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Extreme Mitogenomic Variation in Natural Populations of Chaetognaths

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Abstract

The extent of within-species genetic variation across the diversity of animal life is an underexplored problem in ecology and evolution. Although neutral genetic variation should scale positively with population size, mitochondrial diversity levels are believed to show little variation across animal species. Here, we report an unprecedented case of extreme mitochondrial diversity within natural populations of two morphospecies of chaetognaths (arrow worms). We determine that this diversity is composed of deep sympatric mitochondrial lineages, which are in some cases as divergent as human and platypus. Additionally, based on 54 complete mitogenomes, we observed mitochondrial gene order differences between several of these lineages. We examined nuclear divergence patterns (18S, 28S, and an intron) to determine the possible origin of these lineages, but did not find congruent patterns between mitochondrial and nuclear markers. We also show that extreme mitochondrial divergence in chaetognaths is not driven by positive selection. Hence, we propose that the extreme levels of mitochondrial variation could be the result of either a complex scenario of reproductive isolation, or a combination of large population size and accelerated mitochondrial mutation rate. These findings emphasize the importance of characterizing genome-wide levels of nuclear variation in these species and promote chaetognaths as a remarkable model to study mitochondrial evolution.

Key words: genetic diversity, mitochondrial genomes, molecular evolution, marine invertebrates, Chaetognatha.

Introduction

Genetic diversity estimates provide information about a species’ population history, dynamics, and adaptive potential (Ellegren and Galtier 2016; Leffler et al. 2012; Peijnenburg and Goetze 2013; Reed and Frankham 2003). Mitochondrial genes have been widely used to characterize natural populations at the molecular level, mostly because of technical ease-of-use considerations such as clonality and high mutation rate (Galtier et al. 2009). In particular, the DNA barcoding approach has stimulated the use of mitochondrial gene fragments, such as Cytochrome Oxidase 1 (cox1). The combination of strong interspecific divergence and generally low levels of intraspecific variation, also known as the barcoding “gap”, makes such markers an essential tool for species identification (Bucklin et al. 2011). Many observations of high levels of mitochondrial divergence have consequently been interpreted as evidence of cryptic speciation, the distinct genetic entities being often, but not always, associated with previously overlooked geographic, morphological, or reproductive boundaries (Bickford et al. 2007).

The observation that the range of genetic variation does not scale with population size as predicted by the neutral theory of molecular evolution was referred to by Lewontin as the “paradox of variation” (Hahn 2008; Lewontin 1974).
This paradox is particularly manifest for animal mitochondrial diversity levels, which show surprisingly little variation across species (Bazin et al. 2006). This has been interpreted as the consequence of pervasive positive selection (Bazin et al. 2006), of genetic hitch-hiking (Gillespie 2001), or as the result of an inverse correlation between population size and mutation rate (Piganeau and Eyre-Walker 2009). Similarly, mitochondrial transplantation experiments have demonstrated that the cyto-nuclear interactions in respiratory complexes are an important factor limiting diversity in mitochondrial genes (Kenyon and Moraes 1997; Ellison and Burton 2000). These explanations may account for the fact that the combination of high genetic diversity and signatures of strong purifying selection has rarely been observed in nature for mitochondria, including in species with large population sizes (Meiklejohn et al. 2007).

The genetics and diversity of oceanic invertebrates remain poorly characterized (Bucklin et al. 2011; GIGA Community of Scientists et al. 2014; Peijnenburg and Goetze 2013). Chaetognaths are an enigmatic marine invertebrate phylum, which possess a unique combination of morphological, developmental, and genomic characters (Telford 2004). As a possible sister group to other protostomes, chaetognaths occupy a key phylogenetic position to understand the evolution of bilaterian animals (Marlé taz et al. 2006, 2008; Matus et al. 2006). They also represent a major planktonic group and play important roles in marine food webs as the primary predators of copepods (Bone et al. 1991). Unambiguous fossils of chaetognaths have been described from the lower Cambrian (Vannier et al. 2007), which suggests that the phylum originated during the “Cambrian explosion” and underwent little morphological evolution during the course of animal evolution (Knoll and Carroll 1999). However, the phylum has a poor recent fossil record, which prevents dating divergences of the main extant clades and its taxonomy is still unresolved (Papillon et al. 2006; Gasm i et al. 2014). Transcriptomic studies revealed that chaetognath genomes underwent extensive gene duplication; many genes are transcribed in operons, of- 

