Complete paternally inherited mitogenomes of two freshwater mussels *Unio pictorum* and *Sinanodonta woodiana* (Bivalvia: Unionidae)

Artur Burzyński¹ and Marianna Soroka²

¹ Department of Genetics and Marine Biotechnology, Institute of Oceanology Polish Academy of Sciences, Sopot, Poland
² University of Szczecin, Faculty of Biology, Department of Genetics, Szczecin, Poland

**ABSTRACT**

Freshwater bivalves from the family Unionidae usually have two very divergent mitogenomes, inherited according to the doubly uniparental model. The early divergence of these two mitogenomic lineages gives a unique opportunity to use two mitogenomic data sets in a single phylogenetic context. However, the number of complete sequences of the maternally inherited mitogenomes of these animals available in GenBank greatly exceeds that of the paternally inherited mitogenomes. This is a problem for phylogenetic reconstruction because it limits the use of both mitogenomic data sets. Moreover, since long branch attraction phenomenon can bias reconstructions if only a few but highly divergent taxa are considered, the shortage of the faster evolving paternally inherited mitogenome sequences is a real problem. Here we provide, for the first time, complete sequences of the M mitogenomes sampled from Polish populations of two species: native *Unio pictorum* and invasive *Sinanodonta woodiana*. It increases the available set of mitogenomic pairs to 18 species per family, and allows unambiguous reconstruction of phylogenetic relationships among them. The reconstructions based on M and F mitogenomes which were separated for many millions of years, and subject to differing evolutionary dynamics, are fully congruent.

**INTRODUCTION**

Freshwater mussels from the order Unionoida are known for the widespread occurrence of doubly uniparental inheritance (DUI) of mitochondria (Walker et al., 2006). Under DUI all males are heteroplasmic and pass their mitochondrial DNA to male offspring whereas females pass their mitochondria to all their offspring (Skibinski, Gallagher & Beynon, 1994; Zouros et al., 1994). Consequently, the two independent mitochondrial lineages exist. In freshwater mussels (order Unionoida) the divergence between the two lineages is extreme because their emergence predates the family level radiation (Hoeh, Stewart & Guttman, 2002). The maternal (F) and paternal (M) mitogenomic lineages in Unionidae, the best studied of families from this order, differ slightly in genome organization but the genomes...
are fairly conservative structure-wise within each lineage, with only one rearrangement leading to alternative F gene order in one group of Unionidae (Breton et al., 2009). The two mitogenomes accumulated several unique gender-specific features, in the form of additional ORFs (Milani et al., 2013; Mitchell et al., 2016a) as well as extensions of the existing cox2 CDS (Chapman et al., 2008). These features seem to be associated with DUI and sex determination systems because in species reverting to hermaphroditism both DUI and these peculiar mitogenomic features are eventually lost (Soroka & Burzyński, 2017).

Mitogenomic data are frequently used to reconstruct and date phylogenies (Mitchell et al., 2016b; Burzyński et al., 2017; Kieren et al., 2018; Krzeminska et al., 2018). The existence of two parallel mitochondrial lineages provides a unique opportunity to test the limits of such methodology (Burzyński et al., 2017). This is important in the context of critical re-evaluation of the usability of mitochondrial markers in resolving phylogenies (Wallis et al., 2017).

The first complete mitogenome of a unionid bivalve, Lampsilis ornata, was published more than 15 years ago (Serb & Lydeard, 2003) and the number of complete mitogenome sequences available in public databases is rising rapidly ever since. There are currently complete sequences of F-type mitogenomes of at least 24 species from two major subfamilies: Unionini and Anodontini alone in GenBank (Soroka & Burzyński, 2017). However, there are much fewer M-type mitogenomes present in GenBank, with only nine members of these two subfamilies represented, and a total of 17 M-type mitogenomes available for the whole Unionidae family (Table 1). Notably, two M-type mitogenomes of species from the sister family Margaritiferidae have been published recently (Lopes-Lima et al., 2017a; Guerra et al., 2017).

It has been suggested that sparse and uneven taxon sampling biases the phylogenetic reconstructions, necessitating the need for more M mitogenomic data. To fill this gap, and check the usability of double-mitogenome approach in phylogenies, we sequenced complete M-type mitogenomes of two freshwater unionid species from Poland: the invasive Sinanodonta woodiana from Anodontini and the native Unio pictorum from Unionini. We selected two species for which the F-type mitogenomic data were already published (Table 1). Moreover, for S. woodiana there was also one GenBank record describing the complete M-type from the native range of the species, without formal publication.

