











Genome Evolution and Introgression in the New Zealand mud Snails *Potamopyrgus estuarinus* and *Potamopyrgus kaitunuparaoa*

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Abstract

We have sequenced, assembled, and analyzed the nuclear and mitochondrial genomes and transcriptomes of *Potamopyrgus estuarinus* and *Potamopyrgus kaitunuparaoa*, two prosobranch snail species native to New Zealand that together span the continuum from estuary to freshwater. These two species are the closest known relatives of the freshwater species *Potamopyrgus antipodarum*—a model for studying the evolution of sex, host–parasite coevolution, and biological invasiveness—and thus provide key evolutionary context for understanding its unusual biology. The *P. estuarinus* and *P. kaitunuparaoa* genomes are very similar in size and overall gene content. Comparative analyses of genome content indicate that these two species harbor a near-identical set of genes involved in meiosis and sperm functions, including seven genes with meiosis-specific functions. These results are consistent with obligate sexual reproduction in these two species and provide a framework for future analyses of *P. antipodarum*—a species comprising both obligately sexual and obligately asexual lineages, each separately derived from a sexual ancestor. Genome-wide multigene phylogenetic analyses indicate that *P. kaitunuparaoa* is likely the closest relative to *P. antipodarum*. We nevertheless show that there has been considerable introgression between *P. estuarinus* and *P. kaitunuparaoa*. That introgression does not extend to the mitochondrial genome, which appears to serve as a barrier to hybridization between *P. estuarinus* and *P. kaitunuparaoa*. Nuclear-encoded genes whose products function in joint mitochondrial-nuclear enzyme complexes exhibit similar patterns of nonintrogression, indicating that incompatibilities between the mitochondrial and the nuclear genome may have prevented more extensive gene flow between these two species.

Key words: *Potamopyrgus*, introgression, gastropod evolution.

Significance

No whole-nuclear genome sequences are currently available for snails of the genus *Potamopyrgus*, best known for *P. antipodarum*, an invasive species of rivers and lakes worldwide, and a famous model for the study of the evolution of sex. We have sequenced and analyzed the genome of sexual *P. estuarinus* and *P. kaitunuparaoa*, the closest known relatives of *P. antipodarum*. We show that (i) the genomes are very similar in gene content and size, (ii) *P. kaitunuparaoa* is the closest relative to *P. antipodarum*, (iii) significant introgression has occurred between *P. estuarinus* and *P. kaitunuparaoa*; these genomes set the stage for powerful direct analyses of the genomic features, e.g. sex to asexual transitions and invasive success, that make *P. antipodarum* unique.

Introduction

Groups of closely related species or lineages that feature multiple separate transitions from one character state to another can be powerfully applied to answer general and trait-specific questions regarding evolutionary processes. Here, we present the first high-quality genome assemblies from two species of *Potamopyrgus*. This gastropod genus appears to represent a recent radiation across New Zealand (Haase 2008), and includes multiple examples of transitions from the ancestral ovoviparous and obligately sexual state to ovovivipary (Haase 2005) and obligately asexual lineages (Wallace 1992; Dybdahl and Lively 1995; Paczesniak et al. 2013). These transitions are of broad interest because both of these traits—the modes of egg production (oviparous vs. ovoviviparous) and of reproduction (sexual vs. asexual)—are likely amongst the most important predictors of the likelihood of success in particular environments (Maynard Smith 1978; Bell 1982; Clutton-Brock 1991). The genetic basis of transitions to asexual reproduction has been characterized at least in part in a handful of taxa (reviewed in Neiman et al. 2014; more recent examples include Yagound et al. 2020; Ma et al. 2021; Mau et al. 2021), but this information remains outstanding for the vast majority of asexual taxa. Transitions between ovipary and viviparity (live birth, equivalent to ovoviviparity) have received quite a bit of attention in vertebrates, but comparatively little is known in invertebrates such as gastropods (Mamos et al. 2021). One pattern that has emerged from studies of ovipary–viviparity transitions in mollusks is that derived viviparity is relatively common in freshwater taxa, while oviparity, which appears to be ancestral in mollusks, is the norm in marine species (Köhler et al. 2004; Glaubrecht 2006).

Our focus here is to provide and analyze the first genome assemblies from two *Potamopyrgus* species: *P. estuarinus* and *P. kaitunuparaoa*. These are amongst the closest known evolutionary relatives of *P. antipodarum* (Haase 2008), itself an emerging model species for studying the evolution of asexual reproduction (e.g. McElroy et al. 2021): multiple ($\gg 10$) sexual to asexual transitions have occurred within *P. antipodarum* (Dybdahl and Lively 1995; Paczesniak et al. 2013). Furthermore, a genome

sequencing project is underway for *P. antipodarum*, including representatives from both sexual and asexual lineages (Sharbrough et al. 2018, 2023; McElroy et al. 2021). By completing the genomes of *P. estuarinus* and *P. kaitunuparaoa*, two closely related taxa with the apparently ancestral character states of obligately sexual reproduction involving the production of oviparous eggs, a clear framework is now laid for identifying the underpinnings of reproductive evolution in *Potamopyrgus*. Indeed, these genome assemblies provide key resources required for more detailed comparative analyses of *Potamopyrgus* to understand the genomic and genetic mechanisms involved in the repeated transitions to ovovivipary and asexuality in this genus. To this end, the careful inventory and detailed analyses of genes that are specifically involved in sexual reproduction are potentially key because biological processes such as meiosis and sperm production would be predicted to be under relaxed selective constraint in asexuals (Schurko et al. 2009). Thus, a clear picture of the ancestral states for such genes in a phylogenetically close and obligately sexual relative is both necessary and informative for identifying and understanding the genomic underpinnings—both cause and consequence—of sexual to asexual transitions.

In addition to our specific motivations to understand reproductive transitions, these new genomes contribute phylogenetic diversity to an otherwise sparse set of currently available mollusk genomes. Collectively, mollusks have a 500-million year fossil record, rich in biodiversity and replete with species of both scientific and economic importance. Despite their importance, genome resources for molluscan taxa, especially gastropods, have lagged compared to other bilaterian animals (Davison and Neiman 2021; Ghiselli et al. 2021). As a result, functional genomics in mollusks has been forced to rely upon annotations ported over from model systems millions of years diverged from focal taxa. Phylogenomic analyses can help span this divide by accounting for changes in rates and patterns of evolution across lineages that are often the hallmarks of altered function (Eisen 1998). For example, genes with relatively conserved functions are expected to exhibit relatively slow rates of evolution and can be readily

detected by comparison to more distant relatives. By contrast, genes that gain new functions or that lose their ancestral functions are expected to exhibit relatively rapid rates of evolution and require comparisons to closer relatives for their detection. Thus, by identifying genes with relatively slow and relatively rapid rates of evolution, we can more accurately predict the putative functions of genes identified strictly by bioinformatic inference. We implemented such a comparative approach, using the transcriptomes of the close relative *P. antipodarum* and a more distantly related Littorinimorph snail, *Oncomelania hupensis*, as well as the genomes from even more distant relatives, the apple snails, *Pomacea canaliculata* and *Marisa cornuarietis*, to inform our genome annotations.

As a result of our detailed comparative analyses of these Caenogastropod genomes, initially aimed at providing an accurate and informative genome assembly and annotation, we identified a substantial phylogenetic discordance among the three *Potamopyrgus* species between mitochondrial and nuclear gene trees. Careful analysis of these data indicates that *P. kaitunuparaoa* and *P. antipodarum* are sister taxa to the exclusion of *P. estuarinus* (matching the mitochondrial genome tree topology), but with substantial introgression between *P. estuarinus* and *P. kaitunuparaoa*. Notably, nuclear genes with predicted interactions with mitochondrially encoded genes were particularly resistant to introgression, consistent with a scenario in which mito-nuclear coevolution might contribute to barriers between species in *Potamopyrgus*.

In sum, we have assembled, annotated, and analyzed the genome sequences of two closely related snail species, *P. estuarinus* and *P. kaitunuparaoa*. These genomes add diversity to an otherwise phylogenetically depauperate collection of complete mollusk genomes, aiding future efforts to annotate and characterize genes in this important animal phylum. The *P. estuarinus* and *P. kaitunuparaoa* genomes reveal a clear picture of the ancestral state for sexual reproduction in this genome as a framework for assessing and understanding the evolution of asexuality in their close relative, *P. antipodarum*. Finally, phylogenetic analyses of these genomes reveal a substantial history of introgression associated with the divergence between *P. estuarinus* and *P. kaitunuparaoa*.

Results and Discussion

We have sequenced, assembled, and annotated the genomes of *P. estuarinus* and *P. kaitunuparaoa*. The *P. estuarinus* genome was assembled using a combination of short- and long-read sequences (Illumina and PacBio), whereas the *P. kaitunuparaoa* genome was constructed using only Illumina short-read sequences. Our analyses of these genomes demonstrate considerable similarity in genome size, structure, and gene content, including a conserved set of

genes involved in meiosis. Phylogenetic analyses of both mitochondrial and nuclear genomes indicate a strongly supported sister relationship between *P. antipodarum* (characterized by transcriptome resources) and *P. kaitunuparaoa*, but also reveal a substantial amount of introgression of nuclear genes between *P. estuarinus* and *P. kaitunuparaoa*. We describe and interpret these major findings in detail below.

Genome Assembly, Annotation, and Assessment

Analyses from unassembled short reads indicate that these *Potamopyrgus* species have similar genomic characteristics. We estimated the haploid genome size for both *P. estuarinus* and *P. kaitunuparaoa* at around 500 Mb, with heterozygosity of 3.73% for *P. estuarinus* and 3.63% for *P. kaitunuparaoa* (supplementary fig. S2, Supplementary Material online). Our assembled genome of *P. estuarinus* has a total length of 523.89 Mb ($N = 12,267$ total contigs), an N50 of 80 Kb ($N = 1508$), and a max contig length of 1.1 Mb. Because our *P. kaitunuparaoa* assembly is based on only short-read small-insert Illumina data, we could not obtain as contiguous of an assembly as we did for *P. estuarinus*. The inclusion of comparative scaffolding with our *P. estuarinus* resulted in the following assembly summary for *P. kaitunuparaoa*: total length of 597.46 Mb ($N = 132,386$ total scaffolds and contigs), with an N50 of 48 Kb ($N = 2456$), and a max contig length of 1.04 Mb.

The MAKER annotation pipeline produced similar results between the two species, with 32,238 protein-coding genes predicted from *P. estuarinus* and 30,311 from *P. kaitunuparaoa*. The average coding sequence length was 1,102 bp and 1,253 from the *P. estuarinus* and *P. kaitunuparaoa* annotations, respectively. The proportion of each genome assembly made up of coding sequence was 6.9% in *P. estuarinus* and 6.4% in *P. kaitunuparaoa*. The resulting functional annotations from the BLAST2GO pipeline were also similar between the two species. BLAST hits to the NCBI nr database were obtained for 25,299 (78%) of the *P. estuarinus* genes and 24,964 (82%) of the *P. kaitunuparaoa* genes. Similarly, InterPro functional annotations were found for 26,155 (81%) and 24,928 (82%) genes for *P. estuarinus* and *P. kaitunuparaoa*, respectively. GO terms were applied to 21,482 (66%) of the *P. estuarinus* genes and 20,584 (68%) of the *P. kaitunuparaoa* genes. Together, 19,984 (62%) genes predicted from *P. estuarinus* and 19,220 (63%) from *P. kaitunuparaoa* were annotated with functional information from BLAST against the nr database, GO IDs, and InterPro IDs.

