Mitochondrial Genomes in *Perkinsus* Decode Conserved Frameshifts in All Genes

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Abstract

Mitochondrial genomes of apicomplexans, dinoflagellates, and chromodellids that collectively make up the Myzozoa, encode only three proteins (Cytochrome b [COB], Cytochrome c oxidase subunit 1 [COX1], Cytochrome c oxidase subunit 3 [COX3]), contain fragmented ribosomal RNAs, and display extensive recombination, RNA trans-splicing, and RNA-editing. The early-diverging Perkinsozoa is the final major myzozoan lineage whose mitochondrial genomes remained poorly characterized. Previous reports of *Perkinsus* genes indicated independent acquisition of non-canonical features, namely the occurrence of multiple frameshifts. To determine both ancestral myzozoan and novel perkinsozoan mitochondrial genome features, we sequenced and assembled mitochondrial genomes of four *Perkinsus* species. These data show a simple ancestral genome with the common reduced coding capacity but disposition for rearrangement. We identified 75 frameshifts across the four species that occur as distinct types and that are highly conserved in gene location. A decoding mechanism apparently employs unused codons at the frameshift sites that advance translation either +1 or +2 frames to the next used codon. The locations of frameshifts are seemingly positioned to regulate protein folding of the nascent protein as it emerges from the ribosome. The *cox3* gene is distinct in containing only one frameshift and showing strong selection against residues that are otherwise frequently encoded at the frameshift positions in *cox1* and *cob*. All genes lack cysteine codons implying a reduction to 19 amino acids in these genomes. Furthermore, mitochondrion-encoded rRNA fragment complements are incomplete in *Perkinsus* spp. but some are found in the nuclear DNA suggesting import into the organelle. *Perkinsus* demonstrates further remarkable trajectories of organelle genome evolution including pervasive integration of frameshift translation into genome expression.

Key words: Myzozoa, *Perkinsus*, mitochondrial genome, frameshifts, programmed ribosomal frameshifting.

Introduction

Mitochondria are morphologically distinctive double-membrane organelles best known for their role in ATP synthesis in eukaryotic cells via oxidative phosphorylation (Saraste 1999; Gray et al. 2001; Burger et al. 2003; Wang and Youle 2009; Flegontov et al. 2015; Roger et al. 2017). Mitochondria also contribute to numerous other important cellular functions including iron-sulfur (Fe-S) cluster biogenesis, anabolic metabolism (e.g., heme biosynthesis) and apoptosis (Saraste 1999; Gray et al. 2001; Burger et al. 2003; Wang and Youle 2009; Flegontov et al. 2015; Roger et al. 2017). All extant mitochondria are derived from an ancient endosymbiosis of an alpha-proteobacterium within
the eukaryotic common ancestor, and most have retained some form of the original prokaryotic genome (Roger et al. 2017). However, despite their common origin and generally conserved functions, mitochondrial genomes (mtDNAs) display a remarkable diversity of states (Gray et al. 2001; Burger et al. 2003; Smith and Keeling 2015; Gagat et al. 2017; Roger et al. 2017; Berná et al. 2021).

All mtDNAs are vastly reduced compared to the progenitor prokaryotic genomes. Retained genes typically code for components of the electron transport chain (complexes I, III, IV, and V) and the mitochondrial translation machinery, notably transfer RNAs (tRNAs) and ribosomal RNAs (rRNAs) (Lang et al. 1997; Roger et al. 2017). The protein-coding capacity of mtDNAs ranges from near 100 proteins in jakobid flagellates (Lang et al. 1997; Burger et al. 2003; Flegontov et al. 2015) to a mere two in the chrompodellid Chromera velia, with the typical number across eukaryotes being 40–50 genes (Gray et al. 2001; Burger et al. 2003; Flegontov et al. 2015; Roger et al. 2017). This surprisingly large divergence across extant mtDNAs is the consequence of both organelle function loss and ongoing endosymbiotic gene transfer to the nucleus occurring throughout eukaryotic diversification (Gray et al. 2001; Burger et al. 2003; Herrmann 2003; Roger et al. 2017). The relocated genes are translated in the cytosol and the corresponding proteins imported back into the organelle (Herrmann 2003; Maguire and Richards 2014; Roger et al. 2017). In some instances mitochondria are so derived, reduced and specialized (e.g., hydrogenosomes and mitosomes) that they lack the electron transport chain components and have lost their mtDNAs completely (Maguire and Richards 2014; Roger et al. 2017; Berná et al. 2021). The architecture of the remaining organelle genomes is primarily linear, single chromosomes, which often appear circular in sequence assemblies due to various stabilizing inverted end-structures such as terminal inverted repeats (TIRs) that cause false in-silico circularization (Bendich 1993; Cavalier-Smith 2018; Berná et al. 2021).

The Myzozoa, which comprise dinoflagellates, apicomplexans, chrompodellids and Perkinsids (Flegontov et al. 2015; Roger et al. 2017; Cavalier-Smith 2018), represent some of the most reduced and divergent mtDNAs known to date (Flegontov et al. 2015; Roger et al. 2017). They encode the smallest number of proteins for any organelle genome—Cytochrome b (COB), Cytochrome c oxidase subunit 1 (COX1), Cytochrome c oxidase subunit 3 (COX3)—with the gene for COB apparently completely lost in the chrompodellid Chromera velia (Waller and Jackson 2009; Flegontov et al. 2015; Roger et al. 2017; Berná et al. 2021). Moreover, rRNAs in myzozoan mtDNAs are highly fragmented with no evidence of re-assembly at the RNA level by splicing, and they entirely lack 5S rRNA (Waller and Jackson 2009; Feagin et al. 2012; Berná et al. 2021). The architecture of myzozoan mtDNAs is multiform and complex. For example, most apicomplexan mtDNAs that have been characterized to date have monomeric compacted linear forms, as small as 6 kb in Plasmodium (Feagin et al. 2012; Berná et al. 2021; Namisivayam et al. 2021). Toxoplasma gondii, however, has diverged from other apicomplexans with highly expanded, fragmented, but modularized mtDNAs (Flegontov et al. 2015; Namisivayam et al. 2021). Within the apicomplexan sister group, the chrompodellids, expansion of mtDNAs is also seen as heterogeneous, linear, duplicated and fragmented molecules (Slamovits et al. 2007; Waller and Jackson 2009; Jackson et al. 2012; Flegontov et al. 2015), and a similar pattern is found yet again in dinoflagellates surveyed to date (Jackson et al. 2007; Slamovits et al. 2007; Nash et al. 2008; Waller and Jackson 2009; Jackson et al. 2012; Gagat et al. 2017). This distribution of both simple and complex mtDNAs throughout Myzozoa confuses understanding of the ancestral state of the myzozoan mtDNA: was it monomeric or heterogeneous? Furthermore, dinoflagellate mtDNAs possess additional features and traits, namely: (1) extensive substitutional RNA editing, (2) trans-splicing of cox3 transcripts, (3) use of alternative start codons (also found in apicomplexans); and (4) general loss of encoded stop codons (Jackson et al. 2007; Nash et al. 2008; Waller and Jackson 2009; Jackson et al. 2012; Gagat et al. 2017; Janouskovec et al. 2017). Of particular note, cob and cox1 mRNAs are polyadenylated immediately after the region encoding the conserved C-terminus, presumably resulting in a short read-through poly-lysine tail. However, the molecular evolutionary behavior of cox3 differs in that polyadenylation consistently creates a UAA in-frame stop codon (Waller and Jackson 2009; Jackson et al. 2012).