**METHODS**

Mussels were sampled from the Oder River, 25 km south of the city Szczecin (in the north-western part of Poland, 53.2123N 14.4673E) in 2006. The bivalves were collected in May, June and July when their sex could be reliably identified by microscopic examination of their mature gonads. Tissue samples were immediately collected and processed for DNA isolation. Two males were selected at random from each species for full mitogenome sequencing. DNA was extracted from gonads using established methodology (Burzyński & Soroka, 2018). Material used in this study has been kept in 70% ethanol at the repository of the Department of Genetics, University of Szczecin with voucher numbers: AW5 and AW10 (S. woodiana), UP149 and UP232 (U. pictorum). Long Range Polymerase Chain
| Latin name                       | Accession F | Reference F       | Accession M | Reference M       |
|---------------------------------|-------------|-------------------|-------------|-------------------|
| Aculamprotula tortuosa (Lea, 1865) | KC109779    | Wang, Cao & Li (2013) | KC441487    | –                 |
| Anodonta anatina (Linnaeus, 1758) | KF030964    | Soroka & Burzyński (2015) | KF030963    | Soroka & Burzyński (2016) |
| Arconia lanceolata (Lea, 1856)   | KJ144818    | Wang, Guo & Li (2016) | KJ775864    | –                 |
| Lamprotula leaii (Griffith & Pidgeon, 1833) | JQ691662 | –                  | KC847114    | –                 |
| Margaritifera marocana (Pallary, 1918) | KY131953    | Lopes-Lima et al. (2017a) | KY131954    | F                 |
| Margaritifera monodonta (Say, 1829) | KU873123    | Guerra et al. (2017) | KU873124    | F                 |
| Potamilus alatus (Say, 1817)     | KU559011    | Wen et al. (2017)  | KU559010    | F                 |
| Potomia littoralis (Cuvier, 1798) | KT247374    | Froufe et al. (2016) | KT247375    | F                 |
| Pronodularia japonensis (Lea, 1859) | AB055625 | –                  | AB055624    | –                 |
| Pyganodon grandis (Say, 1829)    | FJ809754    | Breton et al. (2009) | FJ809755    | F                 |
| Quadrula quadrula (Rafinesque, 1820) | FJ809750 | Breton et al. (2009) | FJ809751    | F                 |
| Sinanodonta woodiana (Lea, 1834) CH | KM272949    | Zhang et al. (2016) | KM434235    | –                 |
| Sinanodonta woodiana (Lea, 1834) PL | HQ823346    | Soroka (2010)      | MH349359    | This study        |
| Sinohyriopsis cumingii (Lea, 1852) | KM393224    | Wei et al. (2016)  | KC150028    | –                 |
| Solenia carinata (Heude, 1877)   | KC848654    | Huang et al. (2013) | KC848655    | F                 |
| Unio crassus Retzius, 1788       | KY290447    | Burzyński et al. (2017) | KY290450    | F                 |
| Unio delphinus Spengler, 1793    | KT326917    | Fonseca et al. (2016) | KT326918    | F                 |
| Unio pictorum (Linnaeus, 1758)   | HM014133    | Soroka & Burzyński (2010) | MH349358    | This study        |
| Unio tumidus Retzius, 1788       | KY021076    | Soroka & Burzyński (2018) | KY021075    | F                 |
| Utterbackia peninsularis Bogan & Hoeh, 1995 | HM856636 | Breton et al. (2011) | HM856635    | F                 |
| Venustaconcha ellipsiformis (Conrad, 1836) | FJ809753 | Breton et al. (2009) | FJ809752    | F                 |

Reaction (LR-PCR) primers were designed and used to amplify the complete M and F mitogenomes in two overlapping fragments, followed by primer walking approach (Table S1). Individual PCR products were then sequenced using ABI Big Dye Terminator technology in Macrogen (South Korea). Raw sequencing reads were trimmed and assembled using Staden package (Bonfield, Smith & Staden, 1995).

Complete mitochondrial genomes were annotated as described previously (Zbawicka, Burzyński & Wenne, 2007), using CRITICA (Badger & Olsen, 1999), GLIMMER (Delcher et al., 1999), BLAST (Altschul et al., 1990; Zhang & Madden, 1997), Wise2 (Burney, Clamp & Durbin, 2004), HMMER (Wheeler & Eddy, 2013), and ARWEN (Laslett & Canbäck, 2008). Annotated sequences have been deposited in GenBank under the following accession numbers: MH349356 (AW5), MH349357 (UP232) MH349358 (UP149) and MH349359 (AW10). The annotations present in GenBank record KM434235 were inspected and adjusted to match those of MH349357 and MH349359, for consistency, using the same methodology.

MEGA7 (Kumar, Stecher & Tamura, 2016) was used to align sequences and calculate all diversity indices. Sliding window analysis of genetic diversity was done in DnaSP (Librado & Rozas, 2009). Circular maps and nucleotide composition analyses were produced in
CGView (Stothard & Wishart, 2005). All other sequence manipulations, such as feature extraction and alignment concatenation, were performed in CLC Genomics Workbench version 9.5.4 (https://www.qiagenbioinformatics.com/).

A data set for phylogenetic reconstructions of relationships within Unionidae was selected based on the availability of the complete mitogenomic sequences from both lineages, belonging to Unionidae family and present in GenBank (nr database) at the time of this writing (February 2018). In addition to all such pairs of sequences from Unionidae, two species from Margaritiferidae family were also used to serve as an outgroup. These two families are generally considered to have a sister relationship (Walker et al., 2006; Bogan & Roe, 2008). The details of species names and accession numbers, along with the relevant references, are given in Table 1. All taxonomic names were verified and adjusted according to MUSSELp (2018) (http://mussel-project.uwsp.edu/) database. To overcome the problems associated with potential shifts of mutational pressure due to genomic rearrangements as well as with saturation effects, the analysis should be performed in amino acid space. However, as previously noted (Burzyński et al., 2017), some database sequences do contain suspicious frame shifts leading to unreliable amino acid alignments and biased or plainly wrong reconstructions. To overcome this problem the following procedure was applied. First, all protein coding genes were extracted and translated, following the existing CDS annotations. Then the alignments were produced in both amino acid and nucleotide space and inspected, for each gene and lineage separately. Sequences, for which the nucleotide alignment was inconsistent with the amino acid alignment were deemed suspicious and were removed from the alignments. The curated alignment was then used to produce hmm alignment profile (Finn, Clements & Eddy, 2011), of each protein in both lineages (26 hmm profiles). Finally, these profiles were used to recover protein alignments directly from raw sequence data using genewise (Birney, Clamp & Durbin, 2004). Amino acid alignments were concatenated and used in all subsequent analyses (protein alignments). The rRNA genes were extracted separately and used directly after alignment and concatenation (rDNA alignments). The lengths of all alignments were: 3,907 aa (F), 4,240 aa (M), 2,273 bp (F) and 2,389 bp (M). However, since the gap columns are not used in reconstructions, the alignments were effectively shorter: 3,636 aa (F), 3,524 aa (M), 1,738 bp (F) and 1,694 bp (M), for protein and rDNA alignments of F and M lineages, respectively. The apparently high proportion of gaps in the M protein alignment is caused solely by the unusually short cox2 extension in one species: Potamilus alatus (Wen et al., 2017), and is not indicative of any alignment problems. All four alignments are available as Supplemental Information 1.