### Materials and Methods

#### Animal Sampling and Genotyping

Individuals of the species *Spadella cephaloptera*, were collected from Calanque de Sormiou near Marseille (France) at shallow depth on *Posidonia* seagrass using a plankton net (supplementary fig. S1, Supplementary Material online). Individuals of the species *Sagitta elegans* and *S. setosa* were collected using a multinet on a single daytrip in the Gullmar fjord near Kristineberg (Lysekil, Sweden). All chaetognaths were identified under a microscope while still alive by taxonomic experts (FM and KTCAP). Only mature individuals without food in their guts and without visible parasites were preserved and used for genetic analysis. Genomic DNA was extracted using QIAamp Micro Kit and DNeasy Blood & Tissue Kit (Qiagen) and used as a template for PCR amplification using GoTaq (Promega) or Phire Hot Start II DNA Polymerase (Finzymes). Number of individuals for species and each amplified marker are specified in table 1 and supplementary table S1, Supplementary Material online. Nuclear markers (18S, 28S, nuclear intron of the ribosomal protein L36a) were amplified in the same individuals for which the entire mitogenome was sequenced. Primers and protocols used for 18S and 28S characterization are described in (Papillon et al. 2006). Nuclear loci were cloned in pGEM-T (Promega) and 2–8 clones were picked and sequenced in both directions. All primers employed are specified in supplementary table S5, Supplementary Material online. All sequences were deposited in Genbank (accessions in supplementary table S1, Supplementary Material online).

#### Sequencing of Individual Mitochondrial Genomes

In *S. cephaloptera*, two half-genome fragments (6–8 kb) were amplified using specific primers located in 16S and cox1 genes using the Accuprime Taq (Invitrogen) for five individuals. These fragments were purified using S.N.A.P. gel purification kit (Invitrogen) and subsequently cloned with Topo XL cloning Kit (Invitrogen). Plasmid DNA was prepared with Plasmid Midi Kit (Qiagen). These templates were sequenced using a primer walking strategy using 6–10 Sanger reads in total. In 37 *S. elegans* and 12 *S. setosa* individuals, outward pointing primers were designed within the cox1 and the cox2 genes and used to amplify the mitogenome as a single amplicon using Phusion DNA polymerase (Thermo scientific). For eight individuals long-range PCR fragments of both cox1 and cox2 primer sets were sequenced to verify that the same mitochondrial genome was obtained. A unique library was
Table 1

| Species/Location | Gene | n | K | \( \pi \) | \( \theta_w \) | \( D_f \) |
|------------------|------|---|---|---------|---------|------|
| *Spadella cephaloptera* | *cox1* | 25 | 25 | 0.1385 | 0.1223 | 0.5319 |
| *Coelosoma* & *Cephalocauda* | *cox2* | 29 | 29 | 0.1181 | 0.1149 | 0.1066 |
| *Sormiou, France* | All genes | 5 | 5 | 0.2939 | 0.2470 | 1.4414 |
| *Sweden* | *cox1* | 107 | 96 | 0.1775 | 0.0854 | 3.6069 |
| *Gullmar fjord* & *Sweden* | *cox2* | 108 | 101 | 0.2043 | 0.1113 | 2.7856 |
| *Sagitta elegans* | *cox1* | 37 | 37 | 0.2081 | 0.1304 | 2.1922 |
| *S. cephaloptera* | *cox2* | 24 | 1 | 0.0000 | 0.0000 | NA |
| *Sweden* | *cox1* | 37 | 36 | 0.0411 | 0.0443 | 0.3562 |
| *Sagitta setosa* | *cox1* | 54 | 49 | 0.0088 | 0.0214 | 0.2046 |
| *Gullmar fjord* | *cox2* | 54 | 53 | 0.0138 | 0.0305 | 1.9015 |
| *Sweden* | All genes | 12 | 12 | 0.0088 | 0.0139 | 1.6561 |
| *S. cephaloptera* | *cox1* | 20 | 2 | 0.0001 | 0.0002 | 1.1643 |
| *Gullmar fjord* | *cox2* | 20 | 6 | 0.0006 | 0.0015 | 1.7800 |

\( \pi \), Watterson estimator; \( \theta_w \), synonymous substitution rate; \( D_f \), Tajima’s D. For “all genes,” average values are shown. For more details, see supplementary table S1, Supplementary Material online.

