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# The mitochondrial genomes of Campodea fragilis and C. Iubbocki (Hexapoda: Diplura): high genetic divergence in a morphologically uniform taxon <br> L. Podsiadlowski*, A. Carapelli†, F. Nardi†, R. Dallai $\dagger$, M. Koch*, J.L. Boore $\ddagger$, F. Frati $\dagger$ 

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#### Abstract

Mitochondrial genomes from two dipluran hexapods of the genus Campodea have been sequenced. Gene order is the same as in most other hexapods and crustaceans. Secondary structures of tRNAs reveal specific structural changes in tRNA-C, tRNA-R, tRNA-S1 and tRNA-S2. Comparative analyses of nucleotide and amino acid composition, as well as structural features of both ribosomal RNA subunits, reveal substantial differences among the analysed taxa. Although the two Campodea species are morphologically highly uniform, genetic divergence is larger than expected, suggesting a long evolutionary history under stable ecological conditions.


## Introduction

Mitochondrial sequence data are commonly used in phylogenetic studies from large scale phylogenetics down to the population level. With few exceptions, mitochondrial genomes from multicellular animals are circular DNA molecules, about 16 Kb in size, containing 37 genes encoding for two ribosomal RNA molecules (rrnS and IrrS), 13 protein subunits (cox1-3, cob, nad1-6 and nd4L, atp6 and atp8) and 22 transfer RNA molecules (trnX) Wolstenholme, 1992; Boore, 1999). In addition, mitochondrial genomes of insects bear one large $\mathrm{A}+\mathrm{T}$ rich noncoding region, which is involved in the initiation of mitochondrial replication and transcription (Wolstenholme, 1992), and is therefore commonly referred to as the mitochondrial control region.

Because of the ease of amplification and sequencing, the mitochondrial genome has been extensively studied for phylogenetic purposes, as well as for its peculiar patterns of molecular evolution (Boore, 1999). Concerning the latter, the mitochondrial genome has been found to exhibit extensive nucleotide compositional bias. One of the best examples of mitochondrial compositional bias is that observed in insects (Simon et al., 1994), and some other arthropods (e.g.: Black et al., 1998; Negrisolo et al., 2004), where a significant higher amount of Adenine and Thymine nucleotides ( $\mathrm{A}+\mathrm{T}$ bias) has been found. Another type of compositional bias is that affecting base content at Protein Coding Genes (PCGs) in the different strands of the mitochondrial genome (Perna \& Kocher, 1995). In a thorough investigation of this strand-specific bias across Metazoa, Hassanin et al. (2005) found clear evidence of an asymmetric mutational bias, as well as traces of the reversal of such a bias in some taxa due to an inversion of the control region.

Gene rearrangements of the mitochondrial genome are also well known, and several mechanisms have been proposed to explain gene order changes, involving duplication events followed by the random (Boore, 2000) or non-random (Lavrov et
al., 2004) loss of some duplicated copies. These events sometimes leave unusually large fragments of non-coding DNA, in addition to the canonical control region (Lavrov et al., 2002).

While partial sequences from the control region or from single genes, like cox1, cob, $r r n S$, have been utilised in phylogenetic studies at the species or population level, complete mitochondrial genome data have been used predominantly in phylogenetic analyses above the family level (e.g: Nardi et al., 2003; Negrisolo et al., 2004; Cameron et al., 2005; Cook et al., 2005). In most of the latter cases only PCGs were studied. However, complete mitochondrial genomes provide more data than just sequence information:
(1) Gene order is not conserved among metazoa and gene translocations may serve as characters in phylogenetic analyses. For example the "Pancrustacea" hypothesis a close relationship between Crustacea and Hexapoda (excluding the Myriapoda, which were traditionally viewed as sistergroup to hexapods) - has been suggested on the basis of a gene translocation involving a mitochondrial tRNA gene (Boore et al., 1998).
(2) Secondary structure features of tRNAs and rRNAs are seldomly used as characters in phylogenetic analyses but with a growing amount of data these may be of greater value soon (e.g. Macey et al., 2000).
(3) The mitochondrial genetic code differs in some codons from the standard code and also varies among metazoan taxa. These deviations are of phylogenetic value as well (e.g. Castresana et al., 1998; Telford et al., 2000).

During the last ten years the traditional view of arthropod phylogeny has been strongly challenged by molecular analyses. As a result the Pancrustacea hypothesis is now well supported by mitochondrial gene rearrangement data (Boore et al., 1998) and sequence based analyses of both, mitochondrial (Wilson et al. 2000; Hwang et al. 2001; Lavrov et al., 2004) and nuclear genes (Mallatt et al., 2004, Regier et al.,
2005). On the other hand, crustacean and even hexapod monophyly, as well as their internal relationships, are still under debate (Regier et al., 2005; Babbitt \& Patel, 2005; Podsiadlowski \& Bartolomaeus, 2005). In particular, some analyses using complete mitochondrial datasets do not support the monophyly of Hexapoda, with different crustacean subtaxa forming sistergroups to Collembola and Insecta (Nardi et al., 2003; Cook et al., 2005). In this context, apterygote taxa play a crucial role, and their reciprocal relationships have not yet been resolved with certainty (Kristensen, 1997; Kukalova-Peck, 1987; Stys \& Bilinski, 1990; Carapelli et al., 2000). Recently, the first mitochondrial genome of a member of Diplura, the japygid Japyx solifugus, has been published (Carapelli et al., 2005) and the corresponding phylogenetic analysis suggests a paraphyletic origin of Hexapoda. On the other hand detailed analyses of hexapod relationships based on the nuclear 18S rRNA results in hexapod monophyly (Kjer, 2004; Babitt \& Patel, 2005). The increased use of rRNA sequences in phylogenetic studies has stimulated the debate over the correct alignment procedures. One approach calls for using the information contained in rRNA secondary structures (Kjer, 1995, 2004), which, in turn, are inferred from general models based on the comparison of a large number of sequences (Hickson et al., 1996).