Phylogenetic reconstructions by Bayesian Inference (BI) were done in BEAST2 (Bouckaert et al., 2014), as described previously (Burzyński et al., 2017; Soroka & Burzyński, 2017). Input XML files for BEAST were prepared in BEAUTi. All the analyses were run in quadruplicates, for $10^7$ generations. The resulting log files were inspected in Tracer (Rambaud & Drummond, 2013) to verify the convergence of the runs. There was a good agreement between the repeats, hence all four log files as well as tree files were combined in logcombiner, after removal of 10% burn-in states. Each parameter has reached effective sample size (ESS) of more than 300, in the combined log file. The consensus tree was
obtained from the concatenated treefile in `treeannotator`. There were three such analyses: one using only the M-type data, one using only the F-type data, and one using the data obtained from both mitogenomes in a single reconstruction. Each alignment was added as a separate partition, with individually selected model of substitutions, but with the same tree for all partitions. The optimal model of substitutions was selected, following the recommended Path Sampling procedure (Baele et al., 2012; Baele et al., 2013). Consistently, mtRev model was selected for protein partitions and GTR model for rDNA partitions, both with gamma-distributed heterogeneities of rates. Relaxed, uncorrelated lognormal clock was used following previous recommendations (Burzyński et al., 2017), and the Yule tree prior was assumed in all analyses. The trees were visualized in FigTree (Rambaut, 2009). To allow the comparison of the obtained topologies as well as node heights, all trees were arbitrary normalized to the most recent common ancestor (MRCA) of the family Unionidae.

Phylogenetic reconstructions were also performed under Maximum Likelihood (ML) framework, as implemented in IQ-TREE (Nguyen et al., 2015). First, the optimal model was selected for each individual alignment separately, using `modelfinder` (Kalyaanamoorthy et al., 2017). The following models were chosen by BIC criterion: for protein alignments mtVer+F+R5 (F), mtMet+F+R4 (M) and for both rDNA alignments TIM2+F+I+G4. Then, the three reconstructions were performed, using partitioned data consisted of either two M-type, two F-type or all four alignments in a single reconstruction (Chernomor, Haeseler & Minh, 2016). Finally, to access the stability of the obtained topology, ultrafast bootstrap procedure was used, with 1,000 replicates (Minh, Nguyen & Haeseler, 2013).

Separate analysis was done on a portion of M-type mitogenome containing m-ORF to obtain estimates of relative divergence of the two copies of m-ORF apparently present in *S. woodiana* mitogenome. Since these sequences evolve exceptionally quickly and can only serve comparisons at subfamily level at best (Mitchell et al., 2016a), it is impossible to produce reliable amino acid alignment for them. To overcome this difficulty, m-ORF annotations were extracted from the five closest relatives (FJ809751, KF030963, HM856635, KM434235 and MH349359) and aligned in nucleotide space. This short alignment (seven sequences, 429 positions) was then used in BEAST to reconstruct the phylogeny of duplication, assuming GTR model of substitutions, with gamma-distributed rates and lognormal relaxed clock. Other parameters and run conditions were the same as for the main phylogenetic reconstruction.

**RESULTS**

The two sequenced M mitogenomes of *U. pictorum* are almost identical, with only six substitutions along the whole 16,632 bp, therefore only one was used in all subsequent analyses (Fig. 1). These genomes are very similar both in structure and in sequence to other M mitogenomes from the compared data set. Particularly striking is the similarity to the M mitogenome of *U. delphinus*, with the nucleotide p-distance at the barcoding *cox1* locus of only 0.04.

The structure and pattern of divergence for the second set of sequenced mitogenomes, the M mitogenomes of *S. woodiana* are different. Again, the two sequenced mitogenomes
Figure 1 Genetic map of the sequenced M-type mitogenomes. The mitogenome of *U. pictorum* (A) is structurally very similar to all M-type mitogenomes from this family published to date. The mitogenome of *S. woodiana* (B) is more exceptional because of the two additional features labeled in red. All protein coding genes are labeled by the names of the encoded proteins, two rRNA genes are labeled 16S and 12S for large and small subunit, respectively. The trn genes are labeled by the one letter code of the respective amino acid. Direction of transcription is indicated by the position and direction of the arrows, with clockwise transcribed genes on the outside and anticlockwise transcribed genes on the inside of the circle representing the genome. The two inner circles represent local compositional bias, calculated in a 300 bp long window, in 25 bp long steps. The light gray parts of the first circle represent above average AT skew \[\frac{(A-T)}{(A+T)}\] while the dark gray parts of this circle represent below average AT skews. The inner, black circle represents local GC content. Both indices are calculated with the relation to mitogenome-wide averages which are, for *S. woodiana*: AT-skew -0.3166, GC content 0.3284, for *U. pictorum*: AT-skew -0.2828, GC content 0.3396.

Full-size DOI: 10.7717/peerj.5573/fig-1

were very close to each other (only 22 substitutions, 17,616 bp) allowing the use of a single representative in all subsequent analyses. However, the BLAST search in *nr* database revealed the presence of unpublished mitogenome annotated as M mitogenome of the same species, sampled in China (Table 1). The M mitogenomes of *S. woodiana* sampled in China and sampled in Poland clearly belong to very distant clades because they differ in sequence substantially (average nucleotide p-distance 0.097). The spread of polymorphisms along sequence alignment (Fig. 2) shows fairly typical pattern, with less divergence in rDNA loci but also with several short anomalously divergent regions. However, despite these differences, the mitogenomes are structurally identical and unique among the mitogenomes published to date (Fig. 1). Instead of one supranumerary m-ORF they possess two copies of this gene and one additional *trnMET*. The two copies of m-ORF are quite distinct, clearly their emergence predates the divergence of the two M-clades in *S. woodiana*, as suggested by the phylogenetic reconstruction involving their closest relatives (Fig. 3). The inspection of the *trnMET* suggests that it is utilizing TAT anticodon but otherwise resembles the *trnASP* rather than the canonical *trnMET* serving the start codons (Fig. 4).