The Perkinsozoa represents the deepest-branching lineage of the dinozoan clade diverging relatively close to the split with the sister apicomplexan/chrompodellid clade (Masuda et al. 2010; Zhang et al. 2011; Janouskovec et al. 2017). Thus, Perkinsozoa represents an early branching point of myzozoan diversity. Perkinsus spp. are the best studied representatives of Perkinsozoa owing to their importance as marine parasites and pathogens of commercial important shellfish (Villalba et al. 2007; Choi and Park 2010; Smolowitz 2013). To date, only partial coding sequences for two mitochondrial genes have been reported from Perkinsus spp.—cob and cox1—yet, these already indicate some further novelty in myzozoan mtDNAs (Masuda et al. 2010; Zhang et al. 2011; Bogema et al. 2021). These sequences indicate the recoding of the canonical UGA stop codon as an alternative code for tryptophan, which is a trait seen in other mitochondria including ciliates although not known from other myzozoans. More radical is the observation that both coding sequences contain multiple frameshifts that are apparently not resolved by editing at the mRNA stage. These frameshifts correlate with conserved glycine and proline residues, and the nucleotide sequence at these sites indicates consensus sequences for the glycine-encoded frameshifts (AGGY or TMGGY) and proline-encoded frameshifts (CCCCCT). To date, attempts to identify a cox3 coding sequence have been unsuccessful and it has been speculated as being lost (Koren et al. 2017; Bogema et al. 2021).
While a tantalizing glimpse of the mtDNA in *Perkinsus* has been provided, there are many unanswered questions for this key lineage: What is the form of their mtDNA, monomeric or heterologous? What is its complete coding capacity of *Perkinsus* mitochondria? How are the frameshifts decoded? Can *Perkinsus* inform on the likely ancestral states of the myzozoan mtDNA? To address these questions, we sequenced and assembled the mtDNA from four *Perkinsus* taxa: *Perkinsus atlanticus*, *P. olseni*, *P. chesapeaki* and *P. marinus*, defining the structures, coding capacity, and remarkable use of frameshifts in yet another form of divergent myzozoan mtDNAs.

### Results

**Perkinsus mtDNAs Comprise Unique, Circular-mapping Molecules**

To determine the form of *Perkinsus* mitochondrial genomes (mtDNAs) and its evolutionary stability we selected four taxa for good representation of this genus. A *Perkinsus* molecular phylogeny shows some persistent ambiguity in the taxonomy of this group, nevertheless the isolates used in this study all have distinct nuclear genomic sequence (data not shown) and represent diversity across this genus: *P. atlanticus* (ATCC 50984), *P. olseni* (ATCC PRA-205), *P. chesapeaki* (ATCC 50807), and *P. marinus* (ATCC 50983) (fig. 1A; Supplementary fig. S1). To assemble high-quality mitochondrial genomes (mtDNAs) we used both Illumina short-read and PacBio long read sequencing which were performed on whole-cell extracts due to a lack of reproducible mitochondrial purification methods across these species. Short reads were assembled with the heteroplasmic-sensitive organelle assembler NOVOPlasty, and no heteroplasmy was detected in the assembled mtDNAs. These assemblies were merged with assembled PacBio low G+C contigs. All mtDNAs were circular-mapping single molecules with the exception of the *P. chesapeaki* mtDNAs that comprised two separate molecules (fig. 1B). The sizes of total mitochondrial genomes ranged from 40,900 bp (*P. marinus*) to 95,104 bp (*P. atlanticus*). All *Perkinsus* mtDNAs were strongly AT-skewed ranging from 82.2% (*P. marinus*) to 86.3% AT content (*P. chesapeaki*). The low degree of polymorphism and high degree of similarity between the PacBio- and Illumina-derived mtDNAs and the fact that no heteroplasmic variants were called by NOVOPlasty suggests that all mitochondrial genome exists as near identical copies. However, inter-species structural comparisons of these mtDNAs indicated little synteny between the four taxa (fig. 1B; Supplementary fig. S2), with shared sequence elements limited to protein- and rRNA-coding sequences (see sections below). Greatest
conservation of sequence was seen between the more closely related *P. atlanticus* and *P. olseni* (Supplementary file S1); however, even here there is evidence of considerable genome rearrangement and unique sequence.

Protein Coding Content of the *Perkinsus* mtDNA Genomes

Comparison of the four *Perkinsus* spp. mtDNAs showed conservation of only three putative protein-coding genes (fig. 1B). This is consistent with the highly reduced coding capacity seen in all myzozoan mtDNAs containing only genes for cytochrome B (*cob*) and cytochrome C oxidase subunits 1 and 3 (*cox1* and *cox3*, respectively). No other putative protein-coding sequences were apparent in the *Perkinsus* mtDNAs. *Perkinsus cox1* and *cob* sequences had been previously reported, but sequence for *cox3* had not, and this was suggested to have been lost (Masuda *et al.* 2010; Zhang *et al.* 2011, 2012; Bogema *et al.* 2021). Indeed, searching our mtDNAs with diverse *cox3* sequences failed to identify a putative *Perkinsus* *cox3*. However, the predicted translation of the third common *Perkinsus* mtDNA coding sequence contains the conserved superfamily domain SSF81452 (Cytochrome c oxidase subunit III-like; E value: $8.76 \times 10^{-6}$) and IPR domain IPR013833/G3DSa:1.20.120.80 (Cytochrome c oxidase, subunit III, 4-helical bundle; E value: $1.4 \times 10^{-6}$) consistent with this ORF representing a divergent *cox3*. Protein structural modeling of the *Perkinsus* COX3 sequences (see below) further supports that these sequences represent bona-fide *cox3*.

Alignments of the four *Perkinsus* taxa for the three mitochondrial genes shows very high nucleotide identity within the conserved predicted protein coding sequence, and then dramatic loss of conservation in the immediate flanking sequence (fig. 2; Supplementary figs. S3–S5). No potential ATG start codons are seen at the 5′ regions of these sequences suggesting a likely alternative initiator codon(s). Conserved phenylalanine (F) and semi-conserved asparagine (N) codons occur at the 5′ ends of all three genes, and either of them might be the N-terminal translation initiator of these genes. *Perkinsus* taxa are predicted to use the terminator codon TGA as an alternative tryptophan codon (Masuda *et al.* 2010; Zhang *et al.* 2011), and the use of this codon is highly conserved across the four species in all three mitochondrial genes including *cox3* (fig. 3). At the *cox3* 3′ end, the high sequence identity seen for the four species ends with a conserved TAA, suggesting that this could serve as a translation termination signal (fig. 2). The *cox1* genes, however, lack a common putative stop codon, with *P. olseni* only encoding a canonical stop codon more than 100 nucleotides further downstream (fig. 2; Supplementary fig. S4). The *cob* genes similarly lack a conserved stop codon at the 3′ end of sequence conservation. The *cob* genes, however, do have an internal conserved TAA (Supplementary fig. S3) that was presumed to terminate translation by Zhang *et al.* (2011). Strong sequence identity beyond this codon shared amongst the four *Perkinsus* taxa (fig. 2; Supplementary fig. S3), and conservation of the corresponding predicted protein with other COBs (Supplementary fig. S7), strongly suggests that this TAA is skipped as a terminator, and we discuss a likely explanation for this below. Given that dinoflagellates also lack stop codons in *cob* and *cox1* coding sequences, and polyadenylate these transcripts at the point precisely corresponding to the conserved C-terminus of these proteins (Jackson *et al.* 2012), we sought to determine the 3′ extremities of *cob* and *cox1* transcripts in *Perkinsus*. We use T4 RNA ligase to circularize *P. marinus* and *P. olseni*.