In this study we present two additional mitochondrial genomes from apterygote hexapods: Campodea fragilis and C. lubbocki (Diplura: Campodeidae). We provide detailed analyses of nucleotide and amino acid frequencies, and compositional bias, as well as the reconstruction of the putative secondary structures of tRNA and rRNA genes. Although a lot of mitochondrial genomes from arthropods are published now, only a few congeneric species are compared. The genus Campodea has a worldwide distribution and comprises about 150 soil-dwelling, herbivorous species, sized up to 5 mm and highly uniform in appearance. The two species studied exhibit only a few morphological differences: spatulate (C.I.) or acuminate (C.f.) praetarsal
appendages; presence (C.I.) or absence (C.f.) of mediane macrochaetae on abdominal tergites; annulated (C.I.) or smooth (C.f.) cerci (Palissa, 1964). Both seem to occupy similar ecological niches but their genetic divergence turned out to be rather high, suggesting a long evolutionary history under stable environmental conditions.

## Results and Discussion

## Genome organisation

Both Campodea mitochondrial genomes presented here share the same 37 genes with most bilaterian animals and have the same gene arrangement (Fig. 1) as other apterygote hexapods (Tricholepidion gertschi, NC005437; Nesomachilis australica, NC006895; Japyx solifugus AY771989), insects (Drosophila melanogaster NC001709), and crustaceans (Daphnia pulex NC000844; Penaeus monodon NC002184). This gene order appears to be the ancestral state for Pancrustacea (Crustacea+Hexapoda) and differs from that of chelicerates and myriapods by the translocation of trnL1 (Boore et al., 1998).

In both Campodea species gene overlaps exist between the PCGs atp8/atp6 (7 nucleotides) and nad4/nad4L (7 nucleotides). Other overlaps occur between tRNA genes or between a tRNA- and a protein-coding gene. Only a few overlaps are larger than 3 nucleotides, notably at the trnY/cox1 boundary in both species (8nt in $C$. lubbocki and 5nt in C. fragilis). A large non-coding region is present between rrnS and trnl (558nt in C. fragilis and 561nt in C. lubbocki). This is, by all evidence available, homologous to the mitochondrial control region (CR), which, in insects, is usually called AT-rich region for exhibiting a remarkable A+T-bias (up to $>90 \%$ in hymenopterans; Crozier \& Crozier, 1993). In Campodea, the putative AT-rich region
shows A+T content between 84.2\% (C. fragilis) and 89.1\% (C. lubbocki). More unusual, other non-coding (ncod) sequences are found between other mitochondrial genes of the two Campodea species. One of them, shared by both species, is located between trnS2 and nad1 (56nt in C. fragilis, 80nt in C. lubbocki), while only C. fragilis has an additional ncod region of 111nt between nad2 and trnW (Fig. 1). Interestingly, the two ncod regions can be folded in apparently stable secondary structures (Fig. 2). Most other arthropods lack non-coding sequences larger than a few nucleotides (with the control region as an exception). Although very unusual, the occurrence of large non-coding regions abutting gene junctions has been observed in insects and myriapods (Lavrov et al., 2002; Bae et al., 2004). Sometimes these fragments are mentioned as remnants of gene duplication events due to possible errors during the replication of the mitochondrial genome (Boore, 2000). Extra copies of mitochondrial genes are subsequently eliminated from the compactly arranged mitochondrial genome (Boore, 1999). The occurrence of these structures would suggest that the remnant of an exceeding copy of a tRNA gene is still present in the genomes of the taxa under study. In addition, the ncod fragments found in both Campodea species between trnS2 and nad1 clearly exhibit homologous patterns in either primary sequence and secondary structure. Accordingly, this feature would suggest that the presumed duplication event has occurred before the speciation of the two taxa.

A+T content of complete mitochondrial genomes varies among insects (Table 1). While Japyx solifugus (Japygidae), the only other dipluran species with published mitogenomic data, has the lowest value among hexapods (64.8\%), C. lubbocki has the highest value among apterygote insects (74.9\%), followed by two collembolans (Gomphiocephalus hodgsoni and Tetrodontophora bielanensis) and C. fragilis (72.6\%). Comparatively, high values are also found in Pterygota, while archaeognathan and zygentoman species range between 67-69\%. In all species,
third-codon position sites of PCGs have significantly higher A+T content than either first and second codon positions. Functional reasons for the strong differences of $\mathrm{A}+\mathrm{T}$ content among hexapods are still under debate.

## Protein coding genes

There is no significant difference in the size of the PCGs of $C$. fragilis and C. lubbocki compared to each other and to other apterygote hexapods (Table 2). The three starting codons ATA, ATG and ATT are used in both species, while only in C. lubbocki the starting codon ATC is used for atp8, nad1 and nad5. In some cases stop codons are truncated (cox1-3 in C. fragilis; cox3, nad3, nad4 and nad6 in C. lubbocki) and possibly post-transcriptionally completed to TAA after cleavage (Okimoto et al., 1990, Lavrov et al., 2002).