Following the creation of initial genome assemblies and the resulting annotations for *P. estuarinus* and *P. kaitunuparaoa*, we used BUSCO v.5.4.2 (Manni et al. 2021) and the BlobToolKit pipeline (Challis et al. 2020) to evaluate biological completeness and contamination in both genomes.

Scaffolds with a majority of bacterial, viral, or fungal sequences were identified as contaminants were removed and the resulting assemblies were evaluated using BUSCO comparing both genomes to the metazoa_odb10 BUSCO reference gene set (Fig. 1, [supplementary fig. S4, Supplementary Material](#) online).

BUSCO results showed that a mean of 80.3% of metazoan markers were assembled in single copy in the *P. estuarinus* genome, with an N50 of 77.4 kb (Fig. 1a, [supplementary fig. S3a, Supplementary Material](#) online). A total of 82.9% of metazoan BUSCO markers were assembled in single copy in the *P. kaitunuparaoa* genome. Following contaminant removal, *P. kaitunuparaoa* had an N50 of 47.8 kb (Fig. 1b, [supplementary fig. S3b, Supplementary Material](#) online). Annotated transcripts from both genomes overwhelmingly showed significant similarity, as assessed by BLAST similarity scores, to mollusk or other metazoan records ([supplementary fig. S5a-d, Supplementary Material](#) online), particularly after potentially contaminating sequences were removed ([supplementary fig. S5e-h, Supplementary Material](#) online). A detailed summary of genome statistics before and after the removal of contaminants can be found in [supplementary tables S1 and S2, Supplementary Material](#) online. Importantly, as can be seen with the BUSCO scores, our assemblies are highly biologically complete and thus informative for population and phylogenetic analysis. Our estimates of genomic repeat content using dnaPipeTE were also similar across the two species, with 31.06% and 30.37% repeat portions of the *P. estuarinus* and *P. kaitunuparaoa* genome, respectively ([supplementary table S3, Supplementary Material](#) online). The estimated rDNA amounts for these two species (*P. estuarinus*—0.50%; *P. kaitunuparaoa*—0.32%, [supplementary table S3, Supplementary Material](#) online) fall within the range of rDNA content estimated from diploid sexual *P. antipodarum* (0.21% to 0.83%) using short reads to estimate repeat abundance (McElroy et al. 2021).

Comparative Genomic Assessment of Assembly and Annotation Quality

To evaluate the veracity of our gene models predicted from both *P. estuarinus* and *P. kaitunuparaoa* genomes, we inferred homologous proteins from independently assembled Caenogastropod genomic and transcriptomic resources using OrthoFinder (Fig. 2, [supplementary fig. S6, Supplementary Material](#) online). We found 27,549 homologous gene groups (i.e. groups of genes with significant pairwise BLAST similarity) that contained at least one protein from either *P. estuarinus* (representing all 32,238 *P. estuarinus* genes) or *P. kaitunuparaoa* (representing all 30,311 genes). A vast majority of homologous gene groups (103,672) were specific to either the *P. antipodarum* transcriptome or the *O. hupensis* transcriptome—these apparently

spurious genes were not considered further, as they provided no comparative information. We found 1,713 homologous gene groups that contained exactly one protein from each of *P. estuarinus*, *P. kaitunuparaoa*, *P. antipodarum*, *O. hupensis*, *M. cornuarietis*, and *Pom. canaliculata*. There were 7,731 homologous gene groups for which all species were represented by at least one protein, representing 10,681 (33%) *P. estuarinus* genes and 10,438 (34%) *P. kaitunuparaoa* genes (see Fig. 2). We also identified 6,614 gene groups that were single copy in *Potamopyrgus*, an additional 1,817 that were single copy in both *P. estuarinus* and *P. kaitunuparaoa* but multi-copy in the *P. antipodarum* transcriptome, and 12,202 homologous gene groups (representing 18,616 *P. estuarinus* genes and 19,097 *P. kaitunuparaoa* genes) that had at least one protein from all three *Potamopyrgus* assemblies (the *P. antipodarum* gene set derives from transcriptome resources). For *P. estuarinus*, we found that 26,001 (81%) of all *P. estuarinus* gene models were homologous to one or more proteins from at least one other independent assembly (i.e. excluding *P. kaitunuparaoa*). 25,150 (83%) *P. kaitunuparaoa* gene models satisfied the same criterion (i.e. excluding *P. estuarinus*). There were 1,513 homologous gene groups (representing 1,898 *P. estuarinus* genes and 1,940 *P. kaitunuparaoa* genes) that contained proteins from both *P. estuarinus* and *P. kaitunuparaoa* but no other assembly. We found 3,586 homologous gene groups (representing 4,338 genes) that were specific to *P. estuarinus* and 3,048 homologous gene groups (representing 3,220 genes) that were specific to *P. kaitunuparaoa*. These latter two groups likely represent false-positive gene annotations. There were no *P. estuarinus* or *P. kaitunuparaoa* proteins that were not assigned to homologous gene groups as a result of this process.

To evaluate the quality of gene models, we estimated the rate of synonymous (d_S) and nonsynonymous (d_N) substitutions per site for 1,682 single-copy genes (excluding 30 genes that had internal stop codons in the *Pom. canaliculata* or the *M. cornuarietis* annotations) on all three possible tree topologies (i.e. *Pa-Pk*, *Pe-Pk*, and *Pa-Pe*), plus an analysis excluding *P. antipodarum* (to alleviate challenges caused by topological variation within *Potamopyrgus*), and an analysis including only the *Potamopyrgus* species (to alleviate challenges caused by saturation). Our gene models appear to be overwhelmingly consistent with being correctly annotated, as d_N/d_S estimates for the vast majority of genes (mean $d_N/d_S \pm SD = 0.097 \pm 0.059$; [supplementary fig. S7, Supplementary Material](#) online) were low (i.e. $\ll 1$), as expected under a predominant force of purifying selection. The deep divergence between the apple snails and *Potamopyrgus* means that the observed overall synonymous substitution rates are highly saturated (mean $d_S = 4.35$ synonymous substitutions per site). We can nevertheless use the branch-specific model of codeml (i.e. model 1) to estimate d_S across the tree ([supplementary fig. S8,](#)

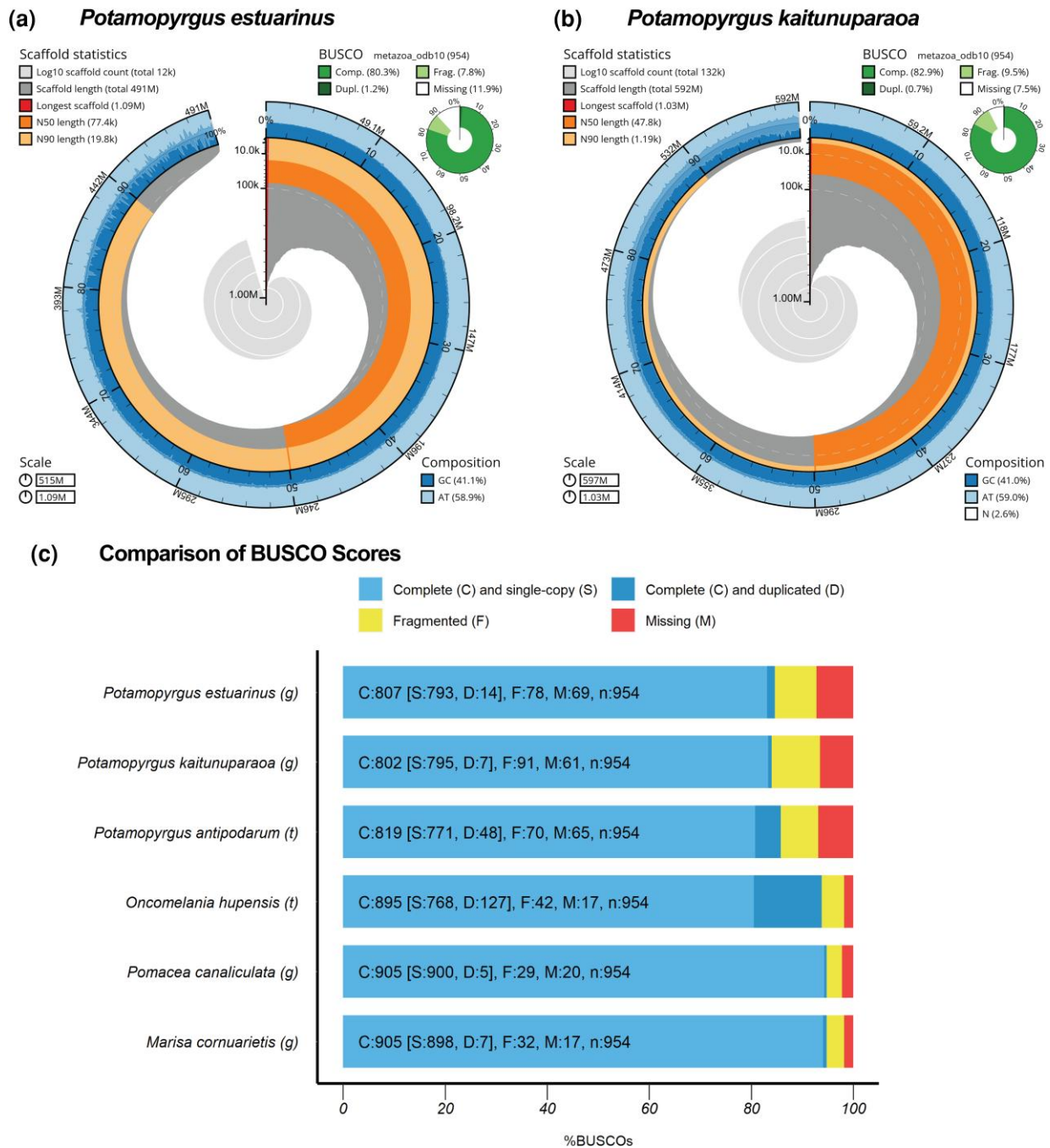


Fig. 1.—Genome assembly statistics for *P. estuarinus* and *P. kaitunuparaoa* genome assemblies. Snail plots describing genome assembly statistics for *P. estuarinus* (a) and *P. kaitunuparaoa* (b) after contaminant removal (i.e. sequences identifying with kingdoms in Bamfordvirae, Bacteria-undef, Viridiplantae or Fungi; or phylum in Pseudomonadota or Bacteroidota). The main plot in the center of each panel is divided into 1,000 size-ordered bins, with each bin representing 0.1% of the assembly (total length *P. estuarinus*—514,633,712 bp; total length *P. kaitunuparaoa*—596,959,456 bp). Sequence length distribution is shown in dark gray, with the radius scaled to the longest present sequence (red). The arcs in the plot represent the N50 sequence length (orange) and N90 sequence length (pale orange). The pale gray spiral shows the cumulative sequence count on a log scale, with orders of magnitude represented by white scale lines. GC, AT, and N percentages are reflected by the dark blue and pale blue area around the outside of the plot, in the same bins as the inner plot. (c) A summary of BUSCO genes in the metazoa_odb10 dataset present after removal of contaminants are shown in the top right of each plot. BUSCO summary statistics for whole assemblies *P. estuarinus*, *P. kaitunuparaoa*, *P. antipodarum* (transcriptome), *O. hupensis* (transcriptome), *M. cornuarietis*, and *Pom. canaliculata* are shown in panel (c).

