Intraspecies variation of the mitochondrial genome: An evaluation for phylogenetic approaches based on the conventional choices of genes and segments on mitogenome

Abstract

Intraspecies nucleotide sequence variation is a key to understanding the evolutionary history of a species, such as the geographic distribution and population structure. To date, numerous phylogenetic and population genetics studies have been conducted based on the sequences of a gene or an intergenic region on the mitochondrial genome (mtDNA), such as cytochrome c oxidase subunits or the D-loop. To evaluate the credibility of the usage of such ‘classic’ markers, we compared the phylogenetic inferences based on the analyses of the partial and entire mtDNA sequences. Importantly, the phylogenetic reconstruction based on the short marker sequences did not necessarily reproduce the tree topologies based on the analyses of the entire mtDNA. In addition, analyses on the datasets of various organisms revealed that the analyses based on the classic markers yielded phylogenetic trees with poor confidence in all tested cases compared to the results based on full-length mtDNA. These results demonstrated that phylogenetic analyses based on complete mtDNA sequences yield more insightful results compared to those based on mitochondrial genes and segments. To ameliorate the shortcomings of the classic markers, we identified a segment of mtDNA that may be used as an ‘approximate marker’ to closely reproduce the phylogenetic inference obtained from the entire mtDNA. In the case of mammalian species, which can be utilized to design amplicon-seq-based studies. Our study demonstrates the importance of the choice of mitochondrial markers for phylogenetic analyses and proposes a novel approach to choosing appropriate markers for mammalian mtDNA that reproduces the phylogenetic inferences obtained from full-length mtDNA.
Introduction

Mitochondria, the organelles present in most contemporary eukaryotic organisms, are considered cellular power plants due to their essential role in energy production. It has been well established that modern mitochondria originated from a bacterial endosymbiont, which probably belongs to an α-proteobacteria ancestor, in a proto-eukaryotic host [1].

As a remnant of being a symbiotic organism, mitochondria possess a genome (mtDNA) that codes for the part of the proteins constituting the organelle. Owing to its evolutionary history as an endosymbiont transformed into innate cellular machinery, the ancestral genome has been substantially altered and reduced into contemporary mtDNA, and many of the original genes have been transferred to the nucleus or lost [2,3]. Consequently, mtDNA genome diversity, with a variety of structures, sizes, and gene contents, emerged across the Eukaryota over the course of evolution. For example, in Metazoa, mtDNAs are circular and relatively small compared to other taxa [4–6]. Generally, metazoan mitochondrial genes lack introns, while a single large noncoding segment, the control region or the D-loop, is found in bilaterian animals [7,8]. The mtDNA typically contains 13 protein-coding genes that code for the electron transport chain and oxidative phosphorylation, 22 tRNA genes, and the genes that code for the large and small rRNA subunits. Gene content, organization, size, translation codes, and uniparental inheritance are preserved among most of the metazoan [4–6], with some exceptions in non-bilaterianism [9]. While animal mtDNA are as small as 15 to 17 kb, plant mtDNAs are much larger [10,11]: angiosperm mtDNAs are typically in the range of 200 to 700 kb, with an extreme example, as large as 11 Mb, found in Silene conica [12]. Fungi mtDNA have been less explored compared to their animal and plant counterparts [13]. Most of the fungi mtDNA studied to date are circular, but there are a few species possessing linear forms [14]. Fungi mtDNAs resemble those of plants in certain features, for example, the presence of a variable number of groups I and II large introns and intergenic regions [15,16], while the kingdom is often associated with Metazoa. The presence of mobile endonuclease open reading frames (ORFs) in these introns represents one of the major sources of variability in fungal mtDNA [17]. In the case of ‘catch-all kingdom’ protists, their mtDNA is known to be diverse [18]. For example, the simplest form of mtDNA is found in myxozoans, alveolates, which encode only subunits 1 and 3 of cytochrome c oxidase (cox), one subunit of cytochrome c reductase (cob), and short, functional fragments of small and large subunits of mitoribosomal RNA [19–21]. In contrast, the most gene-rich mtDNA identified to date belongs to jakobids [22]. The structures of protist mtDNAs are also diverse: while many protists possess a circular or a permuted linear, some dinoflagellate mtDNAs consist of small linear fragments that contain many pseudogenes and many non-functional gene fragments [23–26].

Existing in virtually all eukaryotic organisms and, due to their small sizes in comparison to chromosomes, their conserved genetic components and fast evolutionary rate [5,6,27], mtDNA has been regarded as an excellent marker for phylogeny, phylogenetics, and population structure studies for decades. For example, the non-coding region D-loop in animals is adopted as a marker due to its sequence variation. For animals, fungi, plants, and protists, some well-known core mitochondrial genes, such as cox1, cob, or nad6, have been frequently used to study the speciation processes as well as intraspecific genetic variations. This is because these genes are longer than other mitogenes, and longer sequences presumably better represent the phylogeny of the entire mtDNA. However, several studies have demonstrated that the phylogenetic inferences obtained from the entire mtDNA and the partial sequences, both D-loop and protein-coding sequences, may be discordant in some cases [28–30]. In addition, we identified two hypervariable genes on the mtDNA of a protist, Heterosigma akashiwo, that code for two hypothetical proteins that are homologous to each other [31–33]. Interestingly,
their sequences show association with the geographic origins of the isolates, while the rest of mtDNA did not show clear isolation-by-distance, demonstrating that the phylogenetic studies based on different segments of *H. akashiwo* mtDNA may yield completely different insights. These observations pose a simple but important question: is the traditional choice of gene markers from mtDNA, which have been widely adopted for organisms belonging to different kingdoms, suitable for phylogeographic or population genetic studies?

To answer this question, we surveyed mtDNA sequences of different organisms, with a particular focus on their intraspecific variations and distribution of heterogeneities over different regions. We found that the extent of the sequence variations differed widely depending on both the organism and the regions of mtDNA. The limitations of the usage of traditional phylogenetic markers are discussed in detail. In addition, we attempted to identify an mtDNA segment that can be adopted as an ‘approximate marker’ for mtDNA-based phylogenetic study. We adopted two multiple sequence alignment (MSA) strategies for the phylogenetic reconstructions to test how the choice of the step affects the final output of the entire analyses.