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Molecular Evolution and Population Genetic Analysis

Population genetic parameters were calculated using the EggLib python library (De Mita and Siol 2012). NeighborNet Networks were built using Splitstree4 with K2P distances (Huson and Bryant 2006). Maximum Likelihood (ML) phylogenetic inferences were performed using RAxML assuming the GTR+\( \Gamma \) model for nucleotide data sets and the MtZoa +\( \Gamma \) model for amino acid data sets (Rota-Stabelli et al. 2009; Stamatakis 2014) (fig. 2A). Nonsynonymous and synonymous substitution rates were estimated using the yn00, site-model and branch model using the PAML package (supplementary table S4, Supplementary Material online) (Yang 2007).

Survey of Mitochondrial Genetic Diversity in Metazoans

NCBI popset database was searched for data sets corresponding to *cox1* gene fragments (keyword: *Cox1*, *COI*, *CO1*), including a single species and at least ten sequences. We obtained 1581 records corresponding to 1,079 species. We focused on the 437 data sets associated with a referenced publication (pmid). A codon alignment was built using nucleotide translation discarding partial sequences and those containing degenerate bases (e.g., “N”, “Y”, or “R”). We computed nucleotide diversity statistics using the egg-lib library (De Mita and Siol 2012) and we selected the data set showing the highest diversity in each species (supplementary table S2, Supplementary Material online). The 50 data sets with highest levels of genetic diversity were manually curated for both the sequence alignments and corresponding papers.

Transcriptome Data

RNA was extracted from ~100 mixed embryonic stages and juveniles of *S. cephaloptera* using the Qiagen RNA micro kit (Invitrogen). Normalized cDNA libraries were built by BioS&T Inc. and sequenced according to manufacturer’s instruction on a GS GLX instrument (Roche).

Results

Extreme Mitochondrial Diversity in Two Chaetognath Morphospecies

We characterized mitochondrial variation of individual chaetognaths from three species sampled at single geographic localities in *Sormiou* (France) and Gullmar fjord (Sweden) (supplementary fig. S1, Supplementary Material online). All chaetognaths were examined under a microscope while still alive and identified to species level based on morphological characters. We sequenced mitochondrial gene fragments in more than 180 individuals, and full mitochondrial genomes in 54 representative individuals (table 1 and supplementary table S1, Supplementary Material online). We calculated nucleotide diversity (\( \pi \)), which summarizes the average number of
nucleotide difference per site in a population sample, as well as uncorrected sequence divergence for the cox1 gene. In *S. cephaloptera* (*N* = 25), we found *π* = 0.14 and median sequence divergence of 17.99% (pairwise maximum = 23.68%). In *S. elegans* (*N* = 107), we found *π* = 0.18 and median sequence divergence of 20.80% (pairwise maximum = 24.96%). In both species, phylogenetic analysis of individual mitochondrial genes, such as 16S rRNA or cox1, revealed that this diversity is distributed across 8–11 highly divergent lineages (fig. 1; supplementary fig. S2 and Data set S1, Supplementary Material online). However, we did not detect such high levels of mitochondrial diversity in a third species, *S. setosa* (cox1: *N* = 54, *π* = 0.009, 0.91% median sequence divergence), which is a closely related species to *S. elegans*, and was sampled at the same location in Gullmar fjord (table 1).

We never recovered multiple products in cross-amplification experiments, nor consistent double Sanger chromatogram peaks, which would be indicative of multiple mitochondrial copies per individual (Breton et al. 2007) or nuclear pseudogenes (Bensasson et al. 2001). All mitochondrial genome amplicons included the 13 genes typical of chaetognaths and we also did not find any stop codons or frameshift mutations despite global nucleotide diversities of 0.29 and 0.21 in *S. cephaloptera* and *S. elegans*, respectively (table 1). To further ascertain the validity of our PCR-based sequence data, we examined a transcriptome data set obtained from >100 pooled individuals of *S. cephaloptera*, all sampled from the same location as the genotyped individuals. We found transcripts of multiple mitochondrial genes, including cox1, with unambiguous assignment of all five lineages for which a whole mitogenome was sequenced in *S. cephaloptera* (supplementary Data set S2, Supplementary Material online). These multiple lines of evidence reject explanations such as PCR artifacts, the presence of multiple mitochondrial copies (heteroplasm) or the presence of nuclear pseudogenes as possible explanations for the extreme levels of mitochondrial diversity observed here.