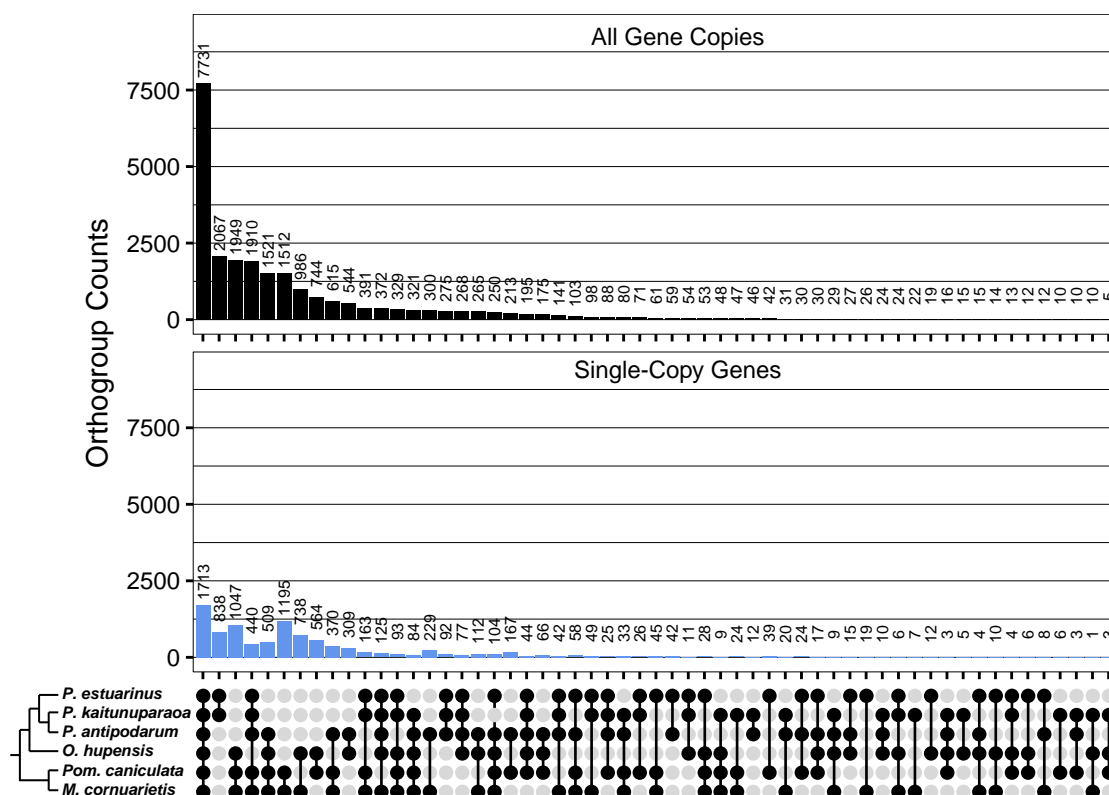


FIG. 2.—Homologous gene groups in assemblies from six Caenogastropod taxa. Upset plot depicting the number of genes fitting into homologous gene groups (group membership specified in bottom panel) for multi-copy genes (top panel) and single-copy genes (bottom panel). Taxa from top to bottom: *Potamopyrgus estuarinus*, *Potamopyrgus kaitunuparaoa*, *Potamopyrgus antipodarum*, *Oncomelania hupensis*, *Pom. canaliculata*, *M. cornuarietis*. Singleton genes that are only present in a single assembly have been removed from this plot for display purposes.

Supplementary Material online). Using this approach, and excluding 23 genes (1.4%) that were clear d_5 outliers (i.e. terminal branch d_5 values >1), we found low d_5 rates (i.e. less than 10% divergence) within *Potamopyrgus* (mode $d_5 \pm SD = 0.0783 \pm 0.10$ synonymous substitutions per site), between *P. estuarinus* and *P. antipodarum* (mode $d_5 \pm SD = 0.041 \pm 0.080$ synonymous substitutions per site), and between *P. kaitunuparaoa* and *P. antipodarum* (mode $d_5 = 0.041 \pm 0.083$ synonymous substitutions per site) assuming the *Pa-Pk* tree. Similar estimates were obtained considering the other two tree topologies. In sum, analyses of molecular evolution indicate that *P. estuarinus* and *P. kaitunuparaoa* have high-quality gene model annotations and that divergence is deep among Caenogastropod species that boast high-quality genomic resources (i.e. the six taxa represented here).

Analysis of Gene Families Associated With Meiotic Function

We inventoried and manually annotated the same, presumed orthologous, 44 meiosis-related genes in three different molluscan gastropod species (*O. hupensis*,

M. cornuarietis, and *Pom. canaliculata*) in addition to the three focal *Potamopyrgus* species, to characterize the gain or loss of these genes in *Potamopyrgus* across a larger phylogenetic framework and to provide a curated assessment of our automated gene annotations. The presence of intact (complete and in-frame CDS) meiosis genes in both focal *Potamopyrgus* taxa indicates the capacity for sexual reproduction is maintained. This finding is of considerable importance in light of the coexistence of obligately asexual lineages that characterizes *P. antipodarum*. Establishing a repertoire of genes involved in sexual reproduction (for sexual lineages) is a necessary prerequisite before focused comparisons across sexual and asexual lineages can take place (Schurko and Logsdon 2008). The meiosis gene comparisons revealed several instances of taxa-specific gene loss and retention (Table 1). First, 17 of the 44 genes are present in the genome or transcriptome assemblies across all six species, and three genes (*CYCA*, *TIM2*, and *RECQ3*) were absent from all six species. Three of the 44 genes were absent in all three *Potamopyrgus* species but present in all three other mollusks (*SEPARASE*, *RECQ2*, and *RECQ4*). *HOP1*, a meiosis-specific gene, is present

Table 1

BLAST comparisons of the aligned portions of MAKER- and hand-annotated models of meiosis genes in the *P. estuarinus* and the *P. kaitunuparaoa* genome assemblies

Gene	<i>P. estuarinus</i>					<i>P. kaitunuparaoa</i>				
	Length	# Nt Diffs	% ID	# Gaps (bp)	% Gaps	Length	# Nt Diffs	% ID	# Gaps (bp)	% Gaps
CDC25 ^a	1,800	0	100	0	0	2,961	0	100	12	1
CYCB	1,329	0	100	0	0	1,329	0	100	0	0
CYCB3	1,329	0	100	0	0	1,425	0	100	0	0
DMC1 ^b	1,020	0	100	0	0	1,041	0	100	21	2
EME1	981	0	100	0	0	981	0	100	0	0
HOP2	498	0	100	0	0	456	0	100	0	0
MLH1 ^c	606	0	100	0	0	525	6	99	0	0
MND1	462	0	100	0	0	552	0	100	0	0
MNS1 ^d	1,416	0	100	0	0	1,167	0	100	26	1
MSH2 ^e	2,628	0	100	0	0	2,808	0	100	84	3
MSH4 ^f	2,259	0	100	0	0	4,578	12	99	40	1
MSH5 ^g	1,302	90	95	27	2	246	0	100	19	1
MSH6 ^h	936	0	100	0	0	1,377	0	100	39	2
MUS81 ^{i,j}	3,477	0	100	6	1	3,303	1	99	0	0
PLK1 ^k	1,797	0	100	0	0	1,797	1	99	0	0
PLK4 ^l	3,204	0	100	0	0	3,198	0	100	12	1
RAD21 ^m	1,119	0	99	9	1	3,792	0	100	24	2
RAD51	822	0	100	0	0	843	0	100	0	0
RAD51 ^{n,o}	1,176	0	100	27	4	873	1	99	0	0
RAD51D	855	0	100	0	0	1,011	0	100	0	0
RAD54	2,058	0	100	0	0	2,271	0	100	0	0
REC8	624	35	93	18	4	740	1	99	0	0
RECQ1 ^p	2,124	0	100	36	4	2,130	0	100	45	3
SCP3	723	0	100	0	0	723	0	100	0	0
SMC1 ^q	3,681	0	100	0	0	3,888	0	100	51	2
SPO11 ^r	1,017	0	100	0	0	987	1	99	1	1
XRCC2 ^s	885	0	100	6	1	2,259	0	100	0	0
XRCC3 ^t	447	37	91	3	1	444	0	100	0	0
ZIP1 ^u	1,176	0	100	0	0	846	5	99	3	1

^a*P. antipodarum* transcript (used as bait) missing 12 bp compared to *P. kaitunuparaoa* transcript (MAKER).

^b*P. antipodarum* transcript (used as bait) missing 21 bp compared to *P. kaitunuparaoa* transcript (MAKER).

^cMAKER split *P. kaitunuparaoa* transcript into four separate genes.

^dMAKER transcript from *P. kaitunuparaoa* includes non-CDS sequence (i.e. gap/missing sequence in *P. antipodarum* transcript UTR).

^e*P. antipodarum* transcript (used as bait) missing 42 bp compared to *P. kaitunuparaoa* transcript (MAKER), while *P. kaitunuparaoa* transcript missing additional 42 bp compared to *P. antipodarum* transcript.

^fMAKER transcript from *P. kaitunuparaoa* includes non-CDS sequence (i.e. gap/missing sequence in *P. antipodarum* transcript UTR).

^gMAKER transcript from *P. estuarinus* and *P. kaitunuparaoa* includes non-CDS sequence (i.e. gap/missing sequence in *P. antipodarum* transcript UTR).

^h*P. antipodarum* transcript (used as bait) missing 39 bp compared to *P. kaitunuparaoa* transcript (MAKER).

ⁱ*P. antipodarum* transcript (used as bait) missing 12 bp compared to *P. estuarinus* transcript (MAKER).

^jMAKER transcript from *P. kaitunuparaoa* has a single-nucleotide difference compared to annotated model.

^kMAKER transcript from *P. kaitunuparaoa* has a single-nucleotide difference compared to annotated model.

^l*P. antipodarum* transcript (used as bait) missing 39 bp compared to *P. kaitunuparaoa* transcript (MAKER).

^m*P. antipodarum* transcript (used as bait) missing 9 bp compared to *P. estuarinus* transcript (MAKER) and 24 bp compared to *P. kaitunuparaoa* transcript (MAKER).

ⁿ*P. antipodarum* transcript (used as bait) missing 27 bp compared to *P. estuarinus* transcript (MAKER).

^oMAKER transcript from *P. kaitunuparaoa* has a single nucleotide difference compared to annotated model.

^p*P. antipodarum* transcript (used as bait) missing 36 bp compared to *P. estuarinus* transcript (MAKER) and 45 bp compared to *P. kaitunuparaoa* transcript (MAKER).