Phylogenetic reconstruction (Fig. 5) revealed good agreement between the M and F data sets. The relationships between all taxa were resolved consistently in all three BI
reconstructions and, after normalization to the Unionidae MRCA node, the node heights were also reasonably congruent. The two divergent haplogroups of *S. woodiana* present in the M data set clearly did not have the counterparts in the F data set but apart from this exceptional case all node heights were estimated more reliably using two data sets simultaneously. ML reconstructions yielded the same topology for M and for joined data sets, with good support for the majority of nodes and notable increase in support in the joined analysis. However, when only the F data set was used, the alternative topology was recovered, placing *Arconaia lanceolata* at the root of Unioninae + Anodontinae clade. The support for this topology was poor and all the other parts of the tree were congruent with the remaining analyses.

**DISCUSSION**

Presence of two divergent mitogenomes following the gender-specific distribution in Unionoida gives the opportunity to analyse structural evolution of the mitogenomes along a long evolutionary timescale. Relatively few structural changes has been revealed in this group so far (*Lopes-Lima et al., 2017a*). In particular no changes in gene repertoire has been noticed to date. Here we report, for the first time a case of major CDS duplication.
involving the m-ORF of *S. woodiana*. Moreover, the single duplication event covered also the adjacent *trn* gene, leading to the emergence of a new *trn*, with TAT anticodon. It is more parsimonious to assume that this new *trnMET* originated from the duplicated *trnASP* by the change of specificity than by independent “jump” of *trnMET* which would also require the change in anticodon sequence (Fig. 4). A *trnMET* gene of the same specificity is present in Mytilidae (*Boore, Medina & Rosenberg, 2004*), but has not been reported in freshwater mussels so far. This shows, that the typical small-scale mitogenomic rearrangements may be more common in Unionoida than could have been expected based on the limited data set currently available. The origin of this duplication is uncertain, but based on the phylogenetic reconstructions (Figs. 3 and 5), it seems to be relatively old, possibly predating species level radiation within the genus. Further mitogenomic data from other members of the genus *Sinanodonta* will no doubt allow more precise analysis of this phenomenon.

Mitochondrial markers were traditionally used in phylogeny reconstructions due to their favorable properties of no recombination and clonal inheritance (*Avise, 1986*). Early attempts to apply this approach to Unionidae, using short fragment (650 bp) of F-type *cox1* sequences produced poorly resolved tree, suggesting non-monophyletic Unionidae (*Bogan & Hoeh, 2000*). Inclusion of more taxa in later analyses allowed for recognition of three well-supported clades within Unionidae (Unioninae, Anodontinae and Ambleminae) as well as confirmation of Margaritiferidae as a sister group to monophyletic Unionidae (*Graf & Ó Foighil, 2000*). However, this marker alone was not sufficient to resolve the internal relationships within Unionidae with any certainty. Increasing the length of mitochondrial sequences by adding 753 bp of *nd1* and 315 bp of *16S* resulted in a limited increase of support for some nodes but the relationship between the major clades remained unsolved (*Campbell et al., 2005*). The addition of a nuclear rDNA marker (473 bp of 28S) along with several morphological characters was also not helpful (*Graf & Cummings, 2006*).
Figure 5  Phylogenetic reconstruction of relationships within Unionidae. Phylogenetic reconstruction of relationships within Unionidae based on the concatenated amino acid alignment of all 13 mitochondrially encoded proteins and nucleotide alignment of the two rRNA genes, from both M and F type mitogenomes (A). Six separate reconstructions were performed: three under BI and three under ML framework, separately for the F and M data sets as well as jointly for both data sets. The presented tree is a result of BI under joined data set but the topology of all three trees obtained under BI was the same. All nodes have posterior probability of 1.0. The 95% CI on node heights is shown as blue bars at nodes for the joined M+F analysis, as red bars above nodes for the analysis based on the M data set and as green bars below nodes for the analysis based on the F data set. The topology obtained under ML framework for M and for joined M+F data sets was the same, with 100% bootstrap support for most of the nodes. The nodes with less than 100% bootstrap support are indicated by the actual percentage numbers next to the CI bars, in the color corresponding to the used data set (red for M, green for F, and blue for M+F). The two “zero” values in green indicate that the topology recovered by ML under the F data set was different, placing A. lanceolata outside the Unioninae clade with poor support, so that these nodes were not present. Relevant fragment of this alternative tree is also shown (B), again only nodes with less than 100% support are labeled. The yellow bar is the 95% CI on the MRCA of the apparent duplication of m-ORF within the S. woodiana lineage, transferred from Fig. 3.