Nucleotide sequence similarity of PCGs between the two Campodea species ranges from $49.5 \%$ (nad6) to $77.9 \%$ (cox2) (Table 2). As an example, the pairwise similarity of cox1-cox3 between C. fragilis and C. lubbocki (77.1\%) is rather low: in the same region, four lxodes species show levels of nucleotide divergence ranging from 75\% to $82 \%$, five Drosophila-species are in the range $92-99 \%$, while two species of the crustacean Triops (T. cancriformis and T. longicaudatus) differ by 81.2\%, and two species of Bombyx (B. mori and B. mandarina) by $96 \%$. In contrast, the divergence between the two Campodea sequences is more similar to that observed between species from different orders, such as in the comparison between Pteronarcys princeps (Pterygota: Plecoptera) and five Drosophila species (78-79\%), or the comparison of $P$. princeps with the zygentoman Tricholepidion gertschi (73.5\%). Frati et al. (1997) provided a comparison between different collembolan species and other hexapods based on cox2 sequences. Levels of nucleotide similarity for cox2 between congeneric species (genera Orchesella and Isotomurus) were only slightly higher (80.4-81\%) than that observed between C. fragilis and C. lubbocki $(77.9 \%$ or
$\mathbf{7 8 . 5} \%$ ?). On the other hand, nucleotide similarity of cox2 between Campodea species and J. solifugus varies between $67.4 \%$ and $67.8 \%$, well in the range of that estimated, in Collembola, between species from different lineages (61.9\%-69.7\%), or between collembolan species and other pterygote insects (60.4\%-68.9\%). Looking at the inferred amino acid sequence similarity, the values estimated in diplurans (68.1\%-69.5\% between Campodea and J. solifugus) are comparable to those found between species from different orders among pterygote insects (Simon et al., 1994). We conclude that, although morphologically very uniform, the taxon Campodea is genetically highly diverse and that, under the molecular clock hypothesis, the split between the two species occurred comparably long ago, perhaps as long as some of the basal splits among pterygote insects. Indeed, taxonomic categories above the species do not necessarily imply similar genetic and morphological differentiation, nor similar age in different lineages. Our results in Campodea parellel those obtained in collembolans (Frati et al., 1997, 2000) in indicating high levels of genetic divergence (in itochondrial sequences) even between congeneric species which do not exhibit comparable morphological differentiation. These results point towards either accelerated rates of evolution or remarkably older age of genera of apterygote hexapods.

As shown by Hassanin et al. (2005), PCGs differ for their A vs. T and C vs. G content according to the strand (J- or N-strands; sensu Simon et al., 1994) where they are encoded. Genes encoded on the J-strand show a slight bias towards Ts over As and a strong bias towards Cs over Gs, while genes encoded on the N -strand show a much stronger bias towards Ts over As and towards Gs over Cs (Table 2, Fig. 3). One reason for this outcome might be the asymmetrical replication process, during which one strand is single-stranded for a longer time than the other, and therefore more susceptible for specific mutations (for more details see: Hassanin et al., 2005).

Relative synonymous codon usage in PCGs (Table 3) reveals that the genome-wide base compositional bias for $\mathrm{A}+\mathrm{T}$ is also reflected in codon usage. The two Campodea species differ in preferred codons for Glycine, Proline and Threonine, whereas both species differ from J . solifugus in codon preferences for Alanine, Histidine and Valine. In the case of Histidine, this reflects the strong differences in A+T content, as in J.solifugus codon the codon CAC is preferred, whereas Campodea species prefer CAT. A similar observation is evident in the case of Leucine, for which TTA is the preferred codon in all three diplurans, but to a much stronger degree in Campodea than in J. solifugus.

## Transfer-RNA coding genes

In both species all 22 tRNA-encoding genes usually found in mitochondrial genomes of metazoans are present. In both species the most likely secondary structure models for tRNA-R, tRNA-S1 and tRNA-S2 lack the D-arm (Fig. 4). In C. lubbocki also the tRNA-C has a shortened (or missing) D-arm. In contrast, all tRNAs mentioned above show the usual cloverleaf secondary structure in J. solifugus (Fig. 4), as well as all remaining tRNAs in C. fragilis and C. lubbocki (data not shown). Accordingly, a reduction of the D-arm of tRNA-R, and the two types of tRNA-S, must have evolved after the split between Campodeidae and Japygidae. Reduction of the D-arm of tRNA-C probably occured even later, after the split between C. Iubbocki and C. fragilis. These data may be of value for phylogenetic inferences, when mitogenomic data will be available from more dipluran species.

Ribosomal-RNA coding genes
The two genes for ribosomal RNA subunits (rrnL for 16 S rRNA and rrnS for 12 S rRNA) can be folded in an inferred secondary structure composed of paired and single strand fragments (Figs. 5, 6). Comparative analysis shows that both rRNA
subunits are composed of a mosaic of variable and conserved sequence fragments, which are distributed in paired and unpaired regions of all domains (Fig. 7). This is probably due to variable functional constraints, acting at different degrees in specific parts of the rRNA subunits, which are necessary to maintain the ribosome functionality. This evidence suggests that different rates of nucleotide substitution affect each domain of rrnL and rrnS. Sequence similarity in aligned (based on secondary structure) rRNAs of C. lubbocki and C. fragilis is within the range of sequence similarity of the PCGs (rrnL: 74.3\%; rrnS: 73.8\%).

In general, core structural elements of C. fragilis and C. lubbocki rRNA subunits closely resemble those proposed for Drosophila melanogaster. The inferred secondary structure of the 12S rRNA of each species (Fig. 5) displays a considerable degree of differentiation in terms of length and shape of most of domains I and II (Fig. 7a), with conserved stretches of sequences present only on helices $h$ and $m$. Conversely, domain III is probably the most conserved fragment of the entire 12 S rRNA subunit, showing many invariant structural elements between $C$. fragilis and $C$. lubbocki (Figs. 5, 7a: all helices from o to a1 with the exception of $u$ ) and some identical stretches of primary sequence. With few exceptions aside (Page, et al. 2002), this evidence is in agreement with the data obtained from broader comparisons that include extended datasets of animal mitochondrial 12 S rRNA sequences (Hickson et al., 1996; Simon et al., 1996).