**Fig. 2.** Nucleotide sequence alignments of the 5′ and 3′ ends of *cob*, *cox1* and *cox3*. Conserved residues as possible initiator codons, and genome-encoded terminator TAA codons, are indicated. cRT-PCR defined 5′ UTR and polyadenylation sequences for *P. marinus* and *P. olseni* are shown. *P. atl*, *P. atlanticus*; *P. che*, *P. chesapeakei*; *P. mar*, *P. marinus*; *P. ols*, *P. olseni*. 

| Gene | 5′ ends | 3′ ends | 5′ UTR | Conserved 3′ codons | Conserved 5′ An-codons | Conserved 5′ Phe-codons | Poly-A tails | Stop codons |
|------|----------|----------|--------|---------------------|------------------------|------------------------|-------------|------------|
| *cob* | | | | | | | | |
| *cox1* | | | | | | | | |
| *cox3* | | | | | | | | |


mRNA and then RT-PCR to amplify and sequence across the fused 5′–3′ boundary for both genes. These circular RT-PCR (cRT-PCR) data determined that both cob and cox1 are polyadenylated at the same points in both species, which is at the end of sequence conservation for both genes (fig. 2; Supplementary fig. S6). The position of polyadenylation is highly conserved with that of dinoflagellates and corresponds to two to five transcript-encoded lysine residues at the termination of COB and COX1, respectively (Supplementary figs. S7 and S8).

Conserved Frameshifts Occur in All Protein Genes But Are Strongly Depleted From cox3
Previous DNA and cDNA sequencing of P. marinus cox1 and cob genes identified frameshifts in these coding sequences (Masuda et al. 2010; Zhang et al. 2011; Erber et al. 2019). Only when the reading frames are shifted at multiple points is a conserved protein translation predicted (Supplementary figs. S7 and S8). The positions of the predicted shifts of either +1 or +2 frames showed nucleotide sequence conservation and coincided with predicted locations for glycine residues (at sites with the consensus sequences AGGY or TMGGY) and proline residues (at sites with the consensus sequence CCCCT; fig. 3A; Supplementary figs. S7 and S8). One interpretation of this is that these four- and five-nucleotide sequences serve as alternative, extended codes for glycine and prolines, although, canonical three-nucleotide codons also specify conserved glycine and proline positions elsewhere in both genes (Supplementary figs. S7 and S8). The positions and sequence of the frameshifts in cox1 and cob is highly conserved in Perkinsus (fig. 3B). Only the position of glycine G384 in P. chesapeakei has lost this frameshift, or never gained it. To exclude a role of RNA-editing in correcting the frameshifts, we aligned RNAseq reads, assembled transcripts and Sanger-sequenced cDNA clones to the

![Diagram](https://doi.org/10.1093/molbev/msac191)
mitochondrial genome sequences (Supplementary file S2; fig. 3A for example). No differences between the genomic and transcript-derived sequences were observed confirming that no RNA-editing occurs in Perkinsus mitochondrial transcripts. In addition to the frameshifts observed by Masuda et al. (2010) and Zhang et al. (2011), the sequence conservation of the 5' region of Perkinsus cob genes downstream of the in frame TAA (Supplementary fig. S3) implies a further, novel, mitochondrial frameshift. The inferred translation of the +1 frame from this TAA shows sequence conservation to the terminus of other COB proteins (Supplementary fig. S7), and this predicted final reading frame is only open for four putative codons upstream of the TAA. These data suggest that a final frameshift occurs in this four-codon window before the TAA. The consensus sequences for the glycine (AGGY or TMGGY) or proline (CCCCCT) frameshifts, however, do not occur in this window and we predict that an alternative frameshift occurs at this site (fig. 3B and E and discussed below).

To query whether the newly identified cox3 genes also contain possible frameshifts, we modeled the sequence of the most abundant frameshift (+1 frame at glycine) to search for candidate frameshift positions in the cox3 sequences. We generated a position-specific matrix model derived from all 55 'AGGY' frameshifts in cox1 and cob from the four taxa, including the flanking 20 nucleotides on either side of the frameshifts. The model identified an enriched consensus sequence: WAWTTAAWWW TAGGTWTAHKTWATAWTW (E value: 2.7e-169) that, when searched against cox3, identified one possible frameshift in this gene in all taxa (fig. 3D). This frameshift alone is sufficient to extend the open reading frame that spans the conserved sequence shared amongst the four Perkinsus spp. (Supplementary fig. S7). Thus, while cox1 and cob contain 10 and 8 frameshifts, respectively, cox3 is distinct in containing only one. An interesting implication for this is a paucity of predicted glycine and proline residues in Perkinsus COX3 proteins. Furthermore, there are no canonical three-codon residues for glycine and proline in the Perkinsus cox3 sequences despite COX3 proteins in other organisms containing several of those residues that are often highly conserved in position (Supplementary fig. S9). These data suggest that some strong selective pressure exclusively on Perkinsus cox3 and not cox1 or cob.

All tRNAs Are Imported into the Mitochondria and None Are Predicted to Decode 4/5-nucleotide Codons

Given the evidence for non-canonical translation processes in the mitochondria of Perkinsus, we examined the tRNA pools of these cells for evidence of tRNAs that might participate in decoding the frameshifts. All known myzozoan mtDNAs lack encoded tRNAs, however, the deep branching point of Perkinsozoa within this large group allowed the possibility of retention of distinct mitochondrion-encoded tRNAs in this lineage. We scanned the mtDNAs of the four Perkinsus taxa with tRNAscan-SE and found no evidence of encoded tRNAs. This suggests that Perkinsus mitochondria are also fully dependent on importing cytosolic tRNAs. To examine the total pool of cellular tRNAs we then performed LOTTE-Seq (long hairpin oligonucleotide based tRNA high-throughput sequencing) for both P. marinus and P. olseni from total cell RNA. This method enriches for the tRNA 3'-CCA ends (Hou 2010; Erber et al. 2019; Chan et al. 2021). We sequenced each of these samples to ×100 genomic coverage and, from this, we identified 50 canonical tRNAs spanning all codon types (Supplementary table S2). None of these Perkinsus tRNAs mapped to the mtDNAs suggesting that all are nucleus-encoded. The only LOTTE-Seq reads that did map to mtDNAs were AT-rich sequences that lacked predicted tRNA structure and often contained the CCA sequence in the genome but which is typically not encoded in the tRNA genes (Hou 2010; Feagin et al. 2012; Jackson et al. 2012; Flegontov et al. 2015; Chan et al. 2021).