**Results and discussion**

**Literature survey for the usage of phylogenetic mtDNA-based markers**

We first surveyed PubMed to understand the trends in the usage of mtDNA-based markers for phylogenetic and population structure studies (Fig 1). Based on the search using different Boolean strings, the number of hits for a search for phylogeographic and population structure studies for non-human organisms using mitochondria-derived markers was 4,310 for the entire period and 541 from July 2018 to July 2021. The number of studies that contained the terms D-loop, cox, or cob were particularly plentiful, suggesting that these genes/segments have been frequently adopted as markers for this purpose. The number of studies based on full-length mtDNA published to date was ~12.9% of the total number of studies. The number of studies based on full-length mtDNA during the past three years was 17.1% of those based on all markers published during the entire period, suggesting that usage of complete mtDNA for phylogenetic or population structure studies became more common recently. At the same time, the studies based on D-loop, cox, and cob during the past three years were 12.7%, 24.5%, and 12.2% of the total number of studies conducted during the same period, respectively, suggesting that these markers were still widely used. cox was frequently used for studies on invertebrates, plants, and fungi, and D-loop was most often used for vertebrate studies (S1 Table).

**Skewed distributions of sequence heterogeneities on mtDNA in various species**

Previously, our group demonstrated that two particularly variable protein-coding regions of *H. akashiwo* showed a strong association with their geographic origins, while other parts of mtDNA did not [31–33]. This may demonstrate that, at least in certain species, analyses on different parts of the mtDNA sequences yield varied phylogenetic insights. To test if this is generally the case, we sampled mtDNA sequences from a variety of taxa, obtained phylogenetic insights based on various parts of the mtDNA sequences, and observed the consistency of the results. To this end, we selected datasets for several organisms that consist of (1) complete mtDNA sequences with no ambiguity (i.e., N, W, M, R, etc., instead of A, T, G, and C), (2) at least 25 independent strains/individuals from the National Center for Biotechnology Information GenBank database for statistical analysis, and (3) sequences published with information on their geographical origin. Twelve organisms, including *Homo sapiens* and *Saccharomyces cerevisiae*, met the criteria. Because the usage of full-length mtDNA has been the norm for *H.*
for a substantial period [34–36], we finally chose ten datasets: *Apodemus agrarius* (striped field mouse [37]), *Camelus bacterianus* (domestic camel [38]), *Canis lupus* (gray wolf [39]), *Capra hircus* (domestic goat [40]), *Fusarium culmorum* (fungal plant pathogen causing seedling rot, foot rot, and ear blight to a wide range of monocots and dicots [41]), *Fusarium graminearum* (fungal plant pathogen causing fusarium head blight on wheat, barley, and rice and ear rot on maize [41]), *Gallus* (domestic red junglefowl [42]), *H. akashiwo* (a protist, causative species of harmful algal bloom [31–33]), *Pan troglodytes* (chimpanzee [43]), and *Ursus arctos* (brown bear [28,29]). Such datasets were not available from the plant kingdom.

First, we surveyed the distribution of sequence heterogeneities over the entire mtDNA of the organisms. Vertebrate species possess a small number of genes that lack introns in a largely conserved order arranged on compact genomes [6,7]. Their sequence heterogeneities were observed to be particularly concentrated in D-loops in all analyzed species, while other parts of the mtDNA, including *cox* and *cob*, had rather uniform heterogeneity distribution at much lower levels (Fig 2A–2G). Maximum heterogeneities observed in D-loops varies among the animal species; that of *A. argarius*, *G. gallus*, and *C. hircus* are particularly small, *C. bacterianus* is intermediate, and *C. lupus* and *U. arctos* are maximal. Protist mtDNAs are known to exhibit highly variable architectures, sometimes composed of multiple chromosomes, both circular and linear [18,22,44]. *H. akashiwo* possesses ~39 kbp circular mtDNA that codes for 17
respiratory genes, 16 ribosomal proteins, 2 rRNA subunits, 1 transporter protein, and 10 conserved hypothetical proteins (Fig 2H [31–33]). Previous studies revealed that two of the hypothetical proteins code for homologous proteins [31–33]. In this study, analysis of the newly sequenced mtDNA of 12 strains combined with the previously analyzed dataset further confirmed that these two protein-coding sequences, hypervariable open reading frames 1 and 2 (MtORFvar1 and 2), expressed particularly high sequence heterogeneities (Fig 2H [31–33]). An intergenic region between large ribosomal RNA and a hypothetical protein, termed the
'intergenic variable region' in previous studies [31–33], is also particularly highly variable (Fig 2H), while the sequences of the region did not express isolation-by-distance [31–33].

Two fungal species, *F. culmorum* and *F. graminearum*, possess mtDNA with larger sizes, and all genes are coded by the same strand (Fig 2I and 2J). In many fungal species, mitogenes contain a variable number of groups I and II large introns that code for mobile endonuclease ORFs [17,41]. In these species, *cox1*, *cox2*, and *cob* in particular express intraspecific variations in exon/intron numbers; also, a couple of truncated versions of a hypothetical protein are found in the isolates. While expressing major variations in length and exon/intron configuration for these genes among strains, the sequence variations within each variant are notably smaller compared to mtDNA from other kingdoms (Fig 2I and 2J). In addition, sequence heterogeneities were particularly high in protein non-coding regions, both in intergenic regions and introns, suggesting that protein sequences are highly conserved among the different isolates (Fig 2I and 2J).

Overall, sequence heterogeneity distribution patterns were significantly different over the analyzed kingdoms, and commonly selected marker genes, *cox1*, *cob* and *nads*, are not particularly variable.