More recent study using similar cox1, 16S, and 28S markers (alignment length 1,900 bp) produced similarly inconclusive reconstruction (Pfeiffer & Graf, 2015). Only recently the complete mitogenomic data has been applied to phylogeny reconstruction within Unionidae. The most straightforward analysis, with 12 protein coding genes aligned separately and concatenated did not result in conclusive resolution of all relationships (Fonseca et al., 2016; Froufe et al., 2016). The support for monophyletic Anodontinae + Unioninae was overwhelming in both M-type and F-type clades. However, the position of Gonideinae on the tree was uncertain and inconsistent between F-type and M-type clades. More recent comprehensive attempts to solve the phylogeny of Unionidae used more
sophisticated methods and much better sampling, but were still based on a very limited set of characters: the same \textit{coxl}, 28S and 16S markers (Lopes-Lima et al., 2017b; Bolotov et al., 2017). Both analyses agree that Ambleminae form a sister relationship with Gonideinae. The same conclusion was supported by our recent mitogenomic-based analysis (Burzyński et al., 2017). The phylogeny reconstructed here (Fig. 5) suggests different relationship of the three major clades: places Ambleminae as a basal clade and Gonideinae as a sister group to Anodontinae + Unioninae. Similar topology was recovered in the most recent reconstruction (Lopes-Lima et al., 2017a), although with relatively poor support. The primary reason for this change is most likely the increased taxonomic sampling, particularly the inclusion of proper outgroup taxa from the sister Margaritiferidae family. The increased support for the problematic nodes is most likely caused by the retention of more informative characters in separate M and F alignments, demonstrating that the presented approach is more efficient. In addition to the proper solution of the relationship between the three major clades, the presented reconstruction places the species \textit{Arconaia lanceolata} on the tree closer to the Unioninae than to the Anodontinae clade. This is also in conflict with most of the previous analyses (Lopes-Lima et al., 2017b), in agreement with the F-type clade presented by (Lopes-Lima et al., 2017a), but with much better support. The small remaining uncertainty regarding the placement of Gonideinae, apparent in the ML tree, is understandable, given the fact that this group shares one particular rearrangement within their F type mitogenome (Breton et al., 2009). Such rearrangements can lead to biased mutational pressure and consequently biased reconstruction. Moreover, the mitogenomic data are still lacking for several Unionidae subfamilies (Lopes-Lima et al., 2017b) and it is likely that when they are available the remaining uncertainty will disappear.

The presence of two apparently very divergent mitochondrial clades in a single species, as noted for \textit{S. woodiana}, is frequently interpreted as indicative of cryptic speciation event. In fact similar interpretation was recently given to explain patterns of deep polymorphisms noted in barcoding \textit{coxl} locus in \textit{S. woodiana} populations worldwide (Bolotov et al., 2016; Bespalaya et al., 2018). This view is probable if one looks at comparable divergence between the two closely related \textit{Unio} species (\textit{U. pictorum} and \textit{U. delphinus}): their separation on the tree (Fig. 5) is comparable to the one observed in \textit{S. woodiana} and yet their status as separate species is supported by all recent phylogeographic analyses (Prie & Puillandre, 2014; Araujo et al., 2018). However, for \textit{S. woodiana} the conclusion must not be straightforward. The fact that the two very divergent M clades are present in the set of populations where only a single F clade is present (PL and CH) argues strongly against such interpretation in this case. It may well be that a complex pattern of incomplete lineage sorting and/or hybridization events makes any classic barcoding attempts, that is species inferences based on F mtDNA \textit{coxl} sequences, questionable in this case. Clearly more data are needed to explain this pattern, including the M mitochondrial markers along with markers based on nuclear loci.

**CONCLUSIONS**

The presented mitogenomic data are a valuable resource for further phylogenetic and population genetic studies.
The presented novel approach to phylogenetic reconstructions using mitogenomic data should help to overcome methodological problems, particularly in solving and dating deeper phylogenies.

### ADDITIONAL INFORMATION AND DECLARATIONS

#### Funding
This work was supported by the Polish Ministry of Science and Higher Education (No. N 303 364 33). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

#### Grant Disclosures
The following grant information was disclosed by the authors:
Polish Ministry of Science and Higher Education: No. N 303 364 33.

#### Competing Interests
The authors declare there are no competing interests.

#### Author Contributions
- Artur Burzyński conceived and designed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Marianna Soroka conceived and designed the experiments, performed the experiments, contributed reagents/materials/analysis tools, approved the final draft.

#### DNA Deposition
The following information was supplied regarding the deposition of DNA sequences:
The mitogenomes described here are accessible via GenBank accession numbers: MH349356 (AW5), MH349357 (UP232) MH349358 (UP149) and MH349359 (AW10).

#### Data Availability
The following information was supplied regarding data availability:
The raw data are provided in the Supplemental Files.

#### Supplemental Information
Supplemental information for this article can be found online at [http://dx.doi.org/10.7717/peerj.5573#supplemental-information](http://dx.doi.org/10.7717/peerj.5573#supplemental-information).

### REFERENCES

Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. *Journal of Molecular Biology* 215:403–410 DOI 10.1016/S0022-2836(05)80360-2.

Araujo R, Buckley D, Nagel K-O, Garcia-Jiménez R, Machordom A. 2018. Species boundaries, geographic distribution and evolutionary history of the Western Palearctic freshwater mussels *Unio* (Bivalvia: Unionidae). *Zoological Journal of the Linnean Society* 182:275–299 DOI 10.1093/zoolinnean/zbek039.
Avise JC. 1986. Mitochondrial DNA and the evolutionary genetics of higher animals. Philosophical Transactions of the Royal Society B: Biological Sciences 312:325–342 DOI 10.1098/rstb.1986.0011.

Badger JH, Olsen GJ. 1999. CRITICA: coding region identification tool invoking comparative analysis. Molecular Biology and Evolution 16:512–524 DOI 10.1093/oxfordjournals.molbev.a026133.

Baele G, Lemey P, Bedford T, Rambaut A, Suchard MA, Alekseyenko AV. 2012. Improving the accuracy of demographic and molecular clock model comparison while accommodating phylogenetic uncertainty. Molecular Biology and Evolution 29:2157–2167 DOI 10.1093/molbev/mss084.

Baele G, Li WLS, Drummond AJ, Suchard MA, Lemey P. 2013. Accurate model selection of relaxed molecular clocks in bayesian phylogenetics. Molecular Biology and Evolution 30:239–243 DOI 10.1093/molbev/mss243.