While we cannot predict which of the 50 Perkinsus tRNAs are imported into the mitochondrion, we examined all for possible explanations for the frameshifts and alternative codons used in Perkinsus mitochondria. We identified two tryptophan-like suppressor tRNAs with a TCA anticodon in the nuclear genome of both P. marinus and P. olseni (Supplementary file S3). These tRNAs are predicted to rescue mtDNA-encoded TGA stop codons providing an alternative anticodon that codes for tryptophan. We also examined tRNAs for potential anticodons complementary to the non-canonical 4–5 bp codes. To this end, we investigated all previously identified RNAs with putative CCA ends from LOTTE-seq for any resemblance to known tRNAs by carrying out BLASTn searches against a database comprising all currently known tRNAs and ncRNAs. No candidates were found, suggesting that the frameshifts in the Perkinsus mitochondria are not translated by specialized tRNAs with novel, extended anticodons.

Unused Codons Likely Trigger Frameshift Translations

Programmed ribosome frameshifting (PRF) is used by many viruses where it is most often facilitated by two sequence elements: a heptanucleotide ribosomal slippery sequence (or slip-site) followed by a downstream RNA structure (Penn et al. 2020). The slip-site typically has the form X XXX YYZ (X is three identical residues, Y is A or U, and Z is A, U or C). In Perkinsus, however, the frameshifting sites to not conform to these slippery sequence motifs (fig. 3A and E). In viruses, RNA secondary structures—either stem-loops or pseudoknots—occur ~5–10 nucleotides downstream of the frameshift. These features induce kinetic and conformational changes to the ribosome during translation and promote the slipping to the alternative frame (Penn et al. 2020). However, we also found no evidence of these predicted secondary structures in relation to the frameshift sites in Perkinsus.

Another mechanism that can contribute to PRF is the relative abundance of tRNAs where relatively depleted
tRNAs can promote ribosome stalling and slippage. To test for evidence of such a mechanism in Perkinsus we calculated the mitochondrial codon usage frequencies for each Perkinsus species. We used the three genes’ predicted reading frames based on the protein alignments (Supplementary figs. S7–S9), only excluding the frameshift sequences from these calculations (AGGY, TMGGY, CCCCT, TAAC) (fig. 3f; Supplementary table S3). We then plotted the codon-usage frequencies onto the predicted reading frames before and after each frameshift for all genes to assess translatability through these regions. From this analysis, a remarkably conspicuous observation common to all frameshifts was made. At the point of each frameshift a codon that is otherwise unused occurs in the current translation frame, and this is often followed by further unused codons in this frame (fig. 4). In the case of the +1 frameshifts, the first codon of the next frame, and those that follow, are commonly used codons restoring translatability (e.g., COX1: G74). For the +2 frameshifts, the first codon of the next reading frame is also an unused codon, and only after sliding two nucleotides do codons used elsewhere occur again, restoring translatability (e.g., COX1: P80). This situation is seen for all frameshifted positions in all four species (Supplementary figs. S10–S13). These data imply that an absence of select amino acid-charged tRNAs in the mitochondrion results in translation arrest when their cognate codons occur. Translation can only be restored by the entry of an available tRNA that can bind to the next available cognate codon. For COB: G41 in three Perkinsus spp., the first frameshifted codon is the TAG stop codon (it is an unused serine codon in P. chesapeaki) which is also otherwise unused in the mitochondrial genomes and likely lacks the corresponding release factor, which would again result in ribosome stalling. This mechanism for frameshift translation also provides a prediction for the location of the last frameshift in COB. Translatability is maintained up to the TAA codon at this site, and single nucleotide slippage allows ongoing translation in the final frame starting with an asparagine (fig. 4: COB:N303). Although the translation termination mechanism remains unclear in Perkinsus mitochondria, we predict that a delay (or absence) of recognition of this TAA codon allows frameshifting as for the other sites.

Conservation of Frameshift Sites Corresponds to Protein Secondary Structure

The frameshifts in Perkinsus spp. mitochondrial genes are predicted to translate as 16 glycines (15 in P. chesapeaki), two prolines and one asparagine; however, these amino acids are also coded for by canonical three-nucleotide codons in all three protein-coding genes (Supplementary figs. S7–S9). This raises the question of why the non-canonical codes are used and, given the conservation of their locations in Perkinsus spp., if they might perform a position-specific function in their respective proteins. To explore this question, we initially examined the conservation of these glycine, proline and asparagine residues in near and distant orthologues. Alignments of Perkinsus COX1 and COB with orthologs shows that frameshifted and canonically coded residues both occur at sites of conserved usage and usage specific to Perkinsus (Supplementary figs. S7 and S8). For COX3 the site of the single frameshifted glycine does not correspond to a conserved glycine position in other homologues (Supplementary fig. S9). Moreover, several widely conserved positions for glycine and proline in COX3 lack these residues in Perkinsus spp. Thus, for all proteins there does not appear to be a link between protein sequence conservation and position of the frameshifts.

The location of the frameshifted residues could contribute to the three-dimensional (3D) properties of the proteins, so we modeled the 3D structure of the P. marinus proteins and compared them to known structures from the model Bos taurus. All structure predictions were of high confidence (c-score >0) and showed near identical matches with the B. taurus protein structures (root-mean-square deviations <2) (fig. 5A). The strong fit of the Perkinsus proteins further substantiates our identification of Perkinsus cox3 s and the 3’ extension of cob coding sequences beyond the TAA frameshift. The locations of the frameshifted residues showed no obvious pattern with respect to the protein structures: they occurred on both sides of the inner mitochondrial membrane that these proteins span, and both within and between predicted transmembrane helices. However, given that the frameshifts likely cause temporary interruption to translation we speculate that they might contribute to some properties of protein folding. Nascent polypeptides can commence folding into secondary and tertiary structures once they emerge from the ribosome exit tunnel which is typically the equivalent of ~28 amino acids of the nascent chain. When the region situated 25–30 residues upstream of each frameshift was plotted on the predicted protein secondary structures, this emergent region typically occurred at the N-terminal boundaries of alpha helices (fig. 5B). These observations suggest a pause in translation occurs before the emergence, folding and/or membrane insertion of helical secondary structures. An interesting exception is for COX3 where the single frameshift occurs less than 20 residues from the predicted protein start, suggesting that this translation pause occurs before the N-terminus emerges from the ribosome.

Fragmented but Incomplete rRNAs in Perkinsus mtDNA

A further feature of myzozoan mtDNAs is the absence of full-length rrna sequences. In Plasmodium, Hematodinium and Chromera the presence of heavily fragmented rRNAs has been reported (Feagin et al. 2012; Jackson et al. 2012; Flegontov et al. 2015). No complete rRNAs were found in the Perkinsus mtDNAs, so we searched for corresponding fragments to those found in other myzozoans. Using 39 rRNA fragments from Plasmodium falciparum, 17 from Hematodinium and 7 from Chromera as queries, we
identified a total of 23 rRNA fragments across the mtDNAs of the four *Perkinsus* species. This is consistent with the rRNA fragmentation having occurred before the divergence of Myzozoa. The representation of rRNA fragments across the four *Perkinsus* taxa, however, was incomplete and not always homogeneous. Only the large ribosomal subunit F fragment (LSUF; as per *P. falciparum* annotation (Russell and Beckenbach 2008; Dinman 2012; Feagin et al. 2012; Seligmann 2012; Haen et al. 2014)) was found in all four mtDNAs (Figures 1 and 6; Supplementary fig. S13). Other fragments were found in three or less taxa; e.g., LSUG, LSUC, and RNA1. Furthermore, several rRNA fragments conserved in other myzozoans were not detected in *Perkinsus* spp. (Fig. 6). In total, eight rRNA fragments were found in the mtDNA for *P. atlanticus*, seven in *P. marinus*, and only four for *P. chesapeaki* and *P. olseni*.