**Correlation of phylogenies between partial and entire mtDNA sequences in different species and effects of the choice of multiple sequence alignment strategies on the phylogenies**

Ideally, partial mtDNA sequences could be adopted as convenient phylogenetic markers when their phylogenies reproduce those of the entire mtDNA. To test if this is the case, we evaluated the relatedness between the phylogenies of each gene and entire mtDNA quantitatively. The extent of the correlation between the phylogenies based on the entire and different genes/segments of mtDNA varies depending on the species (Fig 3 and S1 Fig). Because several studies revealed that the choice of MSA methods and implementation affected the downstream analyses [45–48], we adopted two different MSA methods, PRANK followed by GUIDANCE2 (Fig 3) or MAFFT (S1 Fig), and compared the resulting phylogenetic reconstruction. For example, *A. agrarius* showed high correlations between trees based on the entire mtDNA and all of the tested partial mtDNA sequences in the results from both analyses. In contrast, *C. hircus*, *C. bacterianus*, and *G. gallus* showed relatively low correlations between all analyzed partial and entire mtDNA in both analyses, suggesting that none of the three adopted partial mtDNA sequences would reproduce the phylogeny of the entire mtDNA. It may be noteworthy that the topologies of the phylogenetic trees obtained from D-loop that accumulated sequence heterogeneities at the highest levels in the entire mtDNA of all of the tested animal species did not necessarily correlate well with those of the entire mtDNA sequences, demonstrating that the segment of mtDNA from animal species may not consistently reproduce the phylogeny of the entire mtDNA. Although the correlation between PRANK-based and MAFFT-based full-length mtDNA phylogenies was high (S1 Fig), the extent of the correlation of *cox1*, *cob*, or D-loop to the entire mtDNA deduced from either MAFFT and PRANK were significantly different. To date, several studies have demonstrated that the quality of the phylogenetic reconstructions based on GUIDANCE2-verified PRANK MSA prevails over those based on MAFFT-based MSA [45–47]. Our observation further underscored the importance of the choice of MSA package for downstream analyses. In cases of two fungal species, the phylogenies of all analyzed genes expressed moderate to low correlation with that of the entire mtDNA by both analyses (Fig 3 and S1 Fig). This may be because the strains of the species possess mtDNA consisting of different combinations of various genotypes of *cox1*, *cox2*, and hypothetical protein variants. Here, again, analyses adopting either gene as the representative marker may yield
significantly different results compared to those based on the entire mtDNA. In the case of *H. akashiwo*, the sequence variations of all tested mitochondrial genes correlated to that of entire mtDNA at certain extents. In general, these data revealed that the phylogenies based on the full-length mtDNA and partial sequence of mtDNA do not necessarily correlate with each other, and the gene/segment that phylogenetically correlated best to the entire mtDNA differs depending on the species.

**Comparison of phylogenetic reconstructions based on entire and partial mtDNA sequences**

To gain more detailed information regarding the similarities or discrepancies between the phylogenetic insights obtained from entire and partial mtDNA sequences, we compared the clustering pattern of the phylogenetic trees reconstructed by these analyses. We chose three organisms, *A. agrarius* (the closest correlation between all selected genes and the entire mtDNA is shown in Fig 3 [37]), *P. troglodytes* (the second-highest correlation is shown in Fig 3 [28,29]), and *H. akashiwo* (expressed correlation at intermediate levels [31–33]), and compared the results with previously published observations. For the comparison, the partial mtDNA whose phylogeny exhibited the highest correlation with that of the entire mtDNA (Fig 3) was chosen for each organism.

Phylogenies of *A. agrarius* based on the entire mtDNA and *cob* sequences showed similar branching patterns (Fig 4). Most of the individuals from Denmark’s mainland and islands were grouped together and segregated from individuals who originated in mainland Europe, while two individuals from Denmark’s mainland clustered with individuals from Germany. While smaller datasets that are composed of sequences without any ambiguity were used for the analyses, the phylogenetic insight obtained here was consistent with the one previously published to show the colonization process of the species in Denmark [37].
In the case of *P. troglodytes*, while full-length mtDNA and cob phylogenies showed similar clustering patterns, the posterior probabilities at many branches are lower in cob phylogenies, resulting in phylogenetic reconstruction of lower resolution.

Finally, we conducted a phylogenetic reconstruction of full-length *H. akashiwo* mtDNA and *MtORFvar1* sequences (Fig 6). Previous phylogeographic studies with a smaller number of sequences revealed that *MtORFvar1* and -2 showed significant isolation-by-distance. The sequences of the isolates that were obtained from latitudinal regions higher than the previously defined geographic border, 42˚N, formed a separate clade from others, with some exceptions [31–33]. In this study, when the entire mtDNA and *MtORFvar1* sequences were analyzed, most of the strains obtained from >42˚N of the United States of America (USA) Atlantic, USA
Pacific, and Europe areas were segregated from the strains obtained from other regions (Fig 6A and 6B). Notably, CCAP 934_8 and CCMP1595, isolated from Seattle, Washington, USA, and Rhode Island, USA, respectively, did not cluster with the >42˚N clade (Fig 6A and 6B) in the current study. In addition, Haek9505-1 isolated from Tampa Bay, Florida, USA, was associated with the high latitude longitude as an exception, as previously observed [31–33]. In addition, as observed for A. agrarius and U. arctos datasets, analyses on the entire mtDNA of H. akashiwo yielded trees with higher posterior probabilities compared to those constructed based on the MtORFvar1 analyzed here (Fig 6). These results demonstrate that, even when the phylogenies based on the partial mtDNA sequence and the entire mtDNA correlate well, at least for the datasets analyzed in this study, phylogenetic analyses based on high-quality, entire mtDNAs yielded results with higher resolutions and confidence compared to those based on partial sequences. In addition, the genetic distances estimated based on the entire and partial mtDNA are significantly different in P. troglodytes and H. akashiwo, while estimations based on the A. agrarius dataset were close to each other.

To design a novel mtDNA-based marker that approximates the phylogeny based on the entire mtDNA

Our observations demonstrated that the traditional markers yield fewer phylogenetic insights compared to the entire mtDNA sequences, even for the dataset whose genetic distances among the sequences closely correlate to those of the entire mtDNA dataset. This may be because the longer sequences contain more substitution sites in the dataset and are thus more phylogenetically informative than shorter sequences. Alternatively, it may be because the entire mtDNA consists of both coding and non-coding sequences. The protein-coding sequences evolve under the pressure of functional constraints and thus yield different evolutionary patterns compared to the non-coding region. A segment may serve as a practical phylogenetic marker if it contains both coding and non-coding regions and the phylogenetic inferences of those resemble the one based on entire mtDNA. To identify such a segment, several different lengths of partial mtDNA sequences were selected and the concordance between the phylogenetic trees based on those datasets and the entire mtDNA were compared. To obtain convenient marker candidates that closely reproduce the phylogenetic inferences based on mtDNA in a reproducible manner, we attempted (1) to identify the contiguous and shortest segment of mtDNA that best reproduces the phylogeny of the entire mtDNA, and (2) to test if the randomly selected, smaller datasets of the short sequences reproduce the phylogeny of the entire mtDNA.

