses based on the multiple gene fragments have demonstrated the monophyly of Helicoidea, which comprised the families Helicidae, Bradybaenidae, Xanthonychidae, Hygromiidae, Camaenidae, Polygyridae and Sagiddae (Wade et al. 2006). Our results strongly supported a monophyletic Helicoidea (BS = 100, PP = 1.0). In addition, the superfamilies Urocoptoidea, Succineoidea, Arionoidea, Pupilloidea and Limacoidea were supported to be monophyletic groups. These patterns are congruent with the current taxonomy of land snails (Bouchet et al. 2017).

For the taxon sampling of outgroups, we included four exemplars from Systellommatophora and two from Ellobiida. The monophyly of Systellommatophora and Ellobiida were consistently supported in both ML and BI analyses. Moreover, Systellommatophora and Ellobiida formed a sister-group relationship (BS = 100, PP = 1). Amphipulmonata Schrödl, 2014 was established as a clade containing Systellommatophora and Ellobiida (Bouchet et al. 2017). Amphipulmonata (comprising Ellobioidea and Systellommatophora) was supported in the previous molecular phylogenetic analysis (Dayrat et al. 2011). Our results confirmed the hypothesis of Amphipulmonata.

**Additional information**

**Conflict of interest**

The authors have declared that no competing interests exist.

**Ethical statement**

No ethical statement was reported.

**Funding**

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**Author contributions**

N.S. and T.Z. designed the research; N.S., X.L., Y.Z. and T.Z. performed the research and analyzed the data; N.S., T.Z. and X.L. wrote the paper. All authors have read and agreed to the published version of the manuscript.

**Data availability**

The mitogenome sequences newly generated in this study were deposited in GenBank, with the accession numbers of OQ198714–OQ198715. The dataset used for phylogenetic analyses can be obtained from the corresponding author upon request.

**References**


**Supplementary material 1**

**Taxa included in this study, including GenBank Accession numbers and literature references**

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**Supplementary material 2**

Relative synonymous codon usage (RSCU) for 13 protein-coding genes of Ambigolimax valentianus

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Data type: figure (PDF file)
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Supplementary material 3
The secondary structures of tRNA genes inferred for the mitogenome of *Ambigolimax valentianus*

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Supplementary material 4
The secondary structure of *rrnL* inferred for *Ambigolimax valentianus*

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Supplementary material 5
The secondary structure of *rrnS* inferred for *Ambigolimax valentianus*

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