^qMAKER transcript from *P. kaitunuparaoa* missing 51 bp compared to *P. antipodarum* transcript (used as bait).

^rMAKER transcript from *P. kaitunuparaoa* includes non-CDS sequence (i.e. gap/missing sequence in *P. antipodarum* transcript UTR).

^s*P. antipodarum* transcript (used as bait) missing six bp compared to *P. estuarinus* transcript (MAKER).

^tMAKER split *P. estuarinus* transcript into two separate genes.

^uMAKER annotation of *P. kaitunuparaoa* transcript includes premature stop codons.

only in *Pom. canaliculata* and *M. cornuarietis*. *RAD51D*, a gene involved in homologous recombination and DNA repair, was found in all three *Potamopyrgus* taxa but was absent (or undetectable) from the other three species. That most, but not all, meiosis genes are present for each species

and are not pseudogenes is not surprising in light of previous reports of taxon-specific meiosis gene loss in *bonafide* sexual species (Malik et al. 2008; Schurko and Logsdon 2008). In the current study, at least some of the apparent gene absences are likely a result of genome assemblies

not correctly assembling the reads containing the genes, or in the case of the *O. hupensis* transcriptome, the absent genes were not being expressed in sufficient quantity to be captured by RNA sequencing. In short, the complement of meiosis genes in *Potamopyrgus* provides a framework for further investigation into how genes essential to sex evolve in an asexual background.

Another fundamental expectation for the loss of sex is that while genes critical for sex are under purifying selection and are evolving at similar rates in closely related sexual species, this constraint should be lifted for asexual lineages, resulting in a shift to relaxed selection for those lineages. Relaxed selection can act to degrade and render useless genes that are redundant in the absence of sex. That the majority of meiosis-specific genes remain intact in sexual *Potamopyrgus* species (Table 2) indicates that meiosis—as part of sexual reproduction—has been maintained and these genes are identifiable and retrievable. Indeed, for future studies of relaxed selection in the meiosis genes of asexual lineages, it is critical to establish that these genes are ancestrally operating under purifying selection because other forms of selection can be difficult to distinguish (e.g. differentiating relaxed selection from positive selection in asexuals; see Wertheim et al. 2015).

Here, we establish that the meiosis genes of closely related and sexual *Potamopyrgus* species are highly conserved and nearly identical, establishing the context for which asexual lineages can be compared in the future. We achieved this goal by analyzing the evolution of the 16 meiosis genes that are present in all *Potamopyrgus* and outgroup species. Because the phylogenetic distances between *O. hupensis* and the *Potamopyrgus* species are large (see supplementary fig. S8, Supplementary Material online), and gene comparisons may be close to or at saturation, we assess both the synonymous substitution rates and protein sequence divergence. We found that all three *Potamopyrgus* taxa are statistically identical concerning substitutions ($P=0.993$ for synonymous sites; $P=0.999$ for amino acid distance) when each is compared with the most closely related outgroup, *O. hupensis* (Fig. 3). Synonymous distances are possibly saturated for any *O. hupensis* to *Potamopyrgus* comparison due to uncorrected medians ranging from 0.701 to 0.712, where a theoretical synonymous distance of 0.75 is functionally indistinguishable from a random alignment of nucleotides. However, saturation does not substantially influence alignments at the amino acid level, where medians range from 0.152 to 0.155, highlighting the conserved nature of meiosis genes at the protein, but not nucleotide, level. Our findings align with the expected outcome of meiosis gene evolution in sexual species and now provide a foundation for future studies regarding shifts from purifying to relaxed selection for genes important to sex in *P. antipodarum*.

We also leveraged the hand-curated meiosis gene alignments as a suitable point of comparison for MAKER: the

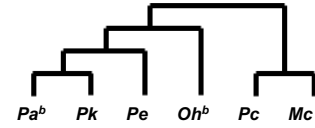
genome-wide, automated pipeline that we used for annotation. Automated gene annotation pipelines can produce erroneous annotation, particularly in genomes with complex structures (e.g. polyploidy) and significant repeat content (Bakke et al. 2009; Salzberg 2019; Ko et al. 2022). Comparing the carefully hand-annotated meiosis genes to the MAKER versions of those genes provides an estimate of the reliability and accuracy of our automated annotations. We find that MAKER transcripts in *P. estuarinus* had 21/29 genes that were indistinguishable from their hand-curated counterpart. Five genes had complete and intact reading frames, but MAKER introduced an in-frame sequence (nucleotide insertions that comprised a multiple of three and did not create a premature stop codon) that was not included in the manual annotation. Two genes were split into two separate transcripts, and one gene included non-CDS at either end of the gene sequence while staying in-frame within the gene. The MAKER transcripts in *P. kaitunuparaoa* had 14/29 genes that were indistinguishable from their hand-curated counterpart. Eight genes had complete and intact reading frame but also featured instances of both insertion and deletion of nucleotides with respect to the query. One gene model had been split into separate transcripts, and four gene models included non-CDS at the ends of the genes (see Table 1 for details). In summary, the MAKER annotations reasonably matched the hand annotations for most meiosis genes, suggesting that the automated approach produced fairly accurate annotations.

Mitochondrial Genome Assembly and Phylogenetic Analysis

Potamopyrgus mitochondrial genomes are comprised of a large single-copy region and a small single-copy region interspersed by a pair of inverted repeats (Sharbrough et al. 2023). As a result, the noncoding region (~1.7 kb in length) thought to function as the control region in *Potamopyrgus* is highly repetitive and intractable with short-read-only approaches. We lacked long reads for *P. kaitunuparaoa* so we were unable to assemble the complete mitochondrial genome. Still, we were able to assemble the large single-copy region (~15.1 kb in length) for two *P. kaitunuparaoa* and two additional *P. estuarinus* individuals. In these short-read assemblies, we found and annotated all 37 genes (13 protein-coding genes, 22 tRNAs, two rRNAs) that are present on the complete and circular reference mitochondrial genome assemblies of *P. antipodarum* (NC_070577) and *P. estuarinus* (NC_070576), and used this region to evaluate phylogenetic relationships between *Potamopyrgus* species from the mitochondrial perspective, using *O. hupensis* as an outgroup (Fig. 4). The mitochondrial genome tree provided strong support for a sister relationship between *P. antipodarum* and *P. kaitunuparaoa*, to the exclusion of *P. estuarinus*.

Table 2

Gene inventory for 44 meiosis genes in six gastropod species



Functional Role	Gene ^a	Pa ^b	Pk	Pe	Oh ^b	Pc	Mc	
Cell cycle control	<i>CDC25^c</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CDK1</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CDK2</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CORT</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CYC A</i>	Black	Black	Black	Black	Black	Black	
	<i>CYC B</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CYC B3</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CYC D</i>	Gray	Black	Black	Black	Black	Gray	
	<i>CYC E</i>	Gray	Black	Black	Black	Black	Gray	
	<i>FZY</i>	Gray	Black	Black	Black	Black	Gray	
	<i>PLK1</i>	Gray	Black	Black	Black	Black	Gray	
	<i>PLK4</i>	Gray	Black	Black	Black	Black	Gray	
	Chromosomal structural maintenance	<i>RAD21</i>	Gray	Black	Black	Black	Black	Gray
		<u>REC8</u>	Gray	Black	Black	Black	Black	Gray
<i>SEPARASE</i>		Black	Black	Black	Black	Black	Black	
<i>SMC1</i>		Black	Black	Black	Black	Black	Black	
<i>TIM2</i>		Black	Black	Black	Black	Black	Black	
Meiotic recombination	<u>DMC1</u>	Gray	Black	Black	Black	Black	Gray	
	<i>EME1</i>	Gray	Black	Black	Black	Black	Gray	
	<u>HOP1</u>	Black	Black	Black	Black	Black	Black	
	<u>HOP2</u>	Black	Black	Black	Black	Black	Black	
	<i>MLH1</i>	Gray	Black	Black	Black	Black	Gray	
	<u>MND1</u>	Gray	Black	Black	Black	Black	Gray	
	<i>MNS1</i>	Gray	Black	Black	Black	Black	Gray	
	<i>MSH2</i>	Gray	Black	Black	Black	Black	Gray	
	<u>MSH4</u>	Black	Black	Black	Black	Black	Black	
	<u>MSH5</u>	Gray	Black	Black	Black	Black	Gray	
	<i>MSH6</i>	Gray	Black	Black	Black	Black	Gray	
	<i>MUS81</i>	Gray	Black	Black	Black	Black	Gray	
	<i>PMS1</i>	Gray	Black	Black	Black	Black	Gray	
	<i>RAD51</i>	Black	Black	Black	Black	Black	Black	
	<i>RAD51C</i>	Gray	Black	Black	Black	Black	Gray	
	<i>RAD51D</i>	Gray	Black	Black	Black	Black	Gray	
	<i>RAD54</i>	Gray	Black	Black	Black	Black	Gray	
	<i>RECQ1</i>	Gray	Black	Black	Black	Black	Gray	
	<i>RECQ2</i>	Black	Black	Black	Black	Black	Black	
	<i>RECQ3</i>	Black	Black	Black	Black	Black	Black	
	<i>RECQ4</i>	Black	Black	Black	Black	Black	Black	
	<i>RECQ5</i>	Gray	Black	Black	Black	Black	Gray	
<i>SCP3</i>	Gray	Black	Black	Black	Black	Gray		
<u>SPO11</u>	Gray	Black	Black	Black	Black	Gray		
<i>XRCC2</i>	Gray	Black	Black	Black	Black	Gray		
<i>XRCC3</i>	Gray	Black	Black	Black	Black	Gray		
<i>ZIP1</i>	Gray	Black	Black	Black	Black	Gray		

^aBold, underlined genes are meiosis-specific.

^bPresence/absence inferred from a transcriptome in this species.

^cPresence indicated by gray box, absence indicated by black box.