To this end, we selected three vertebrates, A. agrarius (Fig 4), P. troglodytes (Fig 5), and U. arctos (Fig 7), that showed clear clustering in the phylogenetic trees based on the entire mtDNA datasets, in which five out of six most ancestral branching were supported by >75% of posterior probabilities (Figs 4–7 and S2–S4 Figs). The mtDNA part whose phylogenetic reconstruction is most concordant with the one based on the entire mtDNA would be a useful phylogenetic marker. The analysis based on the single gene or D-loop showed consistently smaller concordance to the entire mtDNA compared to the analysis based on mtDNA segments containing both coding and non-coding sequences (Fig 8).

Cob, cox, and D-loop showed different levels of concordances to the entire mtDNA depending on the organisms, further demonstrating the risk of adopting the traditional markers. Next, we selected several contiguous parts of mtDNA that contain cox, cob, nads, mS, and mN on their terminals, and spanning through D-loop. Analyses based on cob-rnS showed significantly higher congruence to the results based on the complete mtDNA than cob or D-loop, and the congruence was comparable to those based on the longer sequences, such as cob-cox1
Fig 5. Phylogenetic divergence based on (A) full-length mtDNA and (B) cob and (C) contiguous sequence including cob to mS of P. troglodytes. The phylogenetic trees were reconstructed and rooted with Gorilla gorilla gorilla an outgroup (NCBI accession NC 011120.1). Branches with a posterior probability of 0.5 to 1.0 are expressed in color as indicated. Each branch is color-coded by the clades in (A) as indicated.

https://doi.org/10.1371/journal.pone.0273330.g005
When the analyses were conducted using the MAFFT-aligned MSA, overall congruences of the partial mtDNA datasets were lower than the results obtained from PRANK-MSA and, particularly in *U. arctos*, *cob-rns* showed significantly lower congruence to the full-length mtDNA than *cob-cox1* (S6 Fig). These results suggest that the usage of
PRANK-GUIDANCE2-based MSA may allow simulation of the phylogenetic reconstruction obtained from the entire mtDNA by analyzing shorter partial sequences closely.

Further, the phylogenetic trees based on the entire mtDNA and cob-rnS segments showed substantially higher homology compared to the trees based on the entire mtDNA and an mtDNA gene (Figs 4A, 4C, 5A, 5C, 7A and 7C). It may be noteworthy that the genetic distance estimated from the cob-rnS datasets tended to be closer to those estimated from full-length mtDNA than the single gene or D-loop of the organisms (Figs 4, 5 and 7).

The higher correlations between trees based on full-length mtDNA and cob-rnS compared to the other mtDNA segments were also confirmed for two mammalian organisms, *C. bacterianus* and *C. lupus*, whose branching in mtDNA phylogenetic trees was supported by low posterior probabilities (S2 and S3 Figs). In the case of *C. hircus*, neither of the analyzed mtDNA
segments showed high correlation with the phylogeny based on the entire mtDNA (S4 Fig). This may be, at least partially, because the phylogeny based on the entire mtDNA shows ambiguous clustering among most of the analyzed individuals.

These results suggest that phylogenetic analyses based on the segments may provide ‘approximate markers’ for mtDNA. Because cob and rnS coding sequences are highly conserved, an amplicon-seq-based approach can be adopted to simultaneously obtain the sequences of several individuals at a feasible cost. Designing a set of primers for amplicon-seq that targets the region followed by massively parallel sequencing adopting multiple barcoding may provide a convenient approach to utilize the segments as an approximate marker. Because the cob-rnS segment is 3.5 to 4.2 kb compared to ~16 kbp full-length mtDNA, the sequencing costs will be greatly suppressed by adopting this approach.

Whether utilization of mtDNA-based markers, either entire or partial, for a phylogenetic approach is biologically appropriate remains a debatable issue. The appropriateness of the usage of partial mtDNA sequences for the approach is one point. Our study demonstrated the importance of the choice of the mtDNA segments to reproduce the phylogeny of the entire mtDNA depending on the organisms, and proposed the segments that may be utilized as approximate markers for animals. Further, whether the sequence variation of the entire
mtDNAs, which are only inherited maternally in most organisms and evolve faster than the nuclear genome, properly represents the genetic variation of the tested individuals has yet to be evaluated. For example, mtDNA is maternally inherited in most species, and therefore it may give a biased view of population history [49]. The importance of an integrative approach based on the nuclear genome and mtDNA to estimate the species history has been suggested recently [50]. The several phylogenetic studies that have been conducted to date based on both mitochondrial genes and the nuclear internal transcribed spacer (e.g., [51–53]) may contribute to obtaining an integrative evaluation of both mitochondrial and nuclear molecular evolutions. Utilization of our approximate marker as well as a genome-wide approach, such as RAD-seq, may be useful to incorporate both nuclear and mtDNA variations so as to decipher the evolutionary history of a species in an integrative manner. Such approaches can also be adopted to understand the differential divergence of nuclear and mtDNA sequences in a variety of organisms. For non-vertebrate organisms, including invertebrates, plants, fungi, and protists, the partial mtDNA markers that show congruence with the entire mtDNA should be determined empirically.

**Conclusions**

Our findings may underscore the importance of the marker genes/segments choice on mtDNA to obtain proper information regarding the intraspecific genetic variations to analyze their phylogeny and population structure.

Our observations suggested that the choice of mitogene for phylogeographic or population genetic studies affected the results in a significant manner. This is a reasonable yet frequently disregarded premise. Due to the selection pressure on the function of the products, protein-coding sequences tend to be less divergent, resulting in smaller genetic distances and obscure clustering for phylogenetic analysis. In particular, the use of traditional marker sequences, whose products possess vital functions for the organism, should be considered with great care. Selection of a partial segment based on the comparison among entire mtDNA sequences of several individuals, as well as preliminary phylogenetic analysis based on a small number of samples, may improve the quality of the study. For such studies for mammalian, the segment cob-rnS, length of 3.5 ~ 4.2 kb, may serve as an excellent approximate marker.