Domain I is probably the most variable part of the 16 S rRNA subunit (Fig. 6), differing in terms of length and primary sequence between C. fragilis and C. lubbocki. On domain II the highest levels of sequence variability can be observed in the most peripheral helices ( $g, j, k$ and $o$ ), in the descendant part of helix $I$ and in the single strand segments connecting helices $I, m$ and $n$ (Fig. 7b). An extensive primary sequence homology can be observed in the fragment enclosed between the ascendant and the descendant parts of helix $t$.

## Experimental procedures

## C. lubbocki; DNA isolation and PCR

Campodea lubbocki was collected in the garden of the Institute for Zoology, Freie Universität Berlin. One individual was cut into pieces and directly used for rolling circle amplification with the Templify kit (Amersham) following the manufacturer's protocol. Templify treated material was diluted with water (1:5) and used as DNA template in PCR. Two additional specimens were ethanol fixed for SEM, in order to determine the species according to Palissa (1964).

Initially five partial mitochondrial sequences (cox1, nad5, cob, rrnL, rrnS) were determined with PCR primer pairs designed for this purpose by looking for conserved regions of mitochondrial genes from other hexapod and crustacean sequences (Podsiadlowski \& Bartolomaeus, 2005). PCR primers were purchased from metabion (Germany). PCR was performed on Mastercycler and Mastercycler Gradient (Eppendorf, Hamburg, Germany) using the Eppendorf HotMasterTaq kit. $50 \mu \mathrm{l}$ reaction volumes were set up as follows: $42 \mu \mathrm{l}$ sterilized distilled water, $5 \mu \mathrm{l} 10 \mathrm{x}$ reaction buffer $1 \mu \mathrm{l}$ dNTP mix (Eppendorf) $1 \mu \mathrm{l}$ primer mix ( $10 \mu \mathrm{M}$ each), $1 \mu \mathrm{l}$ DNA template, $0,2 \mu \mathrm{l}(1 \mathrm{u})$ HotMasterTaq polymerase. The cycling protocol included an initial denaturation step ( $94^{\circ} \mathrm{C}, 2 \mathrm{~min}$ ), 40 cycles of denaturation $\left(94^{\circ} \mathrm{C}, 30 \mathrm{sec}\right.$ ), annealing (1 min, primer specific annealing temperature) and extension $\left(68^{\circ} \mathrm{C}, 90\right.$ $\mathrm{sec})$ and a final extension step $\left(68^{\circ} \mathrm{C}, 1 \mathrm{~min}\right)$. After agarose ( $0,9 \%$ ) gel separation and visualization of ethidium-bromide stained PCR products, purification for sequencing (see below) was performed using the PCR purification kit (Qiagen) or when necessary using the Gel extraction kit (Qiagen).

In a second step the determined sequences were used to design five additional PCR primer pairs bridging the gaps between them. PCR was performed as described above, except for using an extension time of seven minutes. PCR products were inspected and purified as described above.

## C. fragilis; $D N A$ isolation and $P C R$

Total DNA was extracted from a specimen of $C$. fragilis collected nearby the Belcaro Castle (Siena; Italia) using the Wizard SV Genomic DNA purification system (Promega). Preliminary amplification of short fragments of nad4 and rrnL, obtained using mitochondrial universal primers (Simon et al., 1994), were used to design species-specific primers to generate two long PCR products corresponding to the entire mitochondrial genome. The first long PCR product (about 6.5 Kb ) amplified the fragment between nd4 and rrnL, with the primer pair Camp-rRNAN (5'-GGTTGTATCGGAAGCTGCAGCTAG-3') and Camp-nd4J (5'-CTGGGTCGATAGTTTTGGCTGC-3'), using the following two-steps PCR conditions: 35 cycles at $96^{\circ} \mathrm{C}$ for 1 min and $68^{\circ} \mathrm{C}$ for 8 min , followed by incubation at $68^{\circ} \mathrm{C}$ for 15 min. The second fragment ( 8.5 Kb ) corresponding to the sequence encompassed between rrnL and nad4, was amplified with the primer pair Camp-rRNAJ and Campnd4R (corresponding to the reverse complementary of the previous pair), using the following PCR condition: 35 cycles at $96^{\circ} \mathrm{C}$ for 1 min and $68^{\circ} \mathrm{C}$ for 12 min , followed by incubation at $68^{\circ} \mathrm{C}$ for 15 min .

PCR reactions were performed on a GeneAmp ${ }^{\circledR}$ PCR System 2700 (Applied Biosystem) in $25 \mu$ reaction volume set up as follows: $10,75 \mu \mathrm{l}$ of sterilized distilled water, $2.5 \mu \mathrm{l}$ 10x reaction buffer, $2.5 \mu \mathrm{l}$ of $25 \mathrm{mM} \mathrm{MgCl} 2,4 \mu \mathrm{l}$ dNTP mix, $1.25 \mu \mathrm{l}$ of each primer $(10 \mu \mathrm{M}), 2.5 \mu \mathrm{l}$ DNA template and $0,25 \mu \mathrm{l}$ (1.25u) of TaKaRa LA Taq polymerase (Takara). Each PCR reaction yielded a single band when visualized with ethidium bromide staining after electrophoresis in a $1 \%$ agarose gel.