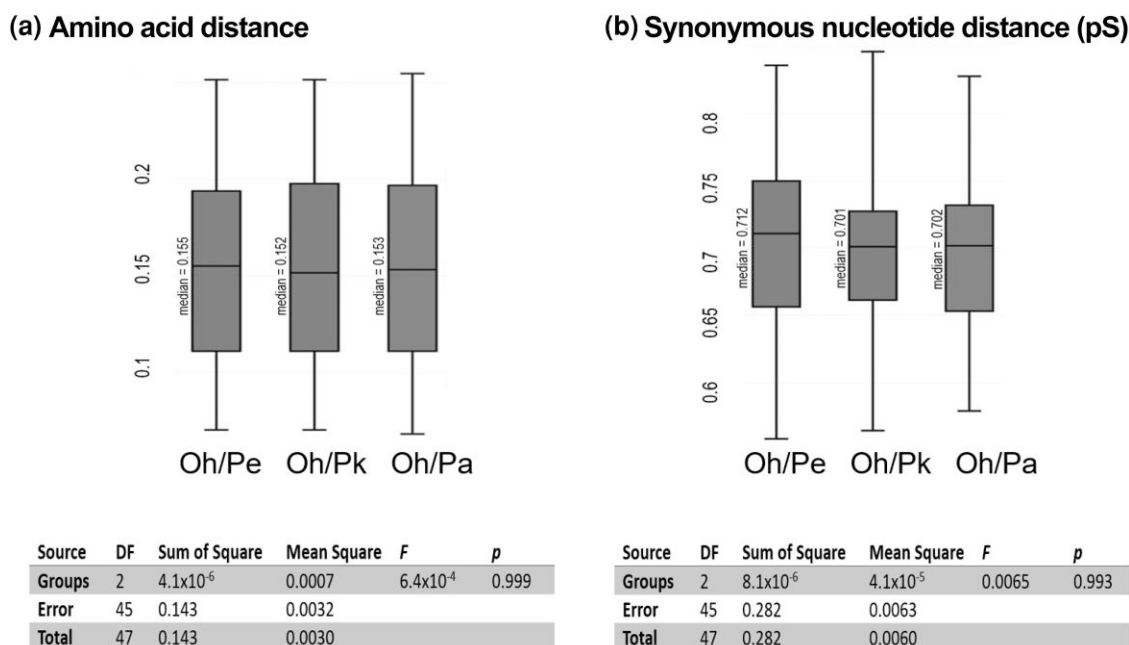


FIG. 3.—Meiosis gene evolution among *Potamopyrgus* species. (a) Pairwise comparison of amino acid distance from each *Potamopyrgus* species (*P. antipodarum*, Pa; *P. estuarinus*, Pe; *P. kaitunuparaoa*, Pk) to single outgroup *O. hupensis* (Oh) ($P=0.999$). (b) Pairwise synonymous nucleotide distance (pS) from each *Potamopyrgus* species to *O. hupensis* ($P=0.995$). Boxplot whiskers extend to minimum and maximum values; box contains Q1 to Q3 and with the median represented by a black line. Statistical significance was assessed using a one-way Anova and Tukey's range test ($\alpha=0.05$). Only meiosis genes with all species represented in single-copy form were used in this analysis.

Signatures of Introgression Across the Nuclear Genomes of *P. estuarinus* and *P. kaitunuparaoa*

Phylogenetic analysis using whole mitochondrial genomes revealed that *P. kaitunuparaoa* appears to be sister to *P. antipodarum*, to the exclusion of *P. estuarinus* (the Pa-Pk tree) (Fig. 4). This pattern was also the most common tree topology inferred in gene trees from 1,639 single-copy nuclear genes (739 gene trees, or 45.1%). However, a sizable fraction of individual gene trees (478, or 29.2%) instead support a sister relationship between *P. kaitunuparaoa* and *P. estuarinus*, to the exclusion of *P. antipodarum* (the Pe-Pk tree). The tree topology featuring *P. estuarinus* and *P. antipodarum* as sister and with *P. kaitunuparaoa* as outgroup (the Pa-Pe tree) was rarer than either of the other two tree topologies (422 gene trees, or 25.7%), likely reflecting baseline levels of incomplete lineage sorting (ILS). The difference in occurrence between the Pe-Pk and the Pa-Pe trees was even more pronounced when only gene trees with bootstrap support values ≥ 80 were considered (Pe-Pk gene trees = 163, Pa-Pe gene trees = 128, binomial test, $P=0.0461$). We also quantified the three tree topologies in 8,077 1:1:1 triplets that were pruned out of larger orthogroup gene trees to test whether this pattern held in a larger set of genes. Midpoint rooting of triplet gene trees yielded a similar result as above, with 3,326 gene trees supporting the Pa-Pk topology (41.2%), 2,658

gene trees supporting the Pe-Pk topology (32.9%), and 2,093 gene trees supporting the Pa-Pe topology (25.9%). Considering only those gene trees with high bootstrap support (i.e. ≥ 80), this pattern was even more pronounced, with 1,566 gene trees supporting the Pa-Pk topology (45.6%), 1,169 gene trees supporting the Pe-Pk topology (34.1%), and only 697 gene trees supporting the Pa-Pe topology (20.3%).

The discrepancy in abundance between the two rarer gene tree topologies (i.e. topology in which *P. estuarinus* and *P. kaitunuparaoa* are sister taxa [Pe-Pk] is much more common than that in which *P. estuarinus* and *P. antipodarum* are sister taxa [Pa-Pe]) is contrary to the expectations of ILS alone (but not conclusive as to the source of phylogenetic discordance). Overall, these observations led us to hypothesize that some degree of introgression had occurred between these three species. We tested this hypothesis using the commonly employed D-statistic, both in its traditional implementation in a four-taxon arrangement (Durand et al. 2011), with *O. hupensis* acting as outgroup, as well as in the more recently developed three-taxon method (Hahn and Hibbins 2019). Both the four-taxon method ($n=1,639$ 1:1:1:1 genes, $D=0.1707$, 95% CIs: (0.0472 to 0.2849), $Z=-2.783$, $P=0.0054$) and the three-taxon method ($n=8,077$ 1:1:1 genes, $D_3=0.0514$, 95% CIs: (0.0383 to 0.0629), $Z=-8.145$, $P<0.001$) identified significant asymmetry across the two

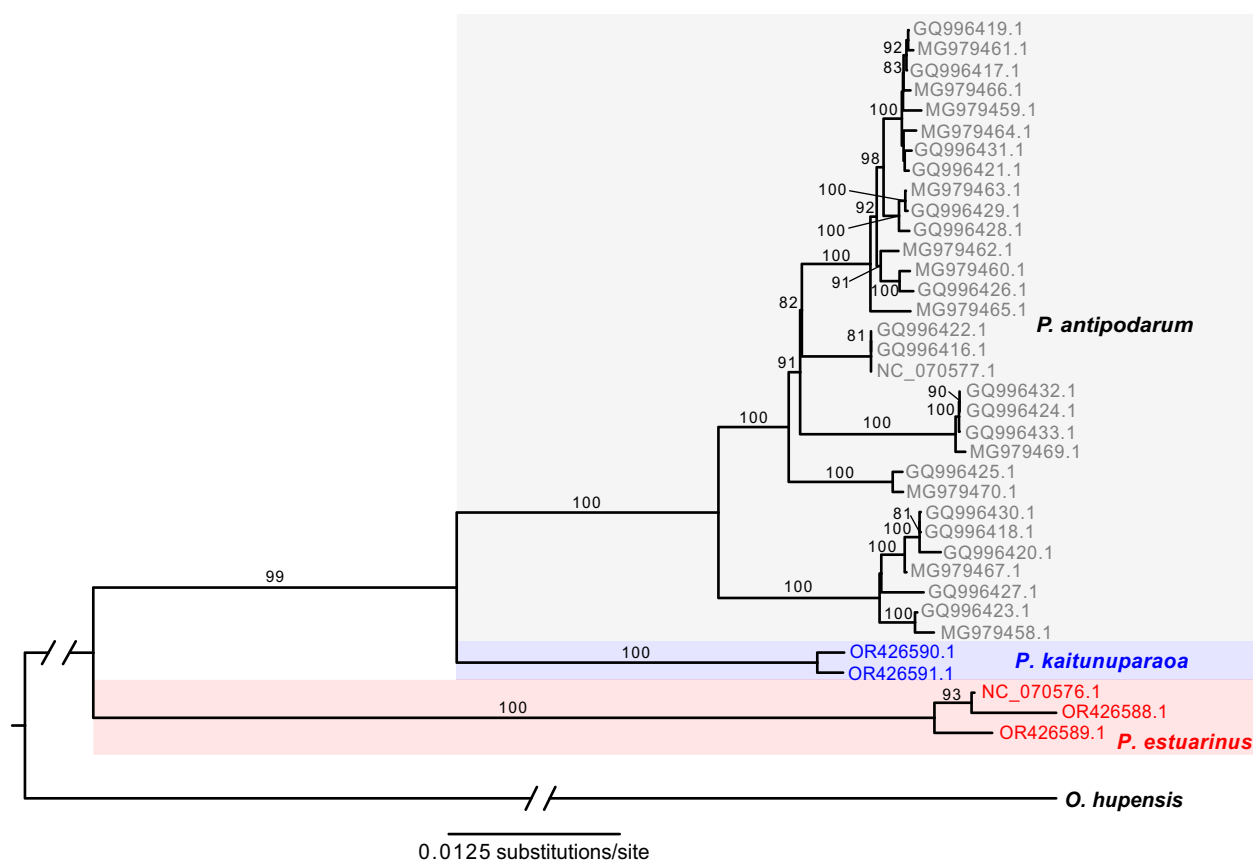


Fig. 4.—Mitochondrial phylogenetic tree inferred by Maximum Likelihood. *Potamopyrgus kaitunuparaoa* (blue) is sister to *P. antipodarum* (gray) to the exclusion of *P. estuarinus* (red) according to mitochondrial genome data. *Oncomelania hupensis* was used as an outgroup in this analysis.

least-common tree topologies, indicating that the distribution of gene tree topologies could not be explained by ILS alone (i.e. some form of introgression appears to have occurred). The result was similar for all three trimming strategies (i.e. untrimmed, ClipKIT-trimmed, and GBLOCKS-trimmed) and all three tree topologies (i.e. *Pa-Pk*, *Pe-Pk*, and *Pa-Pe*), indicating that some degree of introgression between these taxa appears to have occurred, with the identities of the taxa that had experienced introgression depending upon the tree topology assumed.

To characterize the apparent patterns of introgression, it was therefore necessary to establish the correct species tree. We reasoned that post-speciation introgression would be expected to result in the following patterns: (i) the introgression tree would be expected to have the shortest patristic distance from outgroup to ingroup compared to the species tree and the ILS tree (species = ILS \gg introgression), (ii), the introgression trees should be shorter overall than species tree or than the ILS tree if there has been introgression from the ingroup to the outgroup (species = ILS \gg introgression), and (iii) the branch leading to the midpoint-root-selected outgroup (i.e. excluding terminal branches of the ingroup) should be shorter in

unrooted introgression trees than in unrooted species or ILS trees (species = ILS \gg introgression). In agreement with these predictions, we assigned tree topology for single-copy 1:1:1 triplet trees using midpoint rooting and found that outgroup-to-ingroup patristic distance was significantly lower in *Pe-Pk* trees than in *Pa-Pk* and *Pa-Pe* trees (supplementary fig. S9, Supplementary Material online; pairwise Mann–Whitney *U* tests with Holm correction, $P < 0.0001$). Similarly, total tree length was significantly shorter in *Pe-Pk* triplet trees (median = 0.0434 changes per site) than in *Pa-Pk* or *Pa-Pe* trees (pairwise Mann–Whitney *U* tests with Holm correction, $P < 0.0001$; supplementary fig. S10, Supplementary Material online), and outgroup branch length was shorter in *Pe-Pk* trees than in *Pa-Pk* or *Pa-Pe* trees (pairwise Mann–Whitney *U* tests with Holm correction, $P < 0.0001$; supplementary fig. S11, Supplementary Material online). Thus, all three analyses indicate that the mitochondrial tree topology appears to reflect the true species branching order, and that introgression between *P. estuarinus* and *P. kaitunuparaoa* appears to have occurred more recently in the past than the split between the freshwater (*P. kaitunuparaoa* and *P. antipodarum*) and estuarine (*P. estuarinus*) species.