**Materials and methods**

**Literature search**

The MEDLINE database was searched using the PubMed search engine to obtain published articles regarding mtDNA genetic variation, with a particular focus on phylogeographic and population genetics. Because usage of the complete mtDNA sequence for phylogenetic and population structure analysis was already the norm for humans in 2006 [34], these studies were excluded. Specifically, we used the Boolean search string “(mitochondrial AND (DNA or genome)) AND ((phylogeography [MeSH Terms]) OR (genetic, population[MeSH Terms])) NOT human,” then further narrowed the search with the following strings: for tRNA, (transfer RNA) OR (trna); for ribosomal RNA, (ribosomal RNA) OR (rRNA); for cytochrome c oxidase subunits, (cytochrome c oxidase) OR cox OR COI OR COII OR COIII OR CO1; for cytochrome b, (cytochrome b) OR (apocytochrome b) OR cob; for NADH dehydrogenase subunits, (NADH dehydrogenase) OR (nad); for D-loop, (“D Loop”) OR (“control region”); for intergenic regions, (“intergenic region”) OR (“non-coding region”) NOT (“D loop”) OR (“control region”); and for complete mtDNA, complete. The strings were selected to optimize the coverage of MeSH vocabularies.
Datasets

The dataset consisting of complete and unambiguous mtDNA sequences of >25 strains/individuals from the same taxonomic ID (organisms) with information available on their geographic origin was selected for this study. As a result of the process, we finally chose 10 datasets: *Apodemus agrarius* (striped field mouse, 42 individuals), *Camelus bacterianus* (domestic camel, 114 individuals), *Capra hircus* (domestic goat, 112 strains), *Canis lupus* (gray wolf, 31 individuals), *Gallus* (domestic red junglefowl, 96 individuals), *Pan troglodytes* (chimpanzee, 80 individuals), and *Ursus arctos* (brown bear, 115 individuals), *H. akashiwo* (a protist, causative species of harmful algal bloom, 26 isolates), *Fusarium culmorum* (fungal plant pathogen causing seedling rot, foot rot, and ear blight to a wide range of monocots and dicots, 33 isolates), and *F. graminearum* (fungal plant pathogen causing Fusarium head blight on wheat, barley, and rice and ear rot on maize, 75 isolates). An appropriate dataset was not available from the plant kingdom.

Preparation for the analyses

Because all of the sequences analyzed in this study were circular, we first aligned the sequences to be analyzed and redefined the start position of each genome at an identical position. The gene coding sequences (CDSs) on the genomes were re-annotated using MFannot https://megasun.bch.umontreal.ca/cgi-bin/dev_mfa/mfannotInterface.pl using the translation table for the specific taxa, and the CDSs were excised using the EMBOS package (http://emboss.sourceforge.net).

Sequence heterogeneity analysis

To analyze the sequence heterogeneity, the entire mtDNA sequences were aligned by the MAFFT package. The extent of variation at each aligned nucleotide position, heterogeneity (H), was calculated as

$$H = 1 - \sum p_i^2, \quad (i = A, T, G, C, \text{and gap})$$

where $p_i$ is the probability of the occurrence of $i$, and $\sum$ stands for the summation over the five possibilities [54]. H of each segment was further processed using a sliding window approach with window size 10 and sliding by 5 for *U. arctos, C. lupus, C. hircus, G. gallus, C. bacterianus*, and *H. akashiwo*. The H and the open reading frames (ORFs) of the mtDNAs were visualized using ArcWithColor software that is bundled in the GenomeMatcher package [55] (http://www.ige.tohoku.ac.jp/joho/gmProject/gmhomeJP.html).

Phylogenetic reconstruction and the tree correlation and concordance analyses. For the phylogenetic studies, the PRANK and MAFFT package was utilized for the MSA.

The PRANK alignments were conducted using either the codon-conscious mode for protein-coding sequence or the nucleotide mode for the non-coding region [56]. Two *Fusarium* species possess protein-coding sequences that contain both introns and exons, thus the codon-conscious alignment was not adopted. The alignment was verified by the GUIDANCE2 package [57], and the nucleotides aligned at the scores >0.93 were extracted and subjected to further phylogenetic reconstructions using the iqtree [58] or MrBayes [59] packages. In the PRANK-Guidance-based analyses of the sequences of the entire mtDNA or more than one gene and intergenic regions, the aligned sequences were concatenated and subjected to downstream analyses. The MAFFT alignments were directly applied to the analysis by iqtree. The phylogeny trees were visualized using the Geneious package [60] or the FigTree package.
The correlations among the phylogenies based on the different datasets were evaluated using the corrplot function of R package ape (http://ape-package.ird.fr/ [61]). The concordance between the phylogenetic trees constructed based on the full-length mtDNA and partial sequences was evaluated using R package treespace (https://cran.r-project.org/web/packages/treespace/index.html [62]) “treeConcordance” functions. Because the latter analysis is designed to evaluate the concordance of the clade topologies of two phylogenetic trees, three organisms, A. agrarius, P. troglodytes, and U. arctos, that yielded phylogenies supported by high posterior probability values were subjected to the analyses. For analyzing the concordance of the partial mtDNA sequences to the entire mtDNA, 26 individuals of A. agrarius, 47 individuals of P. troglodytes, and 69 individuals of U. arctos, which correspond to ~60% of the available datasets, were randomly selected using the “rand” function of the perl package and subjected to the analyses.

**Supporting information**

S1 Fig. Correlations between MAFFT-aligned MSA-based phylogenies of the full-length mtDNA sequences and those of the indicated genes/segments on the mtDNA and PRANK-aligned full-length mtDNA MSA-based phylogenies.

(TIF)

S2 Fig. Phylogenetic divergence of C. bacterianus full-length mtDNA.

(TIF)

S3 Fig. Phylogenetic divergence of C. lupus full-length mtDNA.

(TIF)

S4 Fig. Phylogenetic divergence of C. hircus full-length mtDNA.