## Sequencing and sequence analysis

Sequencing of C. lubbocki mitochondrial DNA was carried out with a CEQ 8000XL automated DNA Analysis System (Beckman Coulter) using the CEQ DTCS Kit following the manufacturer's protocol, and using initially PCR primers and subsequently new internal primers until completion of sequences (primer walking). In C. fragilis, the two long PCR fragments were purified using a Montage PCR Centrifugal Filter Device (Millipore), and processed for DNA sequencing by the production facility of the DOE Joint Genome Institute (Walnut Creek, California). Flanking regions of the two long PCR sequences were then re-amplified using species-specific primers and sequenced in a CEQ 8000XL. All sequences were assembled using Sequencher 4.2.2 (Gene Codes) and the chromatograms were examined by eye to verify sequencing accuracy. The amount of clones sequenced provided a 10X average coverage for both long PCR fragments.

To determine gene identity BLAST search on NCBI Blast Entrez databases was used. Not determinable by primary sequence information alone, the presumed location of rRNA genes in both Campodea species has been determined according to the observed similarity in primary sequence with other hexapod taxa, and to comparison of 12 S and 16 S rRNA secondary structures between the two dipluran species and D. melanogaster. Start codons in protein coding genes were inferred to be the nearest start codon to the beginning of the sequence alignment of homologous genes with other hexapod species. Most tRNAs were identified using tRNAscan-SE 1.21 (Lowe \& Eddy, 1997) and DOGMA (Wyman et al., 2004), the remaining ones were found by eye inspection of the suspected regions. TransferRNA identity was specified by its anticodon sequence.

The secondary structures of the mitochondrial rRNA subunits (12S and 16S) for both Campodea species were derived by analogy with available models obtained for other
arthropods (Gutell et al., 1994; Misof \& Fleck, 2003; Cannone et al., 2002). Secondary structures of 12 S and 16 S rRNA genes and of intergenic non-coding regions were then visualized using the program RnaViz 2.0 (De Rijk \& DeWachter, 1997).

Nucleotide frequencies and codon usage was determined using DAMBE (version 4.2.13; Xia \& Xie, 2001). PCGs were aligned and compared using Bioedit (version 7.0.1; Hall, 1999).

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## Legends to Figures

Figure 1. Genome organization of the mitochondrial genomes of two Campodea fragilis and C. lubbocki. Arrows indicate direction of coding regions, numbers inside the circle represent the size of the fragments separating two adjacent genes (positive values) or the amount of shared nucleotides between two overlapping genes (negative values). Large non-coding regions are dashed, with the one located between $r r n S$ and $t r n /$ being the putative control region.

Figure 2. Hypothetical secondary structure features found in intergenic non-coding regions: ncod between trnS2 and nad1 in C. fragilis (a) and C. Iubbocki (b); ncod between nad2 and trnW in C. fragilis (c).

Figure 3. $\mathrm{A}+\mathrm{T}$ content, calculated in a sliding window of 100 bp , along the J -strand of the mitochondrial genomes of C. fragilis and C. lubbocki. The position and coding directions of protein- and rRNA-coding genes are shown below.

Figure 4. Putative secondary structures of mitochondrial tRNAs of C. fragilis, C. lubbocki, and Japyx solifugus. Only tRNAs exhibiting secondary structure changes in one or both Campodea species are shown. All other tRNAs of $C$. fragilis and $C$. lubbocki can be folded into cloverleaf-like secondary structures.

Figure 5. Secondary structure model of the mitochondrial 12 S rRNA in the two dipluran species.

Figure 6. Secondary structure model of the mitochondrial 16 S rRNA in the two dipluran species.

Figure 7. Generalized secondary structure model of Campodea mitochondrial 12S (a, domains I-III) and 16S (b, domains I, II, IV and V) rRNAs. Grey beads show the most variable regions, in base composition, sequence length or secondary structure, of the different domains in both species. Black beads represent highly conserved fragments.


C

## $\square \% \mathrm{~A} \quad$ ■T

## C. lubbocki

##  <br> C. fragilis



ND2 COI COII A8 A6 COIII ND3
$\xrightarrow{\text { ND5 ND4 ND4L }} \stackrel{\text { ND6 CytB }}{ }$





## Table 1

| Higher ranking taxon | Species | Accession number | PCG | $1^{\text {st }}$ codon Position | $2^{\text {nd }}$ codon position | $3^{\text {rd }}$ codon position | rRNAs | tRNAs | total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Diplura | Campodea fragilis | XX | 70.6 | 67.0 | 65.8 | 79.0 | 76.5 | 76.4 | 72.6 |
| " | Campodea lubbocki | XX | 73.0 | 68.3 | 66.9 | 84.2 | 80.3 | 76.1 | 74.9* |
| " | Japyx solifugus | AY771989 | 64.1 | 63.5 | 60.4 | 68.5 | 65.0 | 64.9 | 64.8 |
| Collembola | Gomphiocephalus hodgsoni | NC005438 | 72.4 | 66.6 | 65.5 | 86.2 | 77.0 | 76.0 | 74.1 |
| " | Tetrodontophora bielanensis | NC002735 | 71.3 | 66.4 | 66.2 | 81.4 | 77.0 | 75.5 | 72.7 |
| " | Podura aquatica | NC006075 | 64.7 | 60.0 | 64.1 | 70.0 | - | 70.5 | - |
| " | Onychiurus orientalis | NC006074 | 68.5 | 71.5 | 65.8 | 66.2 | - | 73.2 | - |
| Archaeognatha | Petrobius brevistylis | AY956355 | 66.2 | 61.1 | 63.2 | 74.2 | 71.7 | 70.6 | 67.3 |
| " | Nesomachilis australica | NC006895 | 67.2 | 60.8 | 64.4 | 76.8 | 72.6 | 70.3 | 68.8 |
| Zygentoma | Tricholepidion gertschi | NC005437 | 67.7 | 63.0 | 63.0 | 77.1 | 71.2 | 69.0 | 68.6 |
| " | Thermobia domestica | NC006080 | 66.3 | 59.4 | 62.9 | 76.6 | 67.9 | 67.2 | 67.0 |
| Pterygota | Locusta migratoria | NC001712 | 74.1 | 68.9 | 66.1 | 87.3 | 77.7 | 74.0 | 75.3 |
| " | Drosophila yakuba | X03240 | 76.7 | 69.8 | 66.4 | 93.7 | 81.9 | 76.6 | 78.6 |
| " | Apis mellifera | L06178 | 83.3 | 79.3 | 75.4 | 95.2 | 83.4 | 87.1 | 84.9 |