**Nuclear Variation Is Incongruent with Deep Mitochondrial Lineages**

We compared the patterns of mitochondrial variation with those inferred from three nuclear markers: the variable ribosomal protein L36a intron (fig. 1) and the more conserved 18S and 28S rRNA genes (table 1). The ribosomal genes 18S and 28S are commonly used to resolve interspecific relationships within the phylum because of their strong interspecific divergence (Gasmi et al. 2014; Papillon et al. 2006; Telford and Holland 1997). If deep mitochondrial lineages represent ancient, reproductively isolated, species, we would expect to see congruent divergence patterns across independently evolving nuclear loci (Coyne and Orr 2004; Hudson and Turelli 2003).

For *S. elegans*, we characterized variation at three nuclear loci: 18S and 28S markers showed no variation at all, but the nuclear intron of L36a was quite variable with 68 alleles detected in 37 individuals (table 1). A gene tree of the L36a intron shows a complete mixing of individuals with highly divergent mitochondrial genomes (fig. 1) and hence, displays incongruent patterns of divergence with the mitochondrial gene tree for this species. We also found significant levels of recombination in the nuclear intron data set, as would be expected for an interbreeding population of individuals (phi-test, *P* value = 0). Conversely, for *S. cephaloptera* we obtained information for only two nuclear markers and the results are inconclusive. The nuclear intron of L36a was sequenced for 30 individuals and we recovered 36 different alleles. These nuclear alleles arrange in four supported clades, which appear incongruent with deep mitochondrial lineages (fig. 1). We only sequenced eight individuals for 18S, which was quite variable in this species with 27 variable sites, but more genotyping data would be required to determine whether individuals belonging to deep mitochondrial clades in *S. cephaloptera* are diverged at nuclear loci. Nevertheless, phylogenetic trees combining our 18S and 28S sequences from genotyped individuals with those of other available chaetognath species (supplementary figs. S3 and S4, Supplementary Material online, respectively) show that divergences within morphospecies *S. elegans* and, to a lesser extent, also *S. cephaloptera*, are generally much lower than those observed between chaetognath species.

Inconsistency between mitochondrial and nuclear gene trees can be the result of hybridization between, usually closely related, species, resulting in introgression of mtDNA from one species to the other (Toews and Brelsford 2012). However, we consider this unlikely because in a gene tree based on clustered cox1 sequences from 29 chaetognath species (five species of which show deep mitochondrial clades >10% divergent), we found that all mitochondrial lineages in each species share a common ancestor (i.e., represent a monophyletic group) and hence are never shared between species (supplementary fig. S5, Supplementary Material online).

Among the three sampled nuclear markers with varying evolutionary rates, none showed patterns of divergence congruent with the observed deep mitochondrial lineages in either of two chaetognath species. More nuclear loci should be characterized though to fully assess whether completely or partially isolated gene pools are present in natural populations of *S. cephaloptera* and *S. elegans*.

**Extreme Divergence between Mitochondrial Lineages**

Mitochondrial genes annotated in sequenced genomes consistently show high divergence and congruent topologies as expected for a haploid, nonrecombining genome (supplementary Data set S2, Supplementary Material online). We thus examined phylogenetic relationships and divergence
using the complete set of mitochondrial protein-coding sequences along with existing mitogenomic data from other chaetognath species (fig. 2A and supplementary fig. S6, Supplementary Material online) (Helfenbein 2004; Miyamoto et al. 2010a; Papillon et al. 2004). We found that this tree recovers with good support the accepted relationships between chaetognath species (Gasmi et al. 2014; Papillon et al. 2006), and unambiguously supports the monophyly of mitogenomes from all three species analysed in this study.