(TIF)

S5 Fig. Phylogenetic divergence of G. gallus full-length mtDNA.

(TIF)

S6 Fig. Concordances of the phylogenetic trees constructed based on the MAFFT-aligned full-length mtDNA and partial mtDNA sequences of (A) A. agrarius, (B) P. troglodytes, and (C) U. arctos.

(TIF)

S7 Fig. Correlations between the MAFFT-aligned MSA-based phylogenies of full-length mtDNA sequences and those of indicated genes/segments on mtDNA.

(TIF)

S1 Table. The number of phylogeographic and population genetics studies conducted for various organisms based on mitochondrial markers.

(DOCX)

**Acknowledgments**

We would like to thank Ms. Patricia Beristain and Ms. Hilda Peye (Laboratorio de Genética, Acuicultura & Biodiversidad, Universidad de Los Lagos) for their scientific and administrative support.

**Author Contributions**

**Conceptualization:** Masanao Sato, Shoko Ueki.
Data curation: Jesús Morón-López, Karen Vergara, Shoko Ueki.

Formal analysis: Jesús Morón-López, Karen Vergara, Shoko Ueki.

Funding acquisition: Shoko Ueki.

Investigation: Jesús Morón-López, Masanao Sato, Shoko Ueki.

Methodology: Masanao Sato, Shoko Ueki.

Project administration: Shoko Ueki.

Resources: Jesús Morón-López, Karen Vergara, Shoko Ueki.

Software: Masanao Sato, Shoko Ueki.

Supervision: Shoko Ueki.

Validation: Masanao Sato, Shoko Ueki.

Visualization: Shoko Ueki.

Writing – original draft: Jesús Morón-López, Shoko Ueki.

Writing – review & editing: Masanao Sato, Gonzalo Gajardo, Shoko Ueki.

References

1. Gray MW, Burger G, Lang BF. Mitochondrial evolution. Science. 1999; 283(5407):1476–81. https://doi.org/10.1126/science.283.5407.1476 PMID: 10066161

2. Gabaldon T, Huynen MA. Shaping the mitochondrial proteome. Biochim Biophys Acta. 2004; 1659(2–3):212–20. https://doi.org/10.1016/j.bbabio.2004.07.011 PMID: 15576054

3. Smith DR, Keeling PJ. Mitochondrial and plastid genome architecture: Reoccurring themes, but significant differences at the extremes. Proc Natl Acad Sci U S A. 2015; 112(33):10177–84. https://doi.org/10.1073/pnas.1422049112 PMID: 25814499

4. Saccone C, Gissi C, Reyes A, Larizza A, Sbisa E, Pesole G. Mitochondrial DNA in metazoa Degree of freedom in a frozen event. Gene. 2002; 286:3–12. https://doi.org/10.1016/s0378-1119(01)00807-1 PMID: 11943454

5. da Fonseca RR, Johnson WE, O’Brien SJ, Ramos MJ, Antunes A. The adaptive evolution of the mammalian mitochondrial genome. BMC Genomics. 2008; 9:119. https://doi.org/10.1186/1471-2164-9-119 PMID: 18318906

6. James JE, Piganeau G, Eyre-Walker A. The rate of adaptive evolution in animal mitochondria. Mol Ecol. 2016; 25(1):67–78. https://doi.org/10.1111/mec.13475 PMID: 26578312

7. Sbisa E, Tanzariello F, Reyes A, Pesole G, Saccone C. Mammalian mitochondrial D-loop region structural analysis: identification of new conserved sequences and their functional and evolutionary implications. Gene. 1997; 205(1–2):125–40. https://doi.org/10.1016/s0378-1119(97)00404-6 PMID: 9461386

8. Fernandez-Silva P, Enriquez JA, Montoya J. Replication and transcription of mammalian mitochondrial DNA. Exp. Physiol. 2003; 88(1):41–56. https://doi.org/10.1113/eph8802514 PMID: 12525654

9. Lavrov DV, Pett W. Animal mitochondrial DNA as we do not know it: mt-genome organization and evolution in nonbilaterian lineages. Genome Biol Evol. 2016; 8(9):2896–913. https://doi.org/10.1093/gbe/evw195 PMID: 27557826

10. Gualberto KJ., K KJ. Plant mitochondrial genomes: Dynamics and mechanisms of mutation. Annu Rev Plant Biol. 2017; 68:225–52. https://doi.org/10.1146/annurev-arplant-040315-112232 PMID: 28226235

11. Gualberto JM, Mileshina D, Wallet C, Niazi AK, Weber-Lotti F, Dietrich A. The plant mitochondrial genome: Dynamics and maintenance. Biochimie. 2014; 100:107–20. https://doi.org/10.1016/j.biochi.2013.09.016 PMID: 24075874

12. Sloan DB, Alverson AJ, Chuckalovcak JP, Wu M, McCauley DE, Palmer JD, et al. Rapid evolution of enormous, multichromosomal genomes in flowering plant mitochondria with exceptionally high mutation rates. PLoS Biol. 2012; 10(1):e1001241 https://doi.org/10.1371/journal.pbio.1001241 PMID: 22272183