Bespalaya YV, Bolotov IN, Aksenova OV, Gofarov MY, Kondakov AV, Vikhrev IV, Vinarski MV. 2018. DNA barcoding reveals invasion of two cryptic Sinanodonta mussel species (Bivalvia: Unionidae) into the largest Siberian river. Limnologica - Ecology and Management of Inland Waters 69:94–102 DOI 10.1016/j.limno.2017.11.009.

Birney E, Clamp M, Durbin R. 2004. Genewise and genomewise. Genome Research 14:988–995 DOI 10.1101/gr.1865504.

Bogan AE, Hoeh WR. 2000. On becoming cemented: evolutionary relationships among the genera in the freshwater bivalve family Etheriidae (Bivalvia : Unionoida). In: Harper EM, Taylor JD, Crame JA, eds. Evolutionary biology of the bivalvia. Bath: Geological Soc Publishing House, 159–168.

Bogan AE, Roe KJ. 2008. Freshwater bivalve (Unioniformes) diversity, systematics, and evolution: status and future directions. Journal of the North American Benthological Society 27:349–369 DOI 10.1899/07-069.1.

Bolotov IN, Bespalaya YV, Gofarov MY, Kondakov AV, Konopleva ES, Vikhrev IV. 2016. Spreading of the Chinese pond mussel, Sinanodonta woodiana, across Wallacea: one or more lineages invade tropical islands and Europe. Biochemical Systematics and Ecology 67:58–64 DOI 10.1016/j.bse.2016.05.018.

Bolotov IN, Kondakov AV, Vikhrev IV, Aksenova OV, Bespalaya YV, Gofarov MY, Kolosova YS, Konopleva ES, Spitsyn VM, Tanmuangpak K, Tumpeesuwon S. 2017. Ancient river inference explains exceptional oriental freshwater mussel radiations. Scientific Reports 7:2135 DOI 10.1038/s41598-017-02312.

Bonfield JK, Smith KF, Staden R. 1995. A new DNA sequence assembly program. Nucleic Acids Research 23:4992–4999 DOI 10.1093/nar/23.24.4992.

Boore JL, Medina M, Rosenberg LA. 2004. Complete sequences of the highly rearranged molluscan mitochondrial genomes of the scaphopod Graptacme eborea and the bivalve Mytilus edulis. Molecular Biology and Evolution 21:1492–1503 DOI 10.1093/molbev/msh090.

Bouckaert R, Heled J, Kühnert D, Vaughan T, Wu C-H, Xie D, Suchard MA, Rambaut A, Drummond AJ. 2014. BEAST 2: a software platform for bayesian evolutionary
Breton S, Beaupré HD, Stewart DT, Piontkivska H, Karmakar M, Bogan AE, Blier PU, Hoeh WR. 2009. Comparative mitochondrial genomics of freshwater mussels (Bivalvia: Unionoida) with doubly uniparental inheritance of mtDNA: gender-specific open reading frames and putative origins of replication. *Genetics* **183**:1575–1589 DOI 10.1534/genetics.109.110700.

Breton S, Stewart DT, Shepardson S, Trdan RJ, Bogan AE, Chapman EG, Ruminas AJ, Piontkivska H, Hoeh WR. 2011. Novel protein genes in animal mtDNA: a new sex determination system in freshwater mussels (Bivalvia: Unionoida)? *Molecular Biology and Evolution* **28**:1645–1659 DOI 10.1093/molbev/msq345.

Burzyński A, Soroka M, Mioduchowska M, Kaczmarczyk A, Sell J. 2017. The complete maternal and paternal mitochondrial genomes of *Unio crassus*: mitochondrial molecular clock and the overconfidence of molecular dating. *Molecular Phylogenetics and Evolution* **107**:605–608 DOI 10.1016/j.ympev.2016.12.007.

Campbell DC, Serb JM, Buhay JE, Roe KJ, Minton RL, Lydeard C. 2005. Phylogeny of North American amblemines (Bivalvia, Unionoida): prodigious polyphyly proves pervasive across genera. *Invertebrate Biology* **124**:131–164 DOI 10.1111/j.1744-7410.2005.00015.

Chapman EG, Piontkivska H, Walker JM, Stewart DT, Curole JP, Hoeh WR. 2008. Extreme primary and secondary protein structure variability in the chimeric male-transmitted cytochrome c oxidase subunit II protein in freshwater mussels: evidence for an elevated amino acid substitution rate in the face of domain-specific purifying selection. *BMC Evolutionary Biology* **8**:165 DOI 10.1186/1471-2148-8-165.

Chernomor O, Haeseler Avon, Minh BQ. 2016. Terrace aware data structure for phylogenomic inference from supermatrices. *Systematic Biology* **65**:997–1008 DOI 10.1093/sysbio/syw037.

Delcher AL, Harmon D, Kasif S, White O, Salzberg SL. 1999. Improved microbial gene identification with GLIMMER. *Nucleic Acids Research* **27**:4636–4641 DOI 10.1093/nar/27.23.4636.

Finn RD, Clements J, Eddy SR. 2011. HMMER web server: interactive sequence similarity searching. *Nucleic Acids Research* **39**:W29–W37 DOI 10.1093/nar/gkr367.

Fonseca MM, Lopes-Lima M, Eackles MS, King TL, Froufe E. 2016. The female and male mitochondrial genomes of *Unio delphinus* and the phylogeny of freshwater mussels (Bivalvia: Unionoida). *Mitochondrial DNA Part B: Resources* **1**:954–957 DOI 10.1080/23802359.2016.1241677.