To put mitochondrial divergence levels into perspective, we compared the phylogenetic distance between chaetognath mitochondrial lineages with those inferred between the main vertebrate groups (fig. 2B). We found that maximum-likelihood distances between mitochondrial lineages of S. cephaloptera and S. elegans compare with those recovered within amniote and mammalian lineages, which diverged 312 and 170 Myr ago, respectively (Hedges and Kumar 2009). For instance, the distance between the Sr3 lineage to the common ancestor of S. cephaloptera is 0.707 substitution per site,

![Figure 1](https://example.com/fig1.png)
whereas the distance from human to the amniote ancestor is 0.671 substitutions per site. Similarly, the distance between human and platypus is 0.630 and that between lineage A and S. elegans is 0.717 substitutions per site. Although speculative, this analysis suggests that deep mitochondrial lineages in chaetognaths are associated with ancient divergence, strong mutation rate acceleration, or a combination of the two.

FIG. 2.—Mitochondrial divergence in chaetognaths and chordates. Phylogenetic trees based on the concatenation of all protein-coding mitochondrial genes in chaetognath lineages (A) and chordates (B) at the same scale (expected amino acid changes per site). Reconstructions were performed using Maximum Likelihood (MZOA + Γ model). Maximum bootstrap support values are indicated by plain circles on nodes. In chordates, ML branch lengths were inferred from the alignment according to accepted topology. Sample sizes of Sagitta elegans lineages are indicated in brackets (detailed in supplementary fig. S6, Supplementary Material online).
Mitochondrial Diversity Values Represent Extremes Amongst Animals

Nucleotide diversity estimates for two chaetognath species represent the highest values reported so far for any metazoan as established by surveying public databases for all available population cox1 data sets (fig. 3 and supplementary table S2, Supplementary Material online). We classified the data sets as single species or possible cryptic species by analysing the corresponding papers. S. elegans shows the highest level of synonymous diversity ($\pi_S = 0.646$) followed by S. cephaloptera ($\pi_S = 0.476$). Conversely, S. cephaloptera is the more variable species at the coding level ($\pi_N = 0.023$) followed by S. elegans ($\pi_N = 0.018$). A few data sets that were considered to result from cryptic speciation showed slightly higher diversity levels (open circles in fig. 3). We find that both chaetognath species harbor more intraspecific variation than several established cases of extreme mitochondrial diversity driven by (micro-) allopatric divergence, such as the crustacean *Tigriopus californicus* ($\pi_S = 0.404$) and the gastropod *Cepaea nemoralis* ($\pi_S = 0.418$) (Burton and Lee 1994; Thomaz et al. 1996).

Mitochondrial Rearrangements between Lineages

During the annotation of the mitochondrial genomes of *Sagitta elegans*, we observed gene order differences between individual genomes, which appear strictly associated with lineages. This suggests that beyond fast molecular divergence, these lineages underwent unusual structural rearrangements (fig. 4). We confirmed the validity of these structural changes by checking the read alignments. We found no coverage discontinuity associated with breakpoints and we recovered the same arrangements when sequencing multiple genomes from the same individuals (supplementary Data set S2, Supplementary Material online).

In *S. elegans*, at least two lineages exhibit rearrangements compared with the standard gene order involving the genes *nad1*, *nad2*, and *cox3* in one case, and *nad4L* in the other (fig. 4B). Structural changes in mitogenomes also affect the size and proportion of intergenic regions, which are highly variable between lineages, ranging from 8.3% in lineage H to 25.4% in lineage G. Conversely, the fraction of noncoding DNA is remarkably stable within lineages with at most 3% variation (supplementary fig. S7, Supplementary Material online). These intergenic regions do not contain palindromic motifs which would be indicative of a repetitive element origin (Lavrov 2010).

Changes in gene order are commonly observed between species (Miyamoto et al. 2010a; Papillon et al. 2004), but to our knowledge such rearrangements have never been described as part of the diversity of a single animal species before. We also identified partial *cox1* extra-copies inserted in intergenic regions of *S. cephaloptera* mitogenomes Sce-2 and Sce-6, which had accumulated coding substitutions (grey regions in fig. 4A). These duplicates support the model that considers gene duplications as intermediate steps in structural rearrangements of the mitochondrion (Moritz et al. 1987). Such dynamic rearrangements could be responsible for the extreme reduction in size and gene number of chaetognath mitogenomes compared with other bilaterian animals, with the loss of *atp6*, *atp8*, and *tRNA* genes (Helfenbein 2004; Papillon et al. 2004).