But for some taxa only a nuclear version of the rRNA fragment was found (e.g., large subunit rRNA fragment G [LSUG] for *P. olseni*) (Fig. 6C). Moreover, some fragments were only found in nuclear assemblies and not in any of the mtDNAs (e.g. small subunit rRNA fragment B [SSUB]). These data present the intriguing hypothesis that mitochondrial rRNA gene relocation to the nucleus is occurring in *Perkinsus* spp. with their transcripts reimported into the organelle.

**Discussion**

The myzozoan mtDNAs have unprecedented levels of genome reduction, rearrangement, and reconfiguration. Furthermore, they have adopted multiple post-transcriptional processes to restore functional molecules from degenerate genes encoding either proteins or rRNAs. The Perkinsozoa (including the genus *Perkinsus*) were the one major lineage amongst Myzozoa for which the mtDNAs had been largely uncharacterized. Our mtDNA sequences of four *Perkinsus* species have resolved both common ancestral and intermediate derived features of myzozoan mtDNA, but they also reveal yet another level of elaboration of the management of mitochondrial encoded genes.

All *Perkinsus* spp. mtDNAs are relatively simple molecules encoding single copies of genes for the three common myzozoan mitochondrial proteins (COB, COX1, COX3) and fragmented rRNAs, and they lack encoded tRNAs. Thus, the coding capacity is equivalent to other myzozoans and this was likely the ancestral state. The high level of mtDNA amplification, fragmentation, and recombination seen in dinoflagellates, chromerids, and select apicomedians such as *Toxoplasma* suggests that these latter traits are derived and have occurred multiple times.
Mitochondrial Genomes in Perkinsus

independently (Jackson et al. 2007; Nash et al. 2008; Waller and Jackson 2009; Feagin et al. 2012; Jackson et al. 2012; Flegontov et al. 2015; Berná et al. 2021; Mathur et al. 2021; Namasivayam et al. 2021). Nevertheless, genome re-arrangement is seen also in Perkinsus spp. with little common synteny or shared intergenic sequence between the four species, and the P. chesapeaki mtDNA has separated into two separate molecules. This evidence of genome evolution in Perkinsus is consistent with a general myzozoan predisposition for development of mtDNA complexity through rampant recombination. The apparent use of alternative initiator codons to AUG is also shared throughout Myzozoa (and ciliates also). However, we show that both dinoflagellates and Perkinszoa (collectively Dinozoa) polyadenylate cob and cox1 without a termination codon, and that a terminator is exclusive to cox3 in this group. These unusual traits of translation termination are specific to dinozoans, whereas the use of

Fig. 5. Frameshift position conservation and location with respect to protein structures. (A) Frameshifted glycine, asparagine and proline residues (depicted as mesh volumes) in the predicted structures of COB, COX1, and COX3 from P. marinus (pink) overlaid on Bos taurus structures (blue, Protein Data Bank structures: 1OCZ:A, 1OCZ:C and 2A06:C). The non-canonically coded residues are indicated with colored boxes corresponding to their overall conservation (see key and Supplementary figs. S7–S9). Modeling and alignment scores for each protein are shown. (B) Predicted secondary structures of COX1, COB, and COX3 showing frameshifts (circles) and corresponding predicted protein emergence sites from the ribosome exit tunnel when each frameshift is translated (red brackets). RMSD, root-mean-square deviation.
mitochondrial RNA-editing, and trans-splicing of cox3 mRNAs, are evidently further derived characters specific to dinoflagellates. Perkinsus has, therefore, provided further resolution to the sequence of changes that result in the extant diversity of myzozoan mtDNAs. Perkinsus, however, has also independently gained its own mitochondrial oddities that are otherwise absent from any known myzozoan mtDNAs. The canonical TGA stop codon has been recoded to specify tryptophan, as has occurred independently in ciliates also, and we find candidate suppressor tRNAs that likely enable this. More radical is the adoption of programmed ribosomal frameshifting (PRF) in all genes, requiring 18 to 19 corrections across the encoded proteome. Our sequencing of four divergent Perkinsus taxa shows that the number and position of these frameshifts is very stable. The deepest branching taxon from the others, P. chesapeaki, is the only one to show a lost, or possible lack of acquisition, of one frameshift (for COX1 glycine:384). This difference represents only a single nucleotide deletion, and the lack of other such 'simple corrections' in any other taxa or positions implies strong selection for the maintenance of these frameshift features.

PRF plays an important function in many viral genomes where genome compaction can be achieved by overlapping coding sequences occurring in alternative reading frames (Korniﬁ et al. 2019; Penn et al. 2020). Minus one (−1) frameshifting is most common, but −2, +1 and +2 frameshifts also occur. The frameshift signals typically have two elements, a slippery sequence motif followed by an mRNA structure of stem-loop or pseudoknot. Control of the ratios of the alternative protein products fulfills the virus’s protein stoichiometric needs. In Perkinsus, however, we find no evidence of either of the signals of this PRF mechanism at the sites of frameshifting. Nor are there predicted meaningful products of translation in the alternative frames of the up to 10 sites per gene. In eukaryotic mitochondria there are also rare instances of PRF where the relative abundance of tRNAs, particularly depletion of specific tRNAs, is implicated in ribosome pausing that provides the opportunity for translation shippage into alternative frames (Rosengarten et al. 2008; Russell and Beckenbach 2008; Dinman 2012; Seligmann 2012; Haen et al. 2014). This ‘hungry ribosome’ model is implicated in a cox3 frameshift in the hexactinellid sponge.
Aphrocallistés vastus where a rare tryptophan codon (UGG) is followed in the +1 frame by a common glycine codon (GGA) (Farabaugh 1996; Beckenbach et al. 2005; Rosengarten et al. 2008). Similarly, in mtDNAs Polyrhachis ant species a frameshift in one or two positions in cob occurs where a rare codon at the ribosome A-site is combined with a weak wobble pairing of the tRNA bound in the P-site and exact Watson-Crick codon-anticodon pairing in the +1 position that promotes translation in the shifted frame. This mechanism is also consistent with known features of programmed translational frameshifting in the yeast TY1 and TY3 retrotransposons and, furthermore, in bacteria ribosomes are known to slide along a given mRNA and resuming translation several base pairs downstream of a rare codon in certain sequence contexts and/or in physiological conditions of tRNA limitation (Farabaugh 1996; Gallant and Lindsley 1998; Beckenbach et al. 2005).