We also tested whether a ghost lineage (i.e. an unsampled/extinct *Potamopyrgus* lineage) could be responsible for this signature of introgression. If *Pe-Pk* trees are the result of introgression between *P. antipodarum* and an unsampled ghost lineage rather than introgression between *P. estuarinus* and *P. kaitunuparaoa*, then we would expect a similar distance between *P. estuarinus* and *P. kaitunuparaoa* in both *Pa-Pk* and in *Pe-Pk* gene trees. Instead, we found that the distance between *P. estuarinus* and *P. kaitunuparaoa* is substantially and significantly reduced in *Pe-Pk* trees compared to *Pa-Pk* trees (Mann–Whitney $U = 3482938$, $P < 0.001$, [supplementary fig. S12, Supplementary Material](#) online), which is inconsistent with a ghost lineage driving the high preponderance of *Pe-Pk* trees. As such, we can conclude that the true species branching order reflects a sister relationship between *P. antipodarum* and *P. kaitunuparaoa*, with some degree of bidirectional introgression occurring between *P. estuarinus* and *P. kaitunuparaoa*.

Although the majority gene tree does appear to be the true species branching order in these species, a sizeable fraction of the *P. estuarinus* and *P. kaitunuparaoa* genomes exhibit an evolutionary history incongruent with that of the mitochondrial genome. Because interactions between the nuclear genome and the mitochondrial genome are likely to contribute to patterns of co-introgression (Beck et al. 2015) or co-non-introgression (Sharbrough et al. 2017) of nuclear-encoded genes whose products are targeted to the mitochondria (i.e. N-mt genes), we performed enrichment analyses to test whether N-mt genes are enriched for the mitochondrial topology. We used reciprocal best BLASTp hits against the *Drosophila melanogaster* proteome to identify 319 genes in *P. estuarinus* and 293 genes in *P. kaitunuparaoa* that appear to be involved in mitochondrial-nuclear enzyme complexes (i.e. OXPHOS complexes I, III, IV, V, the mitoribosome, and mt-aaRS), of which 121 genes were represented in our set of 8,077 1:1:1 triplets ([supplementary table S4, Supplementary Material](#) online). Among these 121 genes, 59 had gene tree topologies matching the species tree (i.e. *Pe-Pk*), 32 had gene tree topologies that matched the introgression tree (i.e. *Pe-Pk*), and 30 genes matched the ILS topology (i.e. *Pa-Pe*), a pattern that was not different from the genome-wide pattern (Fisher's Exact Test $P = 0.1944$; [Table 3](#)). When we restricted the analysis to include only the 45 gene trees with ≥ 80 bootstrap support, we found significantly fewer *Pe-Pk* trees than the genome-wide patterns would predict (Fisher's Exact Test $P = 0.0043$; [supplementary table S5, Supplementary Material](#) online), potentially indicating that mito-nuclear interactions represent a barrier to gene flow between *P. estuarinus* and *P. kaitunuparaoa*. Investigating individual enzyme complexes, only Complex I (i.e. NADH ubiquinone oxidoreductase) exhibited patterns of gene tree topologies that were significantly different from genome-wide patterns,

Table 3

Gene tree topologies in nuclear-encoded mitochondrially targeted genes by enzyme class

Enzyme Category	<i>Pa-Pk</i>	<i>Pe-Pk</i>	<i>Pa-Pe</i>	FET <i>P-value</i> ^a
Complex I-NADH ubiquinone oxidoreductase	9	1	1	0.0362*
Complex II-Succinate dehydrogenase ^b	1	1	0	1
Complex III-Ubiquinol-cytochrome C oxidoreductase	3	1	0	0.4683
Complex IV-Cytochrome c oxidase	14	12	10	0.9549
Complex V-F ₀ -F ₁ ATP synthase	5	3	3	1
Mitoribosome	25	14	15	0.5413
aaRS	2	0	1	0.4881
Total	58	31	30	0.1944
Rooted Triplets	3326	2658	2093	...

^aFisher's Exact Test was performed against the genome-wide topology counts for rooted triplets, with *P*-values adjusted using the Holm procedure for the Bonferroni correction for multiple comparisons.

^bComplex II is entirely nuclear encoded and does not feature any direct interactions between mitochondrially encoded and nuclear-encoded genes or gene products. This complex is included here as a negative control and excluded from the Total.

*Significant at the $P < 0.05$ level.

with nine genes (82%, Fisher's Exact Test $P = 0.0362$) featuring the *Pa-Pk* topology, and only one gene each that exhibited the *Pe-Pk* and *Pa-Pe* topologies ([Table 3](#)). All three Complex I genes with ≥ 80 bootstrap support exhibited the species topology. Notably, mitochondrially encoded subunits of Complex I exhibit the highest degree of divergence between *P. estuarinus* and *P. kaitunuparaoa* ($\sim 12.5\%$) of all the mitochondrially encoded gene products ([supplementary table S6, Supplementary Material](#) online), potentially indicating that increased mitochondrial genome divergence may have acted as a barrier to interspecific gene flow between *P. estuarinus* and *P. kaitunuparaoa*.

This finding echoes recent work in flies (Camus et al. 2023) and fish (Moran et al. 2024) that Complex I appears to play an outsized role in hybrid incompatibilities. Because Complex I is the largest single contributor of reactive oxygen species (ROS) to the cell (Murphy 2009; Quinlan et al. 2013; Mallay et al. 2019), dysfunction in this complex resulting from cytonuclear incompatibilities may result in elevated and potentially harmful ROS levels in hybrids compared to parental species (Barreto and Burton 2013; Chang et al. 2016; Du et al. 2017), providing a potential mechanism for how gene flow could be impeded specifically for nuclear-encoded genes involved in Complex I, but not for other classes of nuclear genes. The discrepancy between nuclear and mitochondrial introgression is a common phenomenon in natural populations (Sloan et al. 2017), and is likely a consequence of the maternal inheritance of mitochondrial genomes compared to the biparental

inheritance of nuclear genomes (Camus et al. 2022). That is, asymmetry in the success of hybrid matings across reciprocal pairings—termed “Darwin’s corollary to Haldane’s rule (Turelli and Moyle 2007)—can result in differential rates of introgression across nuclear versus mitochondrial genomes. Over time, differential introgression can result in mito-nuclear discordance, as was observed here. Finally, we tested whether any gene functional groups were particularly prone to introgression over others, but none of the GO terms were significantly enriched compared to background rates. Additional population-level statistics (e.g. F_{ST}) will provide a more rigorous test of whether introgression between *P. estuarinus* and *P. kaitunuparaoa* was, at least in part, adaptive.

Conclusions

The genomes of *P. estuarinus* and *P. kaitunuparaoa* have revealed key insights for understanding the evolution of these biologically interesting species. First, both genomes are similar in size and gene content. This observation provides an estimate of the ancestral state of their unique and scientifically prominent congener, *P. antipodarum*, allowing a better understanding of the recent changes in sex, invasiveness, and other derived features in this species. Second, *P. kaitunuparaoa* is clearly the closest relative to *P. antipodarum*. However, *P. estuarinus* and *P. kaitunuparaoa* have undergone considerable exchange of many nuclear genes more recently than their species divergences from each other and with *P. antipodarum*, but with a strong bias against nuclear-encoded genes with mitochondrial function. The presented study adds to evidence from many systems for the evolutionary importance of genomic incompatibilities between nuclear and mitochondrial genomes.

Materials and Methods

Sampling, Ploidy Estimation, and Sequencing

Live specimens of *P. estuarinus* were collected in 2018 from Waikuku Beach estuary in Christchurch, New Zealand, by dragging nets along the surface of the mud in shallow water (<0.5 m depth) and returned alive to the University of Iowa. The *P. kaitunuparaoa* samples were collected in the same manner in February, 2020, from the Mōkau River, New Zealand, where it is sympatric with *P. antipodarum* and *P. estuarinus* (Fig. 5).

For *P. estuarinus*, we conducted both DNA and RNA extraction for genome assembly and transcriptome analysis, respectively (McElroy et al. 2021). To reduce the sequencing of nontarget DNA, we isolated snails for several days without feeding them to ensure that they had evacuated their guts prior to being dissected for DNA extraction. We then dissected body tissue away from the shell for 22

females, snap-froze this tissue in liquid nitrogen, and (i) shipped the frozen tissue from 20 snails to the Arizona Genomics Institute for DNA extraction and sequencing on the Sequel II (Pacific Biosciences) instrument and (ii) extracted genomic DNA for Illumina sequencing from the remaining two snails using a guanidinium thiocyanate and phenol-chloroform-based extraction method (McElroy et al. 2021; Sharbrough et al. 2023).

For PacBio sequencing, DNA was extracted from 20 snails by a modified CTAB protocol optimized to deal with common snail contaminants such as mucopolysaccharides (Arseneau et al. 2017). Extracted high-molecular weight DNA was size checked on pulsed-field gel electrophoresis and the sequencing library was constructed following the manufacturer’s protocols using SMRTbell Express Template Prep kit 2.0. The final library was size selected to 20 kb using a BluePippin (Sage Science, Beverly, USA). The recovered library was quantified with Qubit HS and size checked on Femto Pulse (Agilent). Sequencing was performed on Pacbio Sequel II using all standard protocols from the manufacturer.

We also sequenced *P. estuarinus* and *P. kaitunuparaoa* genomes from individual snails using Illumina short reads. Briefly, DNA was extracted from individual snails using a CHAOS extraction method as described above (following McElroy et al. 2021; Sharbrough et al. 2023). Both sequencing libraries were prepared using a Kapa Hyper Plus preparation and sequenced with Illumina short-insert 2 × 150 bp paired-end (PE) reads on an Illumina HiSeq (*P. estuarinus*) and on a NovaSeq 6000 using S4 chemistry (*P. estuarinus* and *P. kaitunuparaoa*) at the Iowa Institute of Human Genetics (IIHG). All sequencing data, assemblies, and annotations have been deposited at NCBI under BioProject PRJNA717745.

Genome Assembly and Assessment

First, we used our Illumina short-read data to characterize the genomes of *P. estuarinus* and *P. kaitunuparaoa*. Specifically, we estimated genome size using GenomeScope 2.0 (Ranallo-Benavidez et al. 2020), heterozygosity from k-mer distributions ($k = 21$) generated by Jellyfish (v2.3.0 Marçais and Kingsford 2011), and inferred repeat content using RepeatModeler2 (Flynn et al. 2020) for both species’ genomes. This information was then used as priors to guide our genome assembly pipeline and parameters.