* incomplete genome data; missing sequence of control region or control region and rRNAs respectively

Table 2

| Gene <br> (Strand) | identity(\%) | C. fragilis |  |  |  |  |  | C. lubbocki |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | length | start/stop | \%A | \%C | \%G | \%T | lengt | start/stop | \%A | \%C | \%G | \%T |
| atp6 (+) | 73.3 | 675 | ATG/TAA | 31.3 | 20.6 | 10.7 | 37.5 | 675 | ATG/TAA | 30.7 | 18.4 | 9.6 | 41.3 |
| atp8 (+) | 57.0 | 156 | ATT/TAA | 38.5 | 14.7 | 4.5 | 42.3 | 156 | ATC/TAA | 35.3 | 18.0 | 5.8 | 41.0 |
| cox1 (+) | 77.4 | 1540 | ATT/T - - | 29.0 | 19.7 | 15.7 | 35.7 | 1542 | ATA/TAA | 29.0 | 18.6 | 14.6 | 37.8 |
| $\operatorname{cox} 2(+)$ | 77.9 | 679 | ATA/T - - | 33.1 | 19.3 | 11.6 | 35.9 | 684 | ATA/TAA | 33.2 | 17.0 | 10.5 | 39.3 |
| $\operatorname{cox} 3(+)$ | 76.6 | 787 | ATG/T -- | 30.9 | 21.1 | 14.1 | 33.9 | 787 | ATG/T - - | 28.7 | 18.7 | 13.1 | 39.5 |
| cob (+) | 76.5 | 1143 | ATG/TAA | 32.8 | 19.5 | 11.2 | 36.5 | 1140 | ATG/TAA | 32.5 | 17.5 | 10.8 | 39.2 |
| nad1 (-) | 70.7 | 924 | ATT/TAA | 26.4 | 10.5 | 16.3 | 46.8 | 921 | ATA/TAA | 29.0 | 8.3 | 17.3 | 45.5 |
| nad2 (+) | 61.3 | 1005 | ATA/TAA | 32.1 | 20.2 | 9.2 | 38.5 | 1005 | ATT/TAA | 32.4 | 16.9 | 8.0 | 42.7 |
| nad3 (+) | 62.9 | 357 | ATA/TAA | 35.0 | 18.2 | 7.8 | 38.9 | 347 | ATC/TA - | 34.9 | 16.7 | 7.8 | 40.6 |
| nad4 (-) | 68.7 | 1338 | ATG/TAA | 25.7 | 8.2 | 17.6 | 48.5 | 1326 | ATG/T - - | 27.9 | 6.9 | 17.2 | 48.0 |
| nad4l (-) | 74.0 | 285 | ATG/TAA | 28.4 | 4.9 | 20.4 | 46.3 | 285 | ATG/TAG | 27.4 | 1.8 | 21.4 | 49.5 |
| nad5 (-) | 73.7 | 1707 | ATA/TAA | 26.5 | 8.0 | 19.5 | 46.0 | 1710 | ATC/TAG | 30.0 | 7.3 | 17.5 | 45.2 |
| nad6 (+) | 49.5 | 510 | ATT/TAA | 38.4 | 17.1 | 4.7 | 39.8 | 524 | ATT/TA - | 37.6 | 16.8 | 4.8 | 40.8 |