A strong model for Perkinsus mitochondrial PRF driven by tRNA availability is illuminated by the four taxa’s genomes and 75 instances of frameshifts. All frameshifts occur at otherwise unused codons for these mitochondria, and in the case of the +2 frameshifts a further unused codon in the first +1 frame position. Even when there is nucleotide variability between Perkinsus spp. at the +2 glycine frameshift (TMGGY), both alternatives of the last codon of the current frame (TAG:Stop, or TGG:serine) and the +1 first codon (GGG:cysteine, or GGT:cysteine) are all unused codons reinforcing the consistency of this pattern (see fig. 3E). We currently do not know which cytosolic tRNAs are imported into Perkinsus mitochondria, however, non-use of the frameshift codons in any other regions of the genes strongly implies that charged tRNAs are absent for these select codons in the mitochondrion. This is likely to also be true for cysteine tRNAs as no in-frame cysteine codons occur in any of the genes indicating the surprising reduction to 19 amino acids of this genetic system. This is despite cysteine being present in all three proteins in non-Perkinsus taxa, including occurring at relatively conserved positions (Supplementary figs. S7–S9). An implication for PRF relying on absent tRNAs, particularly where up to 10 frameshifts per gene will demand high fidelity decoding, is that it becomes imperative that these frameshift tRNAs are excluded from the mitochondrion. It might be that mechanism for tight import-regulation of permitted tRNAs could also explain the loss of cysteine from this genome where a propensity for tRNA exclusion could have also led to a permissible loss. mRNA secondary structures might also contribute to Perkinsus PRF as our modeling of the most abundant glycine +1 frameshifts (AGGY), used to identify the cox3 frameshift, identified a strong 11–15 nucleotide consensus sequence flanking this frameshift type. While these sequences do not present obvious stem-loop or pseudoknot secondary structures as for viral and some bacterial PRF, it might be that they recruit specific proteins or short binding nucleic acids that assist in translation stalling and promotion of the change of ribosomal reading frame, as for the viral systems (Korniy et al. 2019; Penn et al. 2020).

The properties of Perkinsus mitochondrial ribosomes, while poorly understood, might have contributed to the evolution of abundant frameshifts in their genes. Myzozoan ribosomes are known to be unusual with highly fragmented tRNAs in 20 or more pieces. It is not known how these ribosomes assemble. Conserved rRNA domains of other prokaryote-derived ribosomes are apparently missing in most, and inconsistently present in many, myzozoan groups (Feagin et al. 2012; Jackson et al. 2012; Flegontov et al. 2015). The four Perkinsus mtDNAs even show significant variability between species and, tantalizingly, we observe potential for some fragments to have relocated to the nucleus from where their transcripts might be reinserted as for tRNAs. These modifications could have resulted in ribosomes of altered structural and/or mechanical properties. This could include more permissive access at the ribosome A-site to the mRNA and exploration of alternative frame codons by tRNAs, or a greater ribosome susceptibility to mechanical stresses that might be asserted by mRNA structures and/or binding molecules should these act as frameshift cues. In any case, the density of frameshifts in Perkinsus cox1 and cob, much greater than either cox3 or most other eukaryotic instances of frameshifts, demands that correction of frameshifts is highly efficient as even low frequency errors would be compounded over eight or 10 frameshift sites. Perkinsus, therefore, presents a conundrum of a seemingly degenerate translational system that must be highly efficient, and one that has not developed in other myzozoan mitochondria.

The absence of cognate tRNAs at frameshifts most likely cause a significant pause in translation. The frameshift positional conservation across Perkinsus taxa, and their alignment with the positions of ribosome emergence of the helical domains of the nascent COX1 and COB proteins, suggests that this feature has been adopted to regulate protein folding and/or insertion of the proteins into the inner mitochondrial membrane. All of the frameshifts could be lost by simple one- or two-nucleotide deletions; therefore, their maintenance suggests that they represent a gain of function in this system, rather than being a product of constructive neutral evolution (Lukeš et al. 2011). An outstanding question is why cox3 differs so markedly from cox1 and cob: (1) it contains only one frameshift, (2) this frameshift is sufficiently close to the 5′ end of the mRNA that the nascent polypeptide is likely non-emergent from the ribosome during frameshift translation, and (3) it is otherwise entirely devoid of glycines and prolines despite most COX3 proteins containing 10–15 of these residues. In dinoflagellates, cox3 is also distinguished from cox1 and cob in that it is transcribed as two mRNAs that require trans-splicing to form a mature message, and it acquires a UAA stop to its messages during polyadenylation whereas cox1 and cob are predicted to translate through the short poly-A tail adding a poly-lysine C-terminus. In Perkinsus, the maintenance of a conserved encoded TAA stop in cox3 similarly further distinguishes it from cox1 and cob. These curious features suggest that some aspects
of the synthesis, insertion, and the C-terminus of COX3 are recalcitrant to the divergent genetic properties prevalent in dinozoa, although currently we do not understand the basis of this. However, the peculiar lack of glycines and prolines in Perkinus COX3 suggests that the frameshift-dependent translation of these residues also impacts the use of canonical glycine and proline codons. The mechanism for this is currently unclear because glycine and proline are coded elsewhere in cox1 and cob using codons that are not used at the frameshift sites. Moreover, the asparagine codon AAT, which is encoded at the last frameshift of cob, is otherwise only present in one frameshift but is an abundantly used codon in cox3 and all genes. The cox3 gene, thus, tantalizes us with yet unilluminated aspects of the mechanisms and evolutionary significance of the divergent genetic properties of the mitochondria of Perkinus, and indeed of all myzozoans.

Materials and Methods

Total DNA Extraction, Sequencing, and Assembly of mtDNAs

High molecular weight genomic DNA of P. marinus, P. olseni, and P. chesapeaki was extracted using the Qiagen genomic tip kit for high molecular weight DNA, following the manufacturer's instructions and the nucleic acid integrity control was performed using a Fragment Analyzer™ (Advanced Analytical Technologies). The extracted genomic DNA was quantified with a Qubit® 2.0 Fluorometer and was used for the preparation of SMRTbell Libraries which were sequenced using a PacBio RSII sequencer and a PacBio Sequel sequencing platform (Pacific Biosciences). Following assembly with canu v1.9 (Koren et al. 2017), we noticed a small number of redundant, low G+C contigs, which we identified to exclusively represented full-length and partial mtDNAs. In addition, we sequenced all four taxa using standard Illumina sequencing. Here we extracted genomic DNA for the four Perkinus taxa using a DNeasy Blood and Tissue Kit (Qiagen) according to the manufacturer's instructions. The DNA was quantitated using the Qubit® 2.0 Fluorometer and then sheared on a Covaris E220 (Covaris) to ~500 bp. The DNA libraries were made using the TruSeq Nano DNA Library Prep kit (Illumina), according to the manufacturers’ instructions. The amplified libraries were stored in −20°C. The pooled libraries of P. marinus were sequenced in an Illumina MiSeq Instrument (2 × 300 PE reads), the libraries of P. olseni, P. atlanticus and P. chesapeaki were sequenced in a HiSeq2500 instrument (2 × 251 PE reads) and in a HiSeq4000 instrument (2 × 150 bp PE reads) (Illumina), at KAUST Core Lab facility. A PhiX control library was applied to the sequencing runs as a base balanced sequence for the calibration of the instrument so that each base type is captured during the entire run. Using Illumina reads we then used the organelle assembler NOVOPlasty (Dierckxsens et al. 2017) with a seed-and-extend algorithm using a previously reported coding sequences of cox1 and cob as seed sequences from Zhang et al. (2011) (Zhang et al. 2011) to assemble Perkinus mtDNAs. Each assembly comprised the following number of Illumina read pairs: 2125193 for P. atlanticus, 6238584 for P. chesapeaki, 347194 for P. marinus, and 155873 for P. olseni. This yielded four full-length Perkinus mtDNAs. A total of 3307, 7964, 2332 and 183 PacBio reads were assembled into the mitochondrial genomes of P. atlanticus, P. chesapeaki, P. marinus and P. olseni, respectively. The PacBio reads were aligned to the mitochondrial genomes using minimap2 v2.24 with the option -x map-pb and only the alignments with a phred mapping quality score greater than 60 were retained. The Illumina reads were aligned to the mitochondrial genomes using bwa mem v0.7.17 with the default options, no quality filters were applied after mapping, and discordantly mapping reads were discarded using samtools view v.1.15–7 with the option -f 3.