13. Chatre L, Richetti M. Are mitochondria the Achilles' heel of the Kingdom Fungi? Curr Opin Microbiol. 2014; 20:49–54. https://doi.org/10.1016/j.mib.2014.05.001 PMID: 24908191
14. Forget L, Ustinova J, Wang Z, Huss VAR, Lang BF. Hyaloraphidium curvatum A linear mitochondrial genome, tRNA editing, and an evolutionary link to lower fungi. Mol Biol Evol. 2017; 8:310–9.
15. Aguileta G, de Vienne DM, Ross ON, Hood ME, Giraud T, Petit E, et al. High variability of mitochondrial gene order among fungi. Genome Biol Evol. 2014; 6(2):451–65. https://doi.org/10.1093/gbe/evu028 PMID: 24504088
16. Lambowitz AM, Belfort M. Mobile bacterial group II introns at the crux of eukaryotic evolution.pdf. 2015.
17. Franco MEE, Lopez SMY, Medina R, Lucentini CG, Troncozo MJ, Pastorino GN, et al. The mitochondrial genome of the pathogenic fungus Stemphylium lycopersici uncovers a dynamic structure due to repetitive and mobile elements. PLoS One. 2017; 12(10):e0185545.
18. Wideman JG, Monier A, Rodriguez-Martinez R, Leonard G, Cook E, Poirier C, et al. Unexpected mitochondrial genome diversity revealed by targeted single-cell genomics of heterotrophic flagellated protists. Nat Microbiol. 2020; 5(1):154–65. https://doi.org/10.1038/s41564-019-0605-4 PMID: 31768028
19. Nash EA, Nisbet RER, Barbrook AC, Howe CJ. Dinoflagellate: A mitochondrial genome at all sea. Trends Genet. 2008; 24:328–35. https://doi.org/10.1016/j.tig.2008.04.001 PMID: 18514360
20. Jackson CJ, Gornik SG, Waller RF. The mitochondrial genome and transcriptome of the basal dinoflagellate Hematodinium sp.: Character evolution within the highly derived mitochondrial genomes of dinoflagellates. Genome Biol Evol. 2011; 4:58–72.
21. Feagin JE, Mericle BL, Werner E, Morris M. Identification of additional rRNA fragments encoded by the Plasmodium falciparum 6 kb element. 1997; 25:438–46.
22. Burger G, Gray MW, Forget L, Lang BF. Strikingly bacteria-like and gene-rich mitochondrial genomes throughout jakobid protists. Genome Biol Evol. 2013; 5(2):418–38. https://doi.org/10.1093/gbe/evt008 PMID: 23335123
23. Imanian B, Keeling PJ. The dinoflagellates Durinska baltica and Kryptoperidinium foliaceum retain functionally overlapping mitochondria from two evolutionarily distinct lineages. BMC Evol Biol. 2007; 7 (172).
24. Jackson CJ, Norman JE, Schnare MN, Gray MW, Keeling PJ, Waller RF. Broad genomic and transcriptional analysis reveals a highly derived genome in dinoflagellate mitochondria. 2007; 5:1–17.
25. Kamikawa R, Inagaki Y, Sako Y. Fragmentation of mitochondrial large subunit rRNA in the dinoflagellate Alexandrium catenella and the evolution of rRNA structure in alveolate mitochondria. Protist. 2007; 158:239–45.
26. Waller RF, Jackson CJ. Dinoflagellate mitochondrial genomes: Stretching the rules of molecular biology. BioEssays. 2009; 31:237–45. https://doi.org/10.1002/bies.200800164 PMID: 19204978
27. Avise JC, Arnold J, Martin Bal R, Beringhame E, Lamb T, Neigel JE, et al. Intraspecific Phylogeography: The mitochondrial DNA bridge between population genetics and systematics. BMC Genomics. 2008; 9(119).
28. Keis M, Remm J, Ho SYW, Davison J, Tammeleht E, Tumanov IL, et al. Complete mitochondrial genomes and a novel spatial genetic method reveal cryptic phylogeographical structure and migration patterns among brown bears in north-western Eurasia. Journal of Biogeography. 2013; 40(5):915–27.
29. Hirata D, Mano T, Abramo AV, Baryshnikov GF, Kosintsev PA, Vorobiev AA, et al. Molecular phylogeny of the brown bear (Ursus arctos) in Northeastern Asia based on analyses of complete mitochondrial DNA sequences. Mol Biol Evol. 2013; 30(7):1644–52.
30. Davison J, Ho SYW, Bray SC, Korsten M, Tammeleht E, Hindrikson M, et al. Late-Quaternary biogeographic scenarios for the brown bear (Ursus arctos), a wild mammal model species. Quaternary Science Reviews. 2011; 30(3–4):418–30.
31. Higashi A, Nagai S, Salomon PS, Ueki S. A unique, highly variable mitochondrial gene with coding capacity of Heterosigma akashiwo, class Raphidophyceae. Journal of Applied Phycology. 2017.
32. Higashi A, Nagai S, Seone S, Ueki S. A hypervariable mitochondrial protein coding sequence associated with geographical origin in a cosmopolitan bloom-forming alga, Heterosigma akashiwo. Biol Lett. 2017; 13(4).
33. Ueki S. Phylogeographic characteristics of hypervariable regions in the mitochondrial genome of a cosmopolitan, bloom-forming raphidophyte, Heterosigma akashiwo. J Phycol. 2019; 55(4):858–67.
34. Torroni A, Achilli A, Macaulay V, Richards B, Bandelt HJ. Harvesting the fruit of the human mtDNA tree. Trends Genet. 2006; 22(6):339–45. https://doi.org/10.1016/j.tig.2006.04.001 PMID: 16678300
35. Wolters JF, Chiu K, Fiumeri HL. Population structure of mitochondrial genomes in Saccharomyces cerevisiae. BMC Genomics. 2015; 16:451.
36. Bergstrom A, Simpson JT, Salinas F, Barre B, Parts L, Zia A, et al. A high-definition view of functional genetic variation from natural yeast genomes. Mol Biol Evol. 2014; 31(4):872–88. https://doi.org/10.1093/molbev/msu037 PMID: 24425782
37. Andersen LW, Jacobsen M, Vedel-Smith C, Jensen TS. Mice as stowaways? Colonization history of Danish striped field mice. *Biol Lett.* 2017; 13(7). https://doi.org/10.1098/rsbl.2017.0064 PMID: 28679695

38. Ming L, Yi L, Sa R, Wang ZX, Wang Z, Ji R. Genetic diversity and phylogeographic structure of Bactrian camels shown by mitochondrial sequence variations. *Anim Genet.* 2017; 48(2):217–20. https://doi.org/10.1111/age.12511 PMID: 27775167

39. Pilot M, Dabrowski MJ, Hayrapetyan V, Yavruyan EG, Kopaniani N, Tsingarska E, et al. Genetic variability of the grey wolf *Canis lupus* in the Caucasus in comparison with Europe and the Middle East: Distinct or intermediary population? *PLoS One.* 2014; 9(4):e93828.

40. Li H, Meng X, Zhang H, Duan X, Niu L, Wang L, et al. Complete mitochondrial genome of Nanjiang Yellow goat (*Capra hircus*). *Mitochondrial DNA A DNA Mapp Seq Anal.* 2016; 27(2):1383–4.