Patterns of Selection in Mitochondrial Genomes

Because high levels of coding nucleotide diversity could be the result of positive selection, we examined the patterns of nonsynonymous versus synonymous variation in mitochondrial genes (Oliveira et al. 2008). We found low or moderate $\pi_N/\pi_S$ ratios (average 0.170 for *S. elegans* and 0.049 for *S. cephaloptera*, fig. 5A and supplementary table S3, Supplementary Material online) as well as low intraspecific $d_N/d_S$ ratios between lineages (average 0.086 for *S. elegans* and 0.097 for *S. cephaloptera*, supplementary table S3, Supplementary Material online). We further tested whether individual lineages contribute unequally to the global $d_N/d_S$ estimates by assigning independent $d_N/d_S$ ratios to each lineage in individual genes. No particular lineage or gene showed any evidence of relaxation of selective pressure, though some genes seem more prone to higher coding variation than others in certain lineages, such as *nad6* in Lineage G or *nad3* in Lineage C (fig. 5B). Similarly, likelihood-ratio tests did not provide significant support for positive selection affecting specific lineages or sites (supplementary table S4, Supplementary Material online).
Supplementary Material online). These analyses consistently indicate that positive selection is not responsible for the divergence of mitochondrial lineages in *S. cephaloptera* and *S. elegans*, instead, the low d_{\omega}/d_{S} values suggest that chaetognath mitochondrial lineages evolved mainly under the influence of purifying selection.

**Discussion**

We report the presence of highly divergent mitochondrial lineages in individuals of the chaetognaths *S. cephaloptera* and *S. elegans* sampled at single geographic sites. We further demonstrate that these deep mitochondrial lineages are split by molecular divergences equivalent to those observed among tetrapods (fig. 2) and contain structural rearrangements (fig. 4). The mechanism through which such divergent lineages could appear within these natural populations remains to be ascertained. Mainly, there are two possible interpretations of our results: deep lineages represent cryptic species and are reproductively isolated, or, alternatively, deep lineages are present within interbreeding populations. Although cryptic speciation is a common explanation for such extreme mitochondrial divergence, we did not observe a pattern of divergence at any of the examined nuclear loci consistent with reproductive isolation (fig. 1; supplementary figs. S3 and S4, Supplementary Material online). Moreover, the high number of observed mitochondrial lineages in both species (more than eight) in sympatry would require a complicated scenario of multiple isolation events followed by secondary contact. Our present data set, however, does not incorporate a sufficient number of nuclear loci to decide between these evolutionary scenarios, and particularly, to confidently rule out the possibility that lineages are reproductively isolated. In addition, our sampling was limited to single geographic localities for both species, which makes it difficult to estimate which part of the molecular diversity may be attributable to spatial variation. Hence, extending our geographical sampling and acquiring a broader sampling of nuclear markers across the genome—for instance using RNA-seq or target capture methods (Gayral et al. 2013)—would be necessary to differentiate between the two alternative interpretations of our results.

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**Fig. 4.**—Structural variation between individual mitochondrial genomes of chaetognaths. (A) Annotated mitochondrial genomes with gene positions drawn to scale. The proportion of intergenic regions (white) varies from 8% to 26.4% in *Sagitta elegans* lineages and from 3.7% to 18.4% in *Spadella cephaloptera* lineages. *cox1* duplicates in *S. cephaloptera* are in grey. (B) Schematic representation of inferred mitochondrial gene order rearrangements in two *Sagitta elegans* mitochondrial lineages (B and G) with respect to the most frequent gene order (lineages A, C, D, E, F, and H). See also supplementary fig. S7, Supplementary Material online.
If the alternative interpretation is correct and highly divergent lineages are present within interbreeding populations of chaetognaths, they likely appeared because of an accelerated mitochondrial mutation rate, and were maintained because of large population sizes. By considering synonymous and nonsynonymous rates of substitutions, we ruled out that the extent of mitochondrial variation across animal mitochondrial lineages would fit theoretical expectations predicting that a large population size allows for the maintenance of ancestral polymorphisms (Hahn 2008; Lewontin 1974), and would challenge previous reports suggesting that the extent of mitochondrial variation across animals is limited (Bazin et al. 2006).