Circular Reverse Transcriptase PCR (cRT-PCR)

Total RNA for P. marinus and P. olseni was extracted using Trizol (Thermo Fisher) following the manufacturer’s instructions. Approximately, 11 µg of RNA were first treated with 0.08 U/µl of TURBO DNase and 1× TURBO DNase Buffer (Thermo Fisher) to remove any genomic DNA trace. Then, ~1.2 µg of DNase-treated total RNA was used for transcript's head to tail ligation with 0.25 U/µl of T4 RNA ligase and T4 RNA ligase Buffer 1×, 1 U/µl of RNasin and 20% PEG8000 (all these reagents are from Promega). First strand cDNA synthesis across the ligated transcripts was performed for cob and cox1 (55°C) using 10 U/µl of SuperScript III reverse transcriptase and 5 mM DTT (Invitrogen), 2 U/µl RNasin, 0.8 mM dNTP (Thermo Fisher) and 0.1 µM of specific primers (Supplementary table S1A and S1B). cDNA was treated with 0.1 U/µl of RNase H (Thermo Scientific) prior to PCR. PCR was performed with 2 µl of cDNA template, 0.2 mM dNTPs (NEB), 0.02 U/µl of Phusion DNA polymerase with 1× PCR reaction buffer Mg+ (Thermo Fisher) and 0.2 µM of specific primers (Supplementary table S1A and S1B). All reactions had negative (circularized transcripts but no reverse transcriptase) and blank controls (no circularized transcripts and no reverse transcriptase). PCRs were first cleaned with NucleoSpin clean-up kit (Macherey-Nagel) and then, ligated into the pGEM-T Easy Vector System I (Promega), cloned into XL1-Blue Competent Cells (Agilent) and sequenced. Sequence alignments and visualization were carried out with Sequencer 5.2.3* (Gene Codes)

RNA Extraction and Sequencing

Total RNA for the four Perkinus taxa was extracted using Trizol reagent (Invitrogen) following manufacturers’ instructions and quantitated using the Qubit® 2.0 Fluorometer and was used for library preparation. The strand specific RNA libraries were made using the TruSeq Stranded mRNA Sample Prep Kit according to the manufacturer’s instructions. The amplified libraries were sequenced in an Illumina HiSeq4000 instrument (2 × 150 bp PE reads) at KAUST Core Lab facility. A PhiX...
control library was applied to the sequencing run as a base balanced sequence for the calibration of the instrument so that each base type is captured during the entire run. To capture non-polyadenylated and short RNAs, we also use the non-coding (nc) RNAseq protocol as outlined in Minshall et al. (2022). Briefly, total RNA was extracted from *P. marinus* and *P. olseni* as described above and we applied the TruSeq Stranded Total RNA protocol (Part # 15031048 Rev. E October 2013) with the following changes: (1) to avoid selective loss of shorter RNA from post-rRNA depletion supernatant: in addition to 99 µL AMPure beads, 250 µL isopropanol was added to the ~40 µL supernatant. Following mixing and the recommended 15 min incubation, magnetic bead capture was extended to 15 min; (2) to minimize critical damage to short RNAs while allowing necessary fragmentation of longer transcripts: RNA was fragmented @ 94°C for 3 min instead of the default 8 min; (3) to avoid selective loss of short cDNAs from the 2nd strand synthesis reaction: in addition to 90 µL AMPure beads, 250 µL isopropanol was added to the 50 µL reaction. Following mixing and the recommended 15 min incubation, magnetic bead capture was extended to 15 min; (4) to recover ligated cDNA including shorter inserts: ×1.25 volumes of AMPure beads were used instead of the recommended ×1 volume; and (5) to recover PCR-amplified library spanning shorter inserts: ×1.2 volumes of AMPure beads were used instead of the recommended ×1 volume. Resulting raw reads were then mapped to mitochondrial transcripts for *cox1*, *cox3*, and *cob* as described below.

**PCR Amplification and Sanger-sequencing of *cox1*, *cox3* and *cob***

In addition to the short-read sequencing, both DNA and cDNA versions of *cox1*, *cox3* and *cob* were amplified from *P. marinus* and *P. olseni* and Sanger sequenced. Extracted DNA was processed using GenElute™ Mammalian Genomic DNA Miniprep Kit (Sigma), including the optional RNase A digest. For RNA, pellets were rapidly thawed and lysed in >9 volumes of TRI-Reagent (Sigma) and processed using Direct-zol (Zymo), including an on-column DNase digest. To ensure complete removal of residual DNA, 5 µg of the purified RNA was further treated in solution by 5 units of RQ1 DNase in 100 µL of ×1 RQ1 buffer (Promega) for 1 h at 37°C, re-extracted by phenol/chloroform and ethanol-precipitated. For each species, 300 ng of total RNA was reverse-transcribed in 20 µL using Transcriptor (Roche) primed with 150 pmol random hexamers following the manufacturer’s protocol. Parallel reactions were set up without reverse transcriptase (RT-) to ensure no genomic DNA contribution to RNA-derived amplicons. 2.5 µL reverse transcription reactions or ~10 ng genomic DNA were then subjected to PCR amplification by the high-fidelity Phusion polymerase (NEB) in 50 µL reactions using the primer pairs listed in Supplementary table S1C and the following cycling program: 30 s denaturation @ 95°C, 30 cycles of (30 s denaturation @ 95°C, 30 s annealing @ 50°C, 45 s extension @ 72°C), final blunting for 1’ @ 72°C. Primers were designed across stretches of sequence identical between *P. marinus* and *P. olseni* genes, with a predicted melting temperature of ~51°C (using default parameters in Primer3 http://bioinfo.ut.ee/primer3-0.4.0/primer3/). No PCR product was observed in RT-samples or in no-template controls for either the reverse transcription (RT0) or the PCR step (NTC). Purified PCR products were Sanger-sequenced using Cox1-F, Cox3-R or Cob-F for the corresponding amplicons (Supplementary table S1C).

**RNA Read Mapping**

To determine whether observed framemaps may in fact be sequencing errors and not naturally encoded, we mapped publicly available cDNAs (*cox1*: HQ670240 and cob: HQ670241), sanger-confirmed cDNAs (this study), in-silico transcripts and RNA-seq raw reads (SRA accession numbers: SRR1154652, SRR1154653, SRR1154655, SRR1300219, SRR1300220, SRR2094556, SRR2094558, SRR8390008 and SRR11648389) to the mtDNA references with bwa mem v0.7.17, marking split alignments as secondary alignments. BAM files of this mapping are available publicly via this link: https://fsgshare.com/projects/Mitochondrial_genomes_in_Perkinsus_decode_conserved_framemaps_in_all_genes/139894.