Table 3

|  |  | C.I. | C.f. | J.s. | G.h. | T.b. | P.a. | O.o. | P.b. | N.a. | T.g. | T.d. | L.m. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| GCA | A | 1.15 | 1.03 | 1.68 | 1.22 | 1.13 | 0.82 | 0.75 | 1.29 | 1.66 | 1.70 | 1.33 | 2.11 |
| GCU | A | 2.18 | 1.87 | 1.39 | 2.17 | 2.00 | 1.83 | 2.55 | 1.42 | 1.52 | 1.28 | 1.46 | 1.68 |
| GCG | A | 0.06 | 0.14 | 0.31 | 0.12 | 0.09 | 0.44 | 0.15 | 0.33 | 0.16 | 0.20 | 0.23 | 0.05 |
| GCC | A | 0.62 | 0.96 | 0.62 | 0.49 | 0.78 | 0.91 | 0.55 | 0.96 | 0.67 | 0.83 | 0.99 | 0.16 |
| UGC | C | 0.33 | 0.43 | 0.28 | 0.41 | 0.31 | 0.82 | 0.58 | 0.63 | 0.35 | 0.50 | 0.49 | 0.36 |
| UGU | C | 1.67 | 1.57 | 1.72 | 1.59 | 1.69 | 1.18 | 1.42 | 1.38 | 1.65 | 1.50 | 1.51 | 1.64 |
| GAU | D | 1.46 | 1.40 | 1.09 | 1.53 | 1.44 | 1.22 | 1.20 | 1.44 | 1.18 | 1.32 | 1.49 | 1.68 |
| GAC | D | 0.54 | 0.60 | 0.91 | 0.47 | 0.56 | 0.78 | 0.80 | 0.56 | 0.82 | 0.69 | 0.51 | 0.33 |
| GAG | E | 0.51 | 0.58 | 0.63 | 0.41 | 0.49 | 0.55 | 0.56 | 0.71 | 0.49 | 0.47 | 0.54 | 0.29 |
| GAA | E | 1.49 | 1.42 | 1.37 | 1.59 | 1.51 | 1.45 | 1.44 | 1.29 | 1.51 | 1.53 | 1.46 | 1.71 |
| UUU | F | 1.75 | 1.66 | 1.30 | 1.80 | 1.65 | 1.50 | 1.66 | 1.59 | 1.67 | 1.49 | 1.32 | 1.49 |
| UUC | F | 0.25 | 0.34 | 0.70 | 0.20 | 0.35 | 0.50 | 0.34 | 0.41 | 0.33 | 0.51 | 0.68 | 0.52 |
| GGU | G | 1.21 | 1.48 | 1.39 | 0.87 | 1.02 | 0.71 | 0.96 | 1.27 | 1.23 | 1.22 | 1.24 | 1.77 |
| GGG | G | 1.00 | 0.66 | 0.71 | 1.01 | 0.90 | 1.87 | 1.27 | 0.82 | 0.80 | 0.98 | 0.51 | 0.19 |
| GGC | G | 0.11 | 0.41 | 0.42 | 0.21 | 0.17 | 0.38 | 0.34 | 0.45 | 0.55 | 0.20 | 0.83 | 0.02 |
| GGA | G | 1.68 | 1.45 | 1.48 | 1.92 | 1.91 | 1.03 | 1.43 | 1.46 | 1.43 | 1.60 | 1.42 | 2.02 |
| CAC | H | 0.35 | 0.84 | 1.19 | 0.32 | 0.73 | 1.11 | 0.49 | 0.86 | 0.87 | 0.78 | 0.80 | 0.70 |
| CAU | H | 1.65 | 1.17 | 0.81 | 1.68 | 1.27 | 0.89 | 1.51 | 1.14 | 1.13 | 1.23 | 1.21 | 1.30 |
| ÂUUU | I | 1.75 | 1.61 | 1.46 | 1.77 | 1.65 | 1.49 | 1.48 | 1.49 | 1.63 | 1.59 | 1.63 | 1.78 |
| AUC | 1 | 0.25 | 0.39 | 0.54 | 0.23 | 0.35 | 0.52 | 0.52 | 0.51 | 0.37 | 0.41 | 0.37 | 0.23 |
| AAA | K | 1.64 | 1.60 | 1.22 | 1.64 | 1.44 | 1.44 | 1.58 | 1.41 | 1.45 | 1.51 | 1.21 | 1.41 |
| AAG | K | 0.36 | 0.40 | 0.78 | 0.36 | 0.56 | 0.56 | 0.42 | 0.59 | 0.55 | 0.49 | 0.80 | 0.59 |
| ÜÜG | L | 0.53 | 0.70 | 1.67 | 0.31 | 0.64 | 0.53 | 0.81 | 0.86 | 1.07 | 1.00 | 0.56 | 0.54 |
| UUA | L | 3.77 | 3.50 | 1.93 | 4.04 | 3.39 | 2.54 | 2.85 | 2.62 | 2.46 | 2.65 | 2.61 | 3.91 |
| CUA | L | 0.56 | 0.66 | 1.49 | 0.64 | 1.07 | 1.23 | 0.81 | 1.17 | 1.08 | 1.28 | 1.18 | 0.83 |
| CUC | L | 0.24 | 0.28 | 0.11 | 0.08 | 0.12 | 0.38 | 0.25 | 0.18 | 0.33 | 0.16 | 0.44 | 0.06 |
| CUG | L | 0.08 | 0.07 | 0.25 | 0.04 | 0.07 | 0.31 | 0.21 | 0.26 | 0.13 | 0.23 | 0.13 | 0.05 |
| CUU | L | 0.81 | 0.79 | 0.55 | 0.89 | 0.71 | 1.02 | 1.08 | 0.90 | 0.93 | 0.69 | 1.09 | 0.61 |
| ÄÜG | M | 0.27 | 0.51 | 0.46 | 0.23 | 0.23 | 0.51 | 0.38 | 0.44 | 0.32 | 0.45 | 0.43 | 0.31 |
| AUA | M | 1.73 | 1.49 | 1.54 | 1.78 | 1.77 | 1.49 | 1.62 | 1.56 | 1.68 | 1.55 | 1.57 | 1.69 |
| AAC | N | 0.31 | 0.56 | 1.00 | 0.41 | 0.51 | 0.80 | 0.66 | 0.66 | 0.64 | 0.46 | 0.59 | 0.48 |
| AAU | N | 1.69 | 1.44 | 1.00 | 1.59 | 1.49 | 1.20 | 1.34 | 1.34 | 1.36 | 1.54 | 1.41 | 1.52 |
| CCU | P | 2.06 | 1.38 | 1.58 | 2.05 | 1.80 | 1.70 | 2.05 | 1.67 | 1.70 | 0.94 | 1.83 | 1.50 |