Since S. cephaloptera and S. elegans are distantly related in the chaetognath phylogenetic tree (belonging to the orders Phragmophora and Aphragmophora, respectively (Papillon et al. 2006; Gasiñi et al. 2014)) and have very distinct ecologies (benthic and planktonic, respectively), extreme mitochondrial diversity is probably not related to the particular demographic characteristics of species. Previous reports of deep mitochondrial lineages in other chaetognath species are consistent with this claim, even though these have often been interpreted as evidence of cryptic speciation (Kulagin et al. 2014; Miyamoto et al. 2010b). Not all chaetognath morphospecies harbor extreme levels of mitochondrial diversity though, as we show for the species S. setosa (table 1). The reduced mitochondrial variation in this species (global $\pi = 0.0088$) compared with other chaetognaths has been related to the fragmented, coastal populations of S. setosa, which probably suffered severe population bottlenecks during Pleistocene glacial periods (Peijnenburg et al. 2004, 2005, 2006). As previously stated, more extensive geographic sampling for S. cephaloptera and S. elegans and the genotyping of a broader collection of nuclear markers could shed further light on the origin and maintenance of these deep mitochondrial lineages in chaetognaths.

An accelerated mitochondrial mutation rate could be related to the reduction of chaetognath mitochondrial genomes and their propensity to structural rearrangements (Miyamoto et al. 2010a). Such a combination of extreme size reduction and accelerated mutation rate was also reported for a ctenophore (Pett et al. 2011). The size reduction and peculiar architecture of animal mitochondrial genomes compared with other eukaryotes, such as green plants, was attributed to the combined effect of genetic drift and accelerated mutation rates (Lynch et al. 2006). Hence, the processes that gave rise to the extreme mitochondrial diversity in chaetognaths could be similar to those governing mitochondrial evolution in other metazoans, but only differ by the extent of mutation rate acceleration and (effective) population sizes.

If highly divergent lineages are present within interbreeding populations, this would imply that cyto-nuclear interactions at the respiratory complexes are much less constrained than originally thought (Kenyon and Moraes 1997). Indeed, the interactions between mitochondrial and nuclear subunits should be robust enough to cope with multiple divergent mitochondrial lineages present in interbreeding populations.
of chaetognaths. Mitochondrial transplantation experiments originally demonstrated that cyto-nuclear interactions could be maintained between species, but were dramatically altered when divergence increased (e.g., to human and orang-utan), resulting in a decreased efficiency of respiratory processes (Kenyon and Moraes 1997). Similar effects were reported in hybrids of divergent *Tigriopus californicus* populations, which showed reduced respiratory fitness and altered gene expression (Ellison and Burton 2008a, 2008b). We hypothesize that chaetognaths cope with high levels of mitochondrial variation through an increased robustness in the interaction with nuclear subunits of oxidative phosphorylation complexes. How this increased robustness relates to the original genomic traits of chaetognaths, such as gene duplication, represents an interesting future topic of investigation.

We report the presence of highly divergent mitochondrial genomes in different individuals from the same morphospecies sampled at single geographic localities. Whether these highly divergent genomes are present within interbreeding populations or are attributable to a complex reproductive isolation scenario remains to be determined using a larger collection of nuclear loci and broader geographic sampling. Nevertheless, this finding challenges established views of the amount of mitochondrial diversity in animal species and populations. An assessment of intraspecific mitochondrial diversity in other animal groups, particularly those with dynamic mitochondrial genomes, such as ctenophores, urochordates, or brachiopods, may uncover other extreme cases of intraspecific variation (Pett et al. 2011). Until more of these hyperdiverse taxa are identified, chaetognaths represent an interesting and emerging model to understand the molecular evolution of animal mitochondrial genomes.

### Data Accessibility

- Assembled mitochondrial genomes have been deposited in Genbank under the accessions KP899748-KP899801 (details in supplementary table S1, Supplementary Material online).
- Sequences of genotyped markers are available under the accessions: KP843748-KP843841 and KP857119-KP857568 (details in supplementary table S1, Supplementary Material online).
- Transcriptome data was deposited with the accession PRJNA357934.
- Genotyped markers, alignments, and reconstructed trees are also available as supplementary Data sets S1 and S2, Supplementary Material online at [https://figshare.com/s/a4706e9adc1ecea55880](https://figshare.com/s/a4706e9adc1ecea55880).

### Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

### Author Contribution

F.M., Y.L.P., and K.P. conceived the project and carried out the animal sampling. F.M. and S.L. performed the experiments. F.M., S.L., and Y.L.P. analyzed the sequencing data. F.M. and K.P. wrote the paper.

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