Froufe E, Gan HM, Lee YP, Carneiro J, Varandas S, Teixeira A, Ziertz A, Sousa R, Lopes-Lima M. 2016. The male and female complete mitochondrial genome sequences of the Endangered freshwater mussel *Potomida littoralis* (Cuvier, 1798) (Bivalvia: Unionidae). *Mitochondrial DNA Part A* **27**:3571–3572 DOI 10.3109/19401736.2015.1074223.
Graf DL, Cummings KS. 2006. Palaeoheterodont diversity (Mollusca: Trigonioida + Unionoida): what we know and what we wish we knew about freshwater mussel evolution. *Zoological Journal of the Linnean Society* 148:343–394 DOI 10.1111/j.1096-3642.2006.00259.

Graf DL, Ó Foighil D. 2000. The evolution of brooding characters among the freshwater pearly mussels (Bivalvia: Unionoidea) of North America. *Journal of Molluscan Studies* 66:157–170 DOI 10.1093/mollus/66.2.157.

Guerra D, Plazzi F, Stewart DT, Bogan AE, Hoeh WR, Breton S. 2017. Evolution of sex-dependent mtDNA transmission in freshwater mussels (Bivalvia: Unionida). *Scientific Reports* 7:1551 DOI 10.1038/s41598-017-01708-1.

Hoeh WR, Stewart DT, Guttman SI. 2002. High fidelity of mitochondrial genome transmission under the doubly uniparental mode of inheritance in freshwater mussels (Bivalvia: Unionoidea). *Evolution; International Journal of Organic Evolution* 56:2252–2261 DOI 10.1111/j.0014-3820.2002.tb0149.x.

Huang X-C, Rong J, Liu Y, Zhang M-H, Wan Y, Ouyang S, Zhou C-H, Wu X-P. 2013. The complete maternally and paternally inherited mitochondrial genomes of the endangered freshwater mussel *Solenia carinatus* (Bivalvia: Unionidae) and implications for unionidae taxonomy. *PLOS ONE* 8:e84352 DOI 10.1371/journal.pone.0084352.

Kalyaanamoorthy S, Minh BQ, Wong TKF, Haeseler A von, Jermiin LS. 2017. ModelFinder: fast model selection for accurate phylogenetic estimates. *Nature Methods* 14:587–589 DOI 10.1038/nmeth.4285.

Kieren S, Sparreboom M, Hochkirch A, Veith M. 2018. A biogeographic and ecological perspective to the evolution of reproductive behaviour in the family Salamandridae. *Molecular Phylogenetics and Evolution* 121:98–109 DOI 10.1016/j.ympev.2018.01.006.

Krzeminska U, Morales HE, Greening C, Nyari AS, Wilson R, Song BK, Austin CM, Sunnucks P, Pavlova A, Rahman S. 2018. Population mitogenomics provides insights into evolutionary history, source of invasions and diversifying selection in the House Crow (*Corvus splendens*). *Heredity* 120:296–309 DOI 10.1038/s41437-018-0020-7.

Kumar S, Stecher G, Tamura K. 2016. MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. *Molecular Biology and Evolution* 33:1870–1874 DOI 10.1093/molbev/msw054.

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.

Librado P, Rozas J. 2009. DnaSP v5: a software for comprehensive analysis of DNA polymorphism data. *Bioinformatics* 25:1451–1452 DOI 10.1093/bioinformatics/btp187.

Lopes-Lima M, Fonseca MM, Aldridge DC, Bogan AE, Gan HM, Ghamizi M, Sousa R, Teixeira A, Varandas S, Zanatta D, Zieritz A, Froufe E. 2017a. The first Margaritiferidae male (M-type) mitogenome: mitochondrial gene order as a potential character for determining higher-order phylogeny within Unionida (Bivalvia). *Journal of Molluscan Studies* 83:249–252 DOI 10.1093/mollus/eyx009.
Lopes-Lima M, Froufe E, Do VT, Ghamizi M, Mock KE, Kebapci U, Klishko O, Kovitvadhi S, Kovitvadhi U, Paulo OS, Pfeiffer JM, Raley M, Riccardi N, Sereflisan H, Sousa R, Teixeira A, Varandas S, Wu X, Zanatta DT, Zieritz A, Bogan AE. 2017b. Phylogeny of the most species-rich freshwater bivalve family (Bivalvia: Unionida: Unionidae): defining modern subfamilies and tribes. *Molecular Phylogenetics and Evolution* 106:174–191 DOI 10.1016/j.ympev.2016.08.021.

Milani L, Ghiselli F, Guerra D, Breton S, Passamonti M. 2013. A comparative analysis of mitochondrial orfans: new clues on their origin and role in species with doubly uniparental inheritance of mitochondria. *Genome Biology and Evolution* 5:1408–1434 DOI 10.1093/gbe/evt101.

Minh BQ, Nguyen MAT, Von Haeseler A. 2013. Ultrafast approximation for phylogenetic bootstrap. *Molecular Biology and Evolution* 30:1188–1195 DOI 10.1093/molbev/msr024.

Mitchell A, Guerra D, Stewart D, Breton S. 2016a. In silico analyses of mitochondrial ORFans in freshwater mussels (Bivalvia: Unionoida) provide a framework for future studies of their origin and function. *BMC Genomics* 17:597 DOI 10.1186/s12864-016-2986-6.

Mitchell KJ, Wood JR, Llamas B, McLenachan PA, Kardailsky O, Scofield RP, Worthy TH, Cooper A. 2016b. Ancient mitochondrial genomes clarify the evolutionary history of New Zealand’s enigmatic acanthisittid wrens. *Molecular Phylogenetics and Evolution* 102:295–304 DOI 10.1016/j.ympev.2016.05.038.

MUSSELp. 2018. The MUSSELp database. Available at http://mussel-project.uwsp.edu/ (accessed on 20 February 2018).

Nguyen L-T, Schmidt HA, Von Haeseler A, 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.

Pfeiffer JM, Graf DL. 2015. Evolution of bilaterally asymmetrical larvae in freshwater mussels (Bivalvia: Unionoida: Unionidae). *Zoological Journal of the Linnean Society* 175:307–318 DOI 10.1111/zoj.12282.