| CCG | P | 0.17 | 0.32 | 0.15 | 0.09 | 0.15 | 0.19 | 0.18 | 0.36 | 0.19 | 0.14 | 0.18 | 0.09 |
| CCC | P | 0.78 | 0.43 | 0.59 | 0.39 | 0.46 | 0.99 | 0.43 | 0.93 | 0.69 | 0.61 | 0.85 | 0.12 |
| CCA | $P$ | 0.99 | 1.87 | 1.68 | 1.47 | 1.59 | 1.12 | 1.34 | 1.04 | 1.43 | 2.31 | 1.14 | 2.29 |
| CAG | Q | 0.20 | 0.30 | 0.49 | 0.18 | 0.14 | 0.40 | 0.26 | 0.37 | 0.54 | 0.27 | 0.23 | 0.10 |
| CAA | Q | 1.80 | 1.71 | 1.51 | 1.82 | 1.86 | 1.60 | 1.74 | 1.63 | 1.46 | 1.73 | 1.77 | 1.91 |
| CGA | R | 2.25 | 2.59 | 1.31 | 1.78 | 1.71 | 1.39 | 1.47 | 2.26 | 1.42 | 1.93 | 2.03 | 2.47 |
| CGC | R | 0.17 | 0.31 | 0.30 | 0.44 | 0.33 | 0.77 | 0.33 | 0.26 | 0.14 | 0.21 | 0.49 | 0.07 |
| CGG | R | 0.00 | 0.31 | 0.48 | 0.07 | 0.65 | 0.77 | 0.41 | 0.45 | 0.68 | 0.83 | 0.62 | 0.15 |
| CGU | R | 1.58 | 0.78 | 1.91 | 1.70 | 1.31 | 1.08 | 1.80 | 1.03 | 1.76 | 1.03 | 0.86 | 1.31 |
| AGGC | S | 0.13 | 0.40 | 0.33 | 0.30 | 0.16 | 0.47 | 0.29 | 0.40 | 0.43 | 0.17 | 0.30 | 0.07 |
| AGU | S | 0.61 | 0.87 | 1.18 | 0.91 | 1.03 | 0.99 | 0.73 | 1.10 | 0.86 | 0.95 | 1.28 | 0.53 |
| AGG | S | 0.23 | 0.23 | 0.18 | 0.04 | 0.16 | 0.09 | 0.00 | 0.07 | 0.02 | 0.30 | 0.03 | 0.04 |
| UCU | S | 2.09 | 2.30 | 1.72 | 2.82 | 2.43 | 2.66 | 2.67 | 1.93 | 2.56 | 1.86 | 2.40 | 2.69 |
| UCG | S | 0.13 | 0.11 | 0.23 | 0.11 | 0.16 | 0.18 | 0.31 | 0.47 | 0.26 | 0.11 | 0.20 | 0.09 |
| AGA | S | 1.73 | 1.23 | 0.90 | 1.46 | 1.10 | 1.42 | 1.40 | 1.87 | 1.90 | 1.46 | 1.38 | 1.79 |
| UCC | S | 0.71 | 0.40 | 0.72 | 0.47 | 1.06 | 0.90 | 0.94 | 0.81 | 0.64 | 0.68 | 0.69 | 0.15 |
| UCA | S | 2.38 | 2.45 | 2.74 | 1.89 | 1.91 | 1.29 | 1.65 | 1.35 | 1.33 | 2.48 | 1.73 | 2.65 |
| ACA | T | 1.63 | 1.64 | 1.92 | 1.40 | 1.27 | 1.21 | 1.07 | 1.52 | 1.69 | 1.93 | 1.68 | 2.56 |
| ACU | T | 1.65 | 1.61 | 1.13 | 2.28 | 1.93 | 1.79 | 2.01 | 1.52 | 1.43 | 1.18 | 1.54 | 1.15 |
| ACC | T | 0.67 | 0.73 | 0.74 | 0.30 | 0.73 | 0.74 | 0.78 | 0.75 | 0.76 | 0.80 | 0.68 | 0.23 |
| ACG | T | 0.05 | 0.02 | 0.22 | 0.02 | 0.07 | 0.27 | 0.15 | 0.21 | 0.12 | 0.09 | 0.09 | 0.06 |
| GUU | V | 1.76 | 1.51 | 1.30 | 1.83 | 1.54 | 1.50 | 1.44 | 1.45 | 1.69 | 1.56 | 1.46 | 2.08 |
| GUG | V | 0.29 | 0.48 | 0.82 | 0.42 | 0.28 | 0.92 | 0.74 | 0.57 | 0.47 | 0.48 | 0.35 | 0.07 |
| GUA | V | 1.70 | 1.49 | 1.78 | 1.60 | 1.96 | 1.16 | 1.33 | 1.55 | 1.54 | 1.56 | 1.91 | 1.75 |
| GUC | V | 0.25 | 0.53 | 0.11 | 0.15 | 0.22 | 0.43 | 0.49 | 0.43 | 0.30 | 0.41 | 0.28 | 0.11 |
| ÜGA | W | 1.55 | 1.77 | 1.39 | 1.62 | 1.72 | 1.65 | 1.30 | 1.65 | 1.54 | 1.71 | 1.70 | 1.92 |
| UGG | W | 0.45 | 0.23 | 0.61 | 0.38 | 0.28 | 0.35 | 0.70 | 0.36 | 0.46 | 0.29 | 0.31 | 0.08 |
| UAU | Y | 1.67 | 1.54 | 1.30 | 1.54 | 1.47 | 1.12 | 1.44 | 1.35 | 1.40 | 1.48 | 1.46 | 1.62 |
| UAC | Y | 0.33 | 0.46 | 0.70 | 0.46 | 0.53 | 0.88 | 0.56 | 0.65 | 0.60 | 0.52 | 0.54 | 0.38 |

Bolded numbers represent the codon most commonly used to code the amino acid. C.f. = Campodea fragilis, C.I. $=$ C. lubbocki, J.s. $=$ Japyx solifugus, G.h. $=$ Gomphiocephalus hodgsoni, T.b. $=$ Tetrodontophora bielanensis, P.a. = Podura aquatica, O.o. = Onychurus orientalis, P.b.= Petrobius brevistylis,
N.a. $=$ Nesomachilis australica, T.g. $=$ Tricholepidion gertschi, T.d. $=$ Thermobia domestica, L.m. $=$

Locusta migratoria


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