Genome assembly was conducted using distinct methodologies because the two focal species were sequenced using different technologies, though components of individual inputs did overlap. For *P. estuarinus*, following receipt of raw data in the form of BAM files from the Arizona Genomics Institute, we extracted FASTA and FASTQ files using the bam2fastx utility (<https://github.com/PacificBiosciences/bam2fastx>), and we used Canu

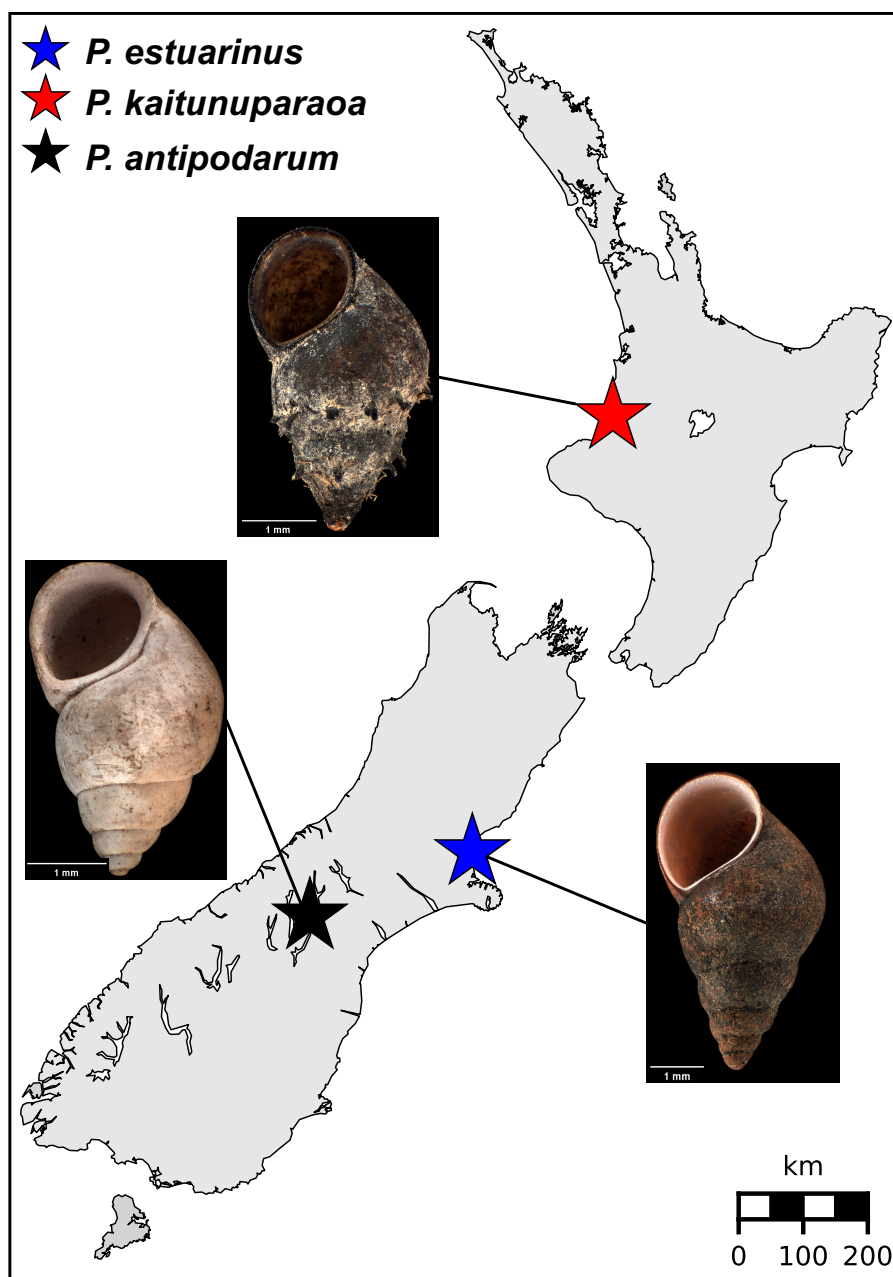


FIG. 5.—Map of New Zealand sampling locations of *P. estuarinus*, *P. kaitunuparaoa*, and *P. antipodarum*. Map locations of *P. estuarinus* (blue star), *P. kaitunuparaoa* (red star), and *P. antipodarum* (black star). Shell morphology for each species is depicted in associated images. Shell photographs were obtained from the Museum of New Zealand (Te Papa Tongarewa) and modified in accordance with the Creative Commons BY 4.0 license. Links to original photographs are provided in the Data Availability statement.

v. 1.8 (Koren et al. 2017) to assemble the first genome draft. Because the sequenced material was derived from 20 individual snail genotypes, default parameters were insufficient to generate either a contiguous or biologically complete assembly. To overcome assembly degradation issues related to large amounts of heterozygosity in the DNA pool, we used an iterative correction of the reads *in silico* to reduce the diversity in raw data. A standard assembly using Canu

will use a single iteration of correction, wherein the stochastic errors that arise during sequencing are removed through a consensus decision. We repeated the correction and subsequent assembly steps until we found an asymptote in assembly length, contiguity, and biological completeness as assessed through BUSCO v5.4.2 (Manni et al. 2021) completeness score. We conducted a total of six correction steps before this asymptote was reached, and

adding the seventh iteration did not qualitatively change the results of the three aforementioned metrics.

In the case of *P. kaitunuparaoa*, only short reads were available, so we assembled contigs using only those data. Following receipt of raw data in compressed FASTQ format from IIHG, we used FASTQC (<https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>) to evaluate read quality and detect adapter contamination. Low-quality reads and adapter contamination were removed using fastp (Chen et al. 2018). We used a second round of FASTQC to verify read trimming was successful. In order to obtain the most accurate assembly, we used both MaSuRCA and SPAdes assemblers and then merged these independent assemblies using the Purge Haplotig approach (Roach et al. 2018). In order to improve the contiguity of our assembly, we used the tool RagTag (Alonge et al. 2022) to conduct a comparative scaffolding using our much more contiguous *P. estuarinus* assembly. Gap filling for both assemblies was done using a combination of Racon (Vaser et al. 2017) and Pilon (Walker et al. 2014).

Once both genomes were assembled, we used BUSCO to reevaluate biological completeness in both genomes, compared to the metazoa_odb10 BUSCO reference gene set. To identify contigs derived from contamination, we employed the BlobToolKit v.4.1.4 (Challis et al. 2020) pipeline with default parameters. This pipeline uses NCBI blastn alignments (Altschul et al. 1997) to the NCBI nucleotide database and DIAMOND blastx alignments (Buchfink et al. 2021) to the NCBI nonredundant protein (nr) databases to assign taxonomic affiliations to each contig with an e-value of 10^{-25} . BlobToolKit also assessed genome completeness and coverage using results from BUSCO evaluation and BWA-MEM aligned sequencing reads (Li 2013). This process was done for both *P. estuarinus* and *P. kaitunuparaoa* genome assemblies and was visualized in snail- and blob-plots. To correct for contamination, scaffolds identified to be bacterial, viral, or fungal were removed from the assembly.

Genome Annotation

We used RepeatModeler2 (Flynn et al. 2020) to build models of repetitive sequence families and masked the assembly using RepeatMasker v.4.0.7 (Smit et al. 2013–2015) classifying the following transposable element families: “DNA”, “RC”, “LTR”, “LINE”, and “SINE”. We then performed three rounds of MAKER2 v.2.31.9 (Holt and Yandell 2011) annotation to predict genes in the *P. estuarinus* genome assembly. We started with evidence-based input data, followed by two iterative rounds of ab initio training based on the previous round(s) of results. As input for the first round of MAKER, we used assembled transcriptomes from three distinct *P. estuarinus* samples (female with embryos, female without embryos, and male) that we combined and

clustered with CD-HIT-EST v.4.7 (Li and Godzik 2006; Fu et al. 2012) to reduce redundancy. Additionally, we used protein models from the *Pom. canaliculata* (GCF_003073045.1) annotation (Liu et al. 2018) and translated open reading frames from *Oncomelania hupensis* (SRR1284718) and *P. antipodarum* (SRR6470170) transcriptomes. Next, we trained SNAP v.0.15.4 (Korf 2004) and AUGUSTUS v. 2.5.5 (Stanke et al. 2006) on the gene models resulting from the transcript and protein-based predictions. To train AUGUSTUS, we ran BUSCO (v.3.0.2; metazoa_odb9 ancestral group) with AUGUSTUS (2.5.5) training enabled and the “–long parameter”. We then reran MAKER, using the SNAP and AUGUSTUS gene models, then repeated the training process and ran a third and final round of MAKER following the same approach as the previous round.

In addition to the RepeatModeler-based results used on the genome assemblies, we compared the repeat content of these genomes based on the reads to account for different assembly qualities. We used dnaPipeTE (v1.3.1c; Goubert et al. 2015), which assembles consensus repeat sequences with Trinity (v2.5.1; Grabherr et al. 2011; Haas et al. 2013) and then classifies the sequences with RepeatMasker (v4.0.7) based on the Repbase libraries (20170127) and estimates repeat family genomic abundance from blastn (BLAST + v2.2.28) hits of reads against classified repeats. The dnaPipeTE pipeline uses a subsample of reads; here, we sampled reads to $0.20 \times$ genome coverage for each species (497000000 for *P. estuarinus* and 510000000 for *P. kaitunuparaoa*). Finally, we used dnaP_utils (https://github.com/clemgoub/dnaPT_utils) to compare the dnaPipeTE results for the two species.

We functionally annotated the completed genome assemblies using DIAMOND Blastx v.2.0.15 (Buchfink et al. 2021) with the following parameters: e-value = $1e-5$, max hsps = 20, very sensitive mode, and searched the predicted transcripts against NCBI’s nonredundant (nr) database. Gene Ontology (GO) terms were then assigned using Blast2GO Basic (v.5.2.5) with default parameters.

Comparative Genomic Assessment of Assembly and Annotation Quality

To evaluate the quality of our assembly and its annotations, we employed a comparative phylogenetic approach incorporating currently available Caenogastropod genomic resources: *Oncomelania hupensis* (tropical freshwater snail—transcriptome), *M. cornuarietis* (common apple snail—genome), and *Pom. canaliculata* (golden apple snail—genome). These three species are the closest related gastropods to *P. estuarinus* and *P. kaitunuparaoa* for which high-quality genomic/transcriptomic resources are publicly available. Including other gastropods allows us to determine whether the presence or the absence of a gene is unique to the *Potamopyrgus* genus or is a

common feature of the gastropods of which sequencing is available. Genomes of *M. cornuarietis* (GCA_004794655.1) and *Pom. canaliculata* (GCF_003073045.1) were downloaded from NCBI. To create the *O. hupensis* transcriptome, we downloaded paired-end Illumina RNA-seq reads for *O. hupensis* and *P. antipodarum* from the SRA database (SRR1284718 and SRR6470170, respectively) and assembled them into a transcriptome using Trinity (v2.6.6; Grabherr et al. 2011; Haas et al. 2013) with enabled (`-trimmomatic`) for read quality control. We then extracted open reading frames with TransDecoder (v5.50; <https://github.com/TransDecoder/TransDecoder>) and clustered the translated peptide sequences with CD-HIT (v4.7; Li and Godzik 2006; Fu et al. 2012). We compared biological completeness for each genome and transcriptome using BUSCO v.5.4.2 (Manni et al. 2021) and the meta-zoa_odb10 BUSCO reference gene set.