**LOTTE-seq for tRNAs**

Total RNA was extracted from *P. marinus* and *P. olseni* and prepared for LOTTE-Seq as previously described (Erber et al. 2019). Upon sequencing, LOTTE-Seq reads with genome coverage greater than 100 × were filtered, clustered, and subjected to the tRNA gene predictor tRNAscan-SE 2.0 (Erber et al. 2019; Chan et al. 2021) to either identify typical tRNAs or atypical tRNAs such as trans-spliced and circularized permuted tRNAs; and to classify them accordingly using the default parameter settings as previously described (Edgar 2004; Erber et al. 2019). Sequences not identified as tRNAs with tRNAscan-SE 2.0 were inspected individually for similarities with tRNAs. Remaining un-assigned reads were analysed with BLASTn (https://blast.ncbi.nlm.nih.gov/Blast.cgi) for signs of homology with tRNAs or other known ncRNAs. In addition, their secondary structures, computed using RNAfold 2.0 (Lorenz et al. 2011), were investigated for similarities to tRNAs.

**Phylogenetic Analysis**

Using a complete *P. marinus* 18S small subunit (SSU) rRNA as search bait (AF126013.1) and BLASTn, an alignment was generated of 146 perkinsid and closely related Alveolata flagellates comprising at least 1221 aligned bases for each taxon using the aligner MUSCLE (Edgar 2004; Capella-Gutierrez et al. 2009) at standard settings within Geneious 8.1.9 (Biomatters). The alignment was automatically trimmed using trimAI (Capella-Gutierrez et al. 2009; Nguyen et al. 2015; Minh et al. 2020) and a maximum likelihood tree was calculated using IQ-TREE at standard
settings (Rice et al. 2000; Nguyen et al. 2015; Minh et al. 2020). Branch support was calculated using the Ultrafast Bootstrap (UFBoot) algorithm with 1,000 replicates and the Shimodaira–Hasegawa approximate likelihood ratio test (SH-aLRT) allowing ultrafast bootstrap support values to first converged. The full and trimmed alignments, raw consensus tree file and accession numbers are provided in PHYLIP format (Supplementary file S1).

Further Bioinformatic Analyses

Genomic G+C content was determined using the geece program from EMBOSS v6.6.0 (Rice et al. 2000; Bernt et al. 2013; Donath et al. 2019). Automated mtDNA annotation was carried out using MITOS2 (Zhang et al. 2000; Bernt et al. 2013; Donath et al. 2019). For confirmation the coding sequences of cox1 and cob were detected using BLASTn v2.11.0 (Zhang et al. 2000; Blum et al. 2021) using the reported sequences of cox1 and cob. To test for cox3, we performed BLASTx searches using the whole mtDNA genome sequences against 74,641 cox3 proteins. Since BLASTx searches did not reveal the coding sequence of cox3, we extracted the ORFs longer than 300 bp using the getorf program from EMBOSS. We then subjected the deduced amino acid sequences to interproscan searches (Harris 2007; Blum et al. 2021) and selected the ORFs that presented the domains IPR000298 (Cytochrome c oxidase subunit III-like) and IPR035973 (Cytochrome c oxidase subunit III-like superfamily). Synteny analysis was performed by searching for highly similar regions among the four mtDNAs with BLASTn (task: megablast, word-size = 4, E value less or equal than 1e-3). Only reciprocal matches were kept for visualization. Whole genome alignments of the mtDNAs were carried out within Geneious R10 (Biomatters) using a LASTZ v1.02.00 (Harris 2007; Krzywinski et al. 2009) plugin applying a step length of 30, a seed pattern of ‘12 of 19’, searching both strands, allowing a single transition in a seed hit and applying a HSP threshold of 3,000. Mitochondrial genome visualization was performed using circos v0.69 (Krzywinski et al. 2009; Ma et al. 2014).

To model the most abundant frameshift sequence occurring in cox1 and cob, we aligned the sequences surrounding such frameshifts and constructed a position specific matrix (PSM) with meme v5.3.0 (Grant et al. 2011; Ma et al. 2014) using the option ‘one occurrence per sequence’. The PSM was employed to search for potential frameshifts in cox3 using fimo v5.3.0 (Gouy et al. 2010; Grant et al. 2011) with a q-value threshold of 0.0003. An updated PSM was constructed from the 59 single-nucleotide frameshifts detected in cox1, cob and cox3 (consensus: WWWYTAGGTWTAWHAKTWT, E value: 4 × 10^{-140}).

COX1, COX3 and COB alignments were constructed using clustal omega (Sievers et al. 2011), manually edited using seaview v5.0.4 (Gouy et al. 2010; Okonechnikov et al. 2012) and visualized using ugene (v34) (Okonechnikov et al. 2012; Yang and Zhang 2015). The 3D protein structures were modeled using the I-TASSER (Wheeler and Eddy 2013; Yang and Zhang 2015) server and the resulting models were visualized using the PyMOL Molecular Graphics System (Version 2.0.0 Schrödinger, LLC) and pdbsum (Laskowski et al. 2018). Codon usage was calculated for each species using the cusp program from EMBOSS (Rice et al. 2000) and the predicted reading frames of the full CDS but excluding the 4- or 5-nucleotide frameshift sites. Codon usage frequencies were then plotted for the alternative frames surrounding each predicted frameshift position (−10 to +5 codons).

Sequences corresponding to possible rRNA fragments were searched in the mtDNA genomes using nhhmer (Masuda et al. 2010; Zhang et al. 2011; Jackson et al. 2012; Wheeler and Eddy 2013; Bogema et al. 2021) with 39 rRNA fragments from Plasmodium falciparum, 17 fragments from Hematodinium and 7 fragments from Chromera velia as queries. The gap-opening and gap-extension probabilities used for constructing the hidden Markov models of nhhmer were adjusted to 0.001 and 0.2, respectively. All heuristic searching parameters were turned off to increase sensitivity. Twenty-three putative rRNA fragments were identified in the mitochondrial genomes. The predicted rRNA fragments and the known rRNA fragments were used as nhhmer queries to search for additional rRNA fragments in the nuclear genome sequences of the four species (manuscript in preparation). To avoid detection of cytosolic rRNAs, the regions corresponding to known rRNA genes and transposable elements up to 1000 nt long with metazoan-only Dfam rRNA domains were excluded from the searches.

Supplementary Material

Supplementary data are available at Molecular Biology and Evolution online.

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Data Availability

Datasets generated for this study are entirely accessible via public databases. The annotated sequences of Perkinsus mtDNA genomes were submitted to GenBank under accession numbers ON035647, ON035649, ON035664, ON035645 and ON035648 (P. atlanticus, P. chesapeakei...
molecules 1 and 2, *P. marinus* and *P. olseni*, respectively). Raw LOTTE-Seq sequences were deposited at the European Nucleotide Archive (ENA) with the following accession numbers: ERR6155172, ERR6155171, ERR6155170 (*P. marinus*), ERR6155169, ERR6153607 and ERR6153604 (*P. olseni*). Raw Illumina and Pacbio sequencing reads were submitted to NCBI SRA under BioProject with accession number PRJNA811585.

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