Prie V, Puillandre N. 2014. Molecular phylogeny, taxonomy, and distribution of French *Unio* species (Bivalvia, Unionidae). *Hydrobiologia* 735:95–110 DOI 10.1007/s10750-013-1571-0.

Rambaut A, Drummond A. 2013. Tracer v1.6. Available at http://tree.bio.ed.ac.uk/software/tracer/ (accessed on 18 February 2018).

Rambaut A. 2009. FigTree, a graphical viewer of phylogenetic trees. Available at http://tree.bio.ed.ac.uk/software/figtree (accessed on 18 February 2018).

Serb JM, Lydeard C. 2003. Complete mtDNA sequence of the North American freshwater mussel, *Lampsilis ornata* (Unionidae): an examination of the evolution and phylogenetic utility of mitochondrial genome organization in Bivalvia (Mollusca). *Molecular Biology and Evolution* 20:1854–1866 DOI 10.1093/molbev/msg218.

Skibinski DOF, Gallagher C, Beynon CM. 1994. Mitochondrial DNA inheritance. *Nature* 368:817–818 DOI 10.1038/368817b0.
Soroka M. 2010. Characteristics of mitochondrial DNA of unionid bivalves (Mollusca: Bivalvia: Unionidae), II. Comparison of complete sequences of maternally inherited mitochondrial genomes of *Sinanodonta woodiana* and *Unio pictorum*. *Folia Malacologica* 18:189–209 DOI 10.2478/v10125-010-0016.

Soroka M, Burzyński A. 2010. Complete sequences of maternally inherited mitochondrial genomes in mussels *Unio pictorum* (Bivalvia, Unionidae). *Journal of Applied Genetics* 51:469–476 DOI 10.1007/BF03208876.

Soroka M, Burzyński A. 2015. Complete female mitochondrial genome of *Anodonta anatina* (Mollusca: Unionidae): confirmation of a novel protein-coding gene (F ORF). *Mitochondrial DNA* 26:267–269 DOI 10.3109/19401736.2013.823176.

Soroka M, Burzyński A. 2016. Complete male mitochondrial genome of *Anodonta anatina* (Mollusca: Unionidae). *Mitochondrial DNA* 27:1679–1680 DOI 10.3109/19401736.2014.958725.

Soroka M, Burzyński A. 2017. Hermaphroditic freshwater mussel *Anodonta cygnea* does not have supranumerary open reading frames in the mitogenome. *Mitochondrial DNA Part B* 2:862–864 DOI 10.1080/23802359.2017.1407705.

Soroka M, Burzyński A. 2018. Doubly uniparental inheritance and highly divergent mitochondrial genomes of the freshwater mussel *Unio tumidus* (Bivalvia: Unionidae). *Hydrobiologia* 810:239–254 DOI 10.1007/s10750-017-3113-7.

Stothard P, Wishart DS. 2005. Circular genome visualization and exploration using CGView. *Bioinformatics* 21:537–539 DOI 10.1093/bioinformatics/bti054.

Walker JM, Curole JP, Wade DE, Chapman EG, Bogan AE, Watters GT, Hoeh WR. 2006. Taxonomic distribution and phylogenetic utility of gender-associated mitochondrial genomes in the Unionoida (Bivalvia). *Malacologia* 48:265–282.

Wallis GP, Cameron-Christie SR, Kennedy HL, Palmer G, Sanders TR, Winter DJ. 2017. Interspecific hybridization causes long-term phylogenetic discordance between nuclear and mitochondrial genomes in freshwater fishes. *Molecular Ecology* 26:3116–3127 DOI 10.1111/mec.14096.

Wang G, Cao X, Li J. 2013. Complete F-type mitochondrial genome of Chinese freshwater mussel *Lamprotula tortuosa*. *Mitochondrial DNA* 24:513–515 DOI 10.3109/19401736.2013.770508.

Wang G, Guo L, Li J. 2016. The F-type complete mitochondrial genome of *Arconaia lanceolata*. *Mitochondrial DNA Part A* 27:322–323 DOI 10.3109/19401736.2014.892098.

Wei M, Yang S, Yu P, Wan Q. 2016. The complete mitochondrial genome of *Hyriopsis cumingii* (Unionoida: Unionidae): genome description and related phylogenetic analyses. *Mitochondrial DNA Part A* 27:1769–1770 DOI 10.3109/19401736.2014.963804.

Wen HB, Cao ZM, Hua D, Xu P, Ma XY, Jin W, Yuan XH, Gu RB. 2017. The complete maternally and paternally inherited mitochondrial genomes of a freshwater mussel *Potamilus alatus* (Bivalvia: Unionidae). *PLOS ONE* 12:e016974 DOI 10.1371/journal.pone.0169749.
Wheeler TJ, Eddy SR. 2013. nhmmer: DNA homology search with profile HMMs. Bioinformatics 29:2487–2489 DOI 10.1093/bioinformatics/btt403.

Zbawicka M, Burzyński A, Wenne R. 2007. Complete sequences of mitochondrial genomes from the Baltic mussel Mytilus trossulus. Gene 406:191–198 DOI 10.1016/j.gene.2007.10.003.

Zhang J, Madden TL. 1997. PowerBLAST: a new network BLAST application for interactive or automated sequence analysis and annotation. Genome Research 7:649–656 DOI 10.1101/gr.7.6.649.

Zhang P, Fang H-Y, Pan W-J, Pan H-C. 2016. The complete mitochondrial genome of Chinese pond mussel Sinanodonta woodiana (Unionoida: Unionidae). Mitochondrial DNA Part A 27:1620–1621 DOI 10.3109/19401736.2014.958697.

Zouros E, Ball AO, Saavedra C, Freeman KR. 1994. Mitochondrial DNA inheritance. Nature 368:818 DOI 10.1038/368818a0.