41. Kulik T, Brankovics B, van Diepeningen AD, Bliska K, Zelechowski M, Mysczynski K, et al. Diversity of mobile genetic elements in the mitogenomes of closely related *Fusarium culmorum* and *F. graminearum* sensu stricto strains and its implication for diagnostic purposes. *Front Microbiol.* 2020; 11:1002.

42. Nishibori M, Shimogori T, Hayashi T, Yasue H. Molecular evidence for hybridization of species in the genus *Gallus* except for *Gallus varius*. *Anim Genet.* 2005; 36(5):367–75.

43. Bjork A, Liu W, Wertheim JO, Hahn BH, Worobey M. Evolutionary history of chimpanzees inferred from complete mitochondrial genomes. *Mol Biol Evol.* 2011; 28(1):615–23. https://doi.org/10.1093/molbev/msq227 PMID: 20802239

44. Flegontov P, Michalek J, Janouskovec J, Lai DH, Jirku M, Hajduska E, et al. Divergent mitochondrial respiratory chains in phototrophic relatives of apicomplexan parasites. *Mol Biol Evol.* 2015; 32(5):1115–31. https://doi.org/10.1093/molbev/msv021 PMID: 25660376

45. Md Mukarram Hossain AS, Blackburne BP, Shah A, Whelan S. Evidence of statistical inconsistency of phylogenetic methods in the presence of multiple sequence alignment uncertainty. *Genome Biol Evol.* 2015; 7(8):2102–16. https://doi.org/10.1093/gbe/evv127 PMID: 26139831

46. Jordan G, Goldman N. The effects of alignment error and alignment filtering on the sitewise detection of positive selection. *Mol Biol Evol.* 2012; 29(4):1125–39. https://doi.org/10.1093/molbev/msq272 PMID: 22049066

47. Blackburne BP, Whelan S. Class of multiple sequence alignment algorithm affects genomic analysis. *Mol Biol Evol.* 2013; 30(3):642–53. https://doi.org/10.1093/molbev/mss256 PMID: 23144040

48. Löytynoja A, Goldman N. Phylogeny-aware gap placement prevents errors in sequence alignment and evolutionary analysis. *Science.* 2008; 320:1632–5. https://doi.org/10.1126/science.1158395 PMID: 18566285

49. de Groot GA, Nowak C, Skrbinský T, Andersen LW, Aspi J, Fumagalli L, et al. Decades of population genetic research reveal the need for harmonization of molecular markers: the grey wolf *Canis lupus* as a case study. *Mammal Review.* 2016; 46(1):44–59.

50. Radzvilavicius A, Layh S, Hall MD, Dowling DK, Johnston IG. Sexually antagonistic evolution of mitochondrial and nuclear linkage. *J Evol Biol.* 2021; 34(5):757–66. https://doi.org/10.1111/jeb.13776 PMID: 33644926

51. Zakeri Z, Otte V, Sipman H, Malicek J, Cubas P, Rico VJ, et al. Discovering cryptic species in the *Aspicilium intermutans* complex (Megasporaceae, Ascomycota)—First results using gene concatenation and coalescent-based species tree approaches. *PLoS One.* 2019; 14(5):e0216675.

52. Baron S, van der Merwe NA, Maritz-Olivier C, Solorzano-Garcia B, Garcia-Varela M. The genetic relationship between *R. microplus* and *R. decoloratus* ticks in South Africa and their population structure. *Mol Phylogenet Evol.* 2018; 129:60–9.

53. Lopez-Jimenez A, Hernandez-Mena DI, Solorzano-Garcia B, Garcia-Varela M. Exploring the genetic structure of *Parastrigea diovadana* Dubois and Macko, 1972 (Digenia; Strigeidae), an endoparasite of the white ibis, *Eudocimus albus*, from the Neotropical region of Mexico. *Parasitol Res.* 2021; 120(6):2065–75.

54. Keim P, Beavis W, Schupp J, Freestone R. Evaluation of soybean RFLP marker diversity in adapted germ plasm. *Theor Appl Genet* 1992; 85:205–12. https://doi.org/10.1007/BF00222861 PMID: 24197306

55. Ohtsubo Y, Ikeda-Ohtsubo W, Nagata Y, Tsuda M. GenomeMatcher: A graphical user interface for DNA sequence comparison. *BMC Bioinformatics* 2008; 9:376. https://doi.org/10.1186/1471-2105-9-376 PMID: 18793444

56. Löytynoja A. Phylogeny-aware alignment with PRANK. In: Russell DJ, editor. *Multiple Sequence Alignment Methods.* Totowa, NJ: Humana Press; 2013. p. 155–70.

57. Sela I, Ashkenazy H, Katoh K, Pupko T. GUIDANCE2: Accurate detection of unreliable alignment regions accounting for the uncertainty of multiple parameters. *Nucleic Acids Res.* 2015; 43(W1):W7–14. https://doi.org/10.1093/nar/gkv318 PMID: 25883146
58. Nguyen LT, Schmidt HA, von Haeseler A, Minh BQ. IQ-TREE: A fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. *Mol Biol Evol*. 2015; 32(1):268–74. https://doi.org/10.1093/molbev/msu300 PMID: 25371430

59. Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Hohna S, et al. MrBayes 3.2: Efficient Bayesian phylogenetic inference and model choice across a large model space. *Syst Biol*. 2012; 61(3):539–42. https://doi.org/10.1093/sysbio/sys029 PMID: 22357727

60. Kumar S, Stecher G, Li M, Knyaz C, Tamura K. MEGA X: Molecular evolutionary genetics analysis across computing platforms. *Mol Biol Evol*. 2018; 35:1547–9. https://doi.org/10.1093/molbev/msy096 PMID: 29722887

61. Paradis E, Schliep K. Ape 5.0: An environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics*. 2019; 35:526–8. https://doi.org/10.1093/bioinformatics/bty633 PMID: 30016406

62. Jombart T, Kendall M, Almagro-Garcia J, Colijn C. treespace: Statistical exploration of landscapes of phylogenetic trees. *Mol Ecol Resour*. 2017; 17(6):1385–92. https://doi.org/10.1111/1755-0998.12676 PMID: 28374552