To investigate patterns of evolution within *Potamopyrgus* (and to evaluate the quality of our genomes *via* comparative genomics), we performed global analyses of phylogenetic patterns across the *P. estuarinus* and *P. kaitunuparaoa* genomes. To accomplish this goal, we used the protein predictions from the *P. estuarinus* and *P. kaitunuparaoa* MAKER genome annotations, as well as predicted proteomes from the *P. antipodarum* transcriptome (this study), the *O. hupensis* transcriptome (this study), the *M. cornuarietis* genome (Liu et al. 2018), and the *Pom. canaliculata* genome (Liu et al. 2018) to identify putative orthologs. Following the approach of Sharbrough et al. (2022), we ran OrthoFinder v2.3.8 (Emms and Kelly 2019) on primary transcripts of all six proteomes. We then aligned CDS sequences from orthologous groups (orthogroups) by translating CDS sequences into amino acids, aligning amino acid sequences with MAFFT v7.480 (Katoh and Standley 2013), and back-translating amino acid sequences into CDS sequences. Alignments were trimmed using three different methods ranging from low to high stringency: untrimmed (low stringency), ClipKIT v1.2.0 (Steenwyk et al. 2020) (medium stringency), and GBLOCKS v0.91b (Castresana 2000) (high stringency). Downstream phylogenomic methods were performed at all three stringency levels. Gene trees were inferred from both amino acid and CDS sequence data. Protein-based trees were inferred by RAxML v8.2.12 (Stamatakis 2014), using the -GTRGAMMAIX model of molecular evolution and 100 bootstrap replicates, while CDS trees used gene-specific models of molecular evolution (inferred by jModelTest2 (Darriba et al. 2012)). The maximum-likelihood tree was inferred using five random tree starts and 100 bootstrap replicates in PhyML v3.3.20211021 (Guindon et al. 2010).

To evaluate rates and patterns of evolution across orthologs and identify potentially spurious annotations, we also estimated substitution rates under a variety of parameters using the codeml package as part of the PAML

v4.9j (Yang 2007). Briefly, we employed both the model 0 (single ω value for the whole tree) and the model 1 (branch-specific ω values) branch models of codeml (i.e. using CDS sequences), for all three possible tree topologies (see Signatures of Introgression section below) for all genes. For all PAML analyses, we also set the RateAncestor parameter to 1, the getSE parameter to 1, the clean_data parameter to 1, the optimization method to 0 (all branches simultaneously), initial kappa = 2, with kappa estimated as part of the model, and initial omega = 0.4.

Mitochondrial Genome Assembly and Phylogenetic Analysis

The complete *P. estuarinus* mitochondrial genome was assembled using a combination of Illumina HiSeq, Illumina MiSeq, and PacBio reads as described in Sharbrough et al. (2023). *Potamopyrgus* mitochondrial genomes feature a long inverted repeat interspersed by a large dinucleotide repeat (Sharbrough et al. 2023), so assembling the complete sequence of the *P. kaitunuparaoa* mitochondrial genome was not possible. Instead, we assembled the nonrepetitive portion of the mitochondrial genome (i.e. the large single-copy region, see Sharbrough et al. 2023 for more details) using MEANGS v1.0 (Song et al. 2022) for two *P. kaitunuparaoa* individuals and two *P. estuarinus* individuals with the -n flag set to 0.1 (10% subsample of reads used) and otherwise default parameters. Mitochondrial genomes were first auto-annotated using MITOS2 (Donath et al. 2019) and then manually edited.

After assembling mitochondrial genomes of both species, we aligned whole mitochondrial genomes from *P. estuarinus* (NC_070576, OR426588, OR426589), *P. kaitunuparaoa* (OR426590, OR426591), *P. antipodarum* (NC_070577, GQ996416-GQ996433, MG979458-MG979467, MG979469-MG979470), and *O. hupensis* (NC_013073), and manually edited the alignment in MEGA11 (Tamura et al. 2021). We used jModelTest2 v2.1.10 (Darriba et al. 2012) to choose the best model of molecular evolution using the best-scoring AICc model, and inferred the maximum-likelihood tree using the fast hill-climbing algorithm implemented by RAxML v8.2.12 (Stamatakis 2014) assuming the GTRGAMMAIX model of molecular evolution, and assessed tree topology with 100 bootstrap replicates. Tree topology was visually inspected to determine relationships among *Potamopyrgus* species in FigTree v.1.4.4 (<https://github.com/rambaut/figtree/releases>).

Analysis of Gene Families Associated with Meiotic Function

A catalog of meiosis genes present in the genomes of sexually reproducing *P. estuarinus* and *P. kaitunuparaoa* provides the basis for studying the maintenance and/or decay of sex-specific genes in the multiple separately derived

obligately asexual lineages that characterize *P. antipodarum*. To this end, we generated and curated a meiosis gene set using both the original (Villeneuve and Hillers 2001; Schurko and Logsdon 2008) and more up-to-date (Tvedte et al. 2017; Berdieva et al. 2021) inventories of conserved meiosis genes for all six species. These inventories discriminate between genes that specifically function in meiosis and genes that are pleiotropic with respect to function outside of meiosis. A combination of NCBI database searches and previously sequenced meiosis genes (Rice 2015) were used to find homologous meiosis genes. We queried the genomes of *P. estuarinus* and *P. kaitunuparaoa* using BLASTx to locate each gene's scaffold position. We exported the sequence to Sequencher 5.1 (Gene Codes, Ann Arbor, USA) to manually annotate each gene's start codon, exons, introns, and stop codon. Each annotation was evaluated by (i) comparison to the *P. antipodarum* transcriptome and (ii) gene identity by using BLAST. Gene presence was defined as a BLAST query covering >70% of the gene and an e-value <10⁻²⁰ (as in Tvedte et al. 2017).

We also used the hand-curated meiosis gene inventory as an opportunity to assess the accuracy of transcript models generated by the automated MAKER pipeline. We compared the hand-curated models of meiosis genes to their MAKER-generated counterparts. We aligned the annotated meiosis genes derived from *P. antipodarum* transcriptomes to compare each of the 29 meiosis genes present in all three focal *Potamopyrgus* species. We generated a local BLAST database for the MAKER transcripts (for both *P. estuarinus* and *P. kaitunuparaoa*) to make these calls. Then, we used the annotated meiosis genes as queries for BLASTn sequence retrieval. Those retrieved sequences were aligned and scored for nucleotide identity and whether sequence gaps were introduced in MAKER versus annotated genes. The proportion of genes that exactly or closely match the hand-curated annotations compared with genes that have errors gives insight into the quality of the *P. estuarinus* and *P. kaitunuparaoa* genome assemblies and the accuracy of our annotation pipeline approach. To understand the phylogenetic context for meiosis gene presence or absence, we also compared meiosis gene toolkits with the transcriptomes and genomes of *O. hupensis*, *M. cornuarietis*, and *Pom. canaliculata*.

Signatures of Introgression Across the Genome

To evaluate phylogenetic relationships between the three *Potamopyrgus* species (*P. antipodarum* (Pa), *P. estuarinus* (Pe), and *P. kaitunuparaoa* (Pk)), we first extracted 1:1:1 triplets from orthogroup trees using a custom Python script (rootedTriplets.py). Because these triplet relationships were rooted by the other sequences in the alignment, we initially quantified the number of gene trees that supported each of the three possible gene tree

topologies: *Pa-Pe*, *Pa-Pk*, and *Pe-Pk* (supplementary fig. S1, Supplementary Material online). Tree topology and branch lengths were assessed using custom Python scripts that take advantage of the DendroPy package in Python (Sukumaran and Holder 2010).

The rooted triplets provided evidence of significant asymmetry among the two rarer tree classes (*Pe-Pk* and *Pa-Pe*); we reasoned that post-speciation introgression might have played a role in the evolution of these three species. Accordingly, we employed several distinct tests of introgression by taking advantage of amino acid alignments, CDS alignments, and tree branch lengths. These tests are all derived from various implementations of the D-statistic (also known as the ABBA-BABA test), which test for deviations of nonspecies-tree site patterns from 50 to 50 in four- (Durand et al. 2011) and three-taxon sequence alignments (Hahn and Hibbins 2019). Bootstrapping of D-statistics was performed at the gene level to avoid effects of pseudoreplication that result from linkage. Because ghost lineages (i.e. unsampled taxa, whether extinct or extant) can contribute to false inferences of introgression between species (Tricou et al. 2022), we also tested whether a potential ghost lineage could explain our tree topology patterns.

In order to determine whether the species tree reflected the mitochondrial gene tree (i.e. *Pa-Pk* topology) versus the second most common tree topology (i.e. *Pe-Pk* topology), we developed several predictions for our triplet trees that would enable us to differentiate between various scenarios. First, we expected the introgression tree to exhibit the shortest patristic distance from outgroup to ingroup compared to the species tree and the incomplete-lineage-sorting (ILS) tree (species = ILS ≫ introgression). Second, the overall length of the triplet trees should be shorter for the introgression trees than for the species or ILS trees (species = ILS ≫ introgression). After selecting an outgroup using midpoint rooting, the selected outgroup would be expected to have a shorter branch in unrooted introgression trees than in unrooted species or ILS trees (species = ILS ≫ introgression). We used Mann–Whitney *U* tests to compare these statistics across tree topologies to identify the true species tree. While the mitochondrial genome appeared to follow the inferred species relationship with *P. antipodarum* and *P. kaitunuparaoa* as sister taxa (*Pa-Pk*), much of the nuclear genome (i.e. introgressed regions) exhibited a more recent common ancestor between *P. kaitunuparaoa* and *P. estuarinus* (*Pe-Pk*). Because evolutionary mismatches between nuclear-encoded genes that directly interact with mitochondrially encoded genes can produce incompatibilities resulting in reduced fitness, we reasoned that such genes would be particularly resistant to introgression from *P. estuarinus* into *P. kaitunuparaoa*. To test this hypothesis, we reciprocally BLASTed *Drosophila melanogaster* proteins that are involved in mitochondrial-nuclear interactions (i.e. proteins

involved in OXPHOS Complexes I, III, IV, and V, the mitochondrion, and mitochondrial tRNA aminoacyl synthetases, manually obtained from flybase.org) against the inferred *P. estuarinus* and *P. kaitunuparaoa* proteomes to identify nuclear-encoded mitochondrially interacting genes in *Potamopyrgus*. We then tested whether tree topologies from these complexes were more likely to match the species topology (i.e. *Pa-Pk*) than the rest of the genome using a Fisher's Exact Test, correcting for multiple comparisons using the Holm procedure for the Bonferroni correction (Holm 1979).

Supplementary Material

Supplementary material is available at *Genome Biology and Evolution* online.

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Data Availability

Raw reads associated with this project are available at NCBI under BioProject PRJNA717745. Assemblies, annotations, and curated gene datasets are available at [Zenodo.org](https://zenodo.org/doi/10.5281/zenodo.10023075) (DOI:10.5281/zenodo.10023075). Shell photograph for *P. estuarinus* is available at <https://collections.tepapa.govt.nz/object/147506>. Shell photograph for *P. kaitunuparaoa* is available at <https://collections.tepapa.govt.nz/object/642730>. Shell photograph for *P. antipodarum* is available at <https://collections.tepapa.govt.nz/object/602110>. Genomic data derived from taoka (treasured) species such as freshwater

pūpū are taoka (treasures) in their own right. Accordingly, the raw reads of *Potamopyrgus kaitunuparaoa* associated with this project will be made available (via BioProject PRJNA717745 and [Zenodo.org](https://zenodo.org)) on recommendation of Ngāti Maniapoto that affiliate as kaitiaki (guardians) for this species.

Statement of Ethics

While snails are invertebrates and thus, excepted from IACUC guidelines regarding the ethical and humane treatment of "animals", we make every effort to treat the snails we study humanely and ethically.

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