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### **Publication Date**

2005-12-01

Peer reviewed

**LBNL-59181**

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# **The Complete Sequence of the Mitochondrial Genome of the Chambered Nautilus (Mollusca: Cephalopoda)**

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## Abstract

### Background

Mitochondria contain small genomes that are physically separate from those of nuclei. Their comparison serves as a model system for understanding the processes of genome evolution. Although complete mitochondrial genome sequences have been reported for more than 600 animals, the taxonomic sampling is highly biased toward vertebrates and arthropods, leaving much of the diversity yet uncharacterized.

### Results

The mitochondrial genome of a cephalopod mollusk, the Chambered Nautilus, is 16,258 nts in length and 59.5% A+T, both values that are typical of animal mitochondrial genomes. It contains the 37 genes that are typical for animal mtDNAs, with 15 on one DNA strand and 22 on the other. The arrangement of these genes can be derived from that of the distantly related *Katharina tunicata* (Mollusca: Polyplacophora) by a switch in position of two large blocks of genes and transpositions of four tRNA genes. There is strong skew in the distribution of nucleotides between the two strands. There are an unusual number of non-coding regions and their function, if any, is not known; however, several of these demark abrupt shifts in nucleotide skew, suggesting that they may play roles in transcription and/or replication. One of the non-coding regions contains multiple repeats of a tRNA-like sequence. Some of the tRNA genes appear to overlap on the same strand, but this could be resolved if the polycistron were cleaved at the beginning of the downstream gene, followed by polyadenylation of the product of the upstream gene to form a fully paired structure.

## **Conclusions**

*Nautilus* sp. mtDNA contains an expected gene content that has experienced few rearrangements since the evolutionary split between cephalopods and polyplacophorans. It contains an unusual number of non-coding regions, especially considering that these otherwise often are generated by the same processes that produce gene rearrangements. This appears to be yet another case where polyadenylation of mitochondrial tRNAs restores what would otherwise be an incomplete structure.

## Background

Animal mitochondrial DNA (mtDNA) is nearly always a closed circular molecule and, with a few exceptions [e.g. 1-4], contains the same 37 genes, specifying 13 proteins, two ribosomal RNAs, and 22 tRNAs [5]. Sequences of these diminutive genomes have been broadly used to address phylogenetic questions ranging from the population [6,7] to the interphylum [8-11] levels and to model many processes of genome evolution [12,13]. Although there are exceptions, most mtDNAs contain no introns and are between 14 and 17 kb. Typically there are few intergenic nucleotides except for a single large non-coding region generally thought to contain elements that regulate replication and transcription [14]. Occasionally non-coding regions have been found that contain repeated elements [15] or contain pseudogenes [12,16] or that may be remnants of duplicated regions, perhaps those that mediate gene rearrangements [12,16,17]. Gene rearrangements tend to be uncommon and to occur in a saltatory manner [see 10]. The “universal” genetic code has been modified in many animal lineages, to include the use of alternative start codons and abbreviated stop codons [18,19]. In some mtDNAs there is pronounced skew in nucleotide composition, often with one strand being rich in G and T and the other in A and C [20]. Post-transcriptional modification of nucleotides has been observed for tRNAs [21,22].

Little study has been done to date on mollusk mtDNAs compared to those of vertebrates or arthropods [23], but it is already apparent that mollusks exhibit much variation in the features of their mitochondrial genomes, including losses and gains of genes [2], atypically large amounts of duplicated or non-coding nucleotides [15,24], highly rearranged genomes [2,25], and an unusual pattern of passage termed doubly uniparental inheritance [26,27]. We further this here by

reporting and comparing the features of the mitochondrial genome of the Chambered Nautilus (Mollusca: Cephalopoda).

Nautiloids were once abundant and diverse in the Paleozoic seas, but only a handful of species remain, all in the genus *Nautilus*. They are part of the molluscan class Cephalopoda, which otherwise contains octopi, squid, and cuttlefish. They are the earliest diverging lineage of this group and are often considered to be “living fossils” since living forms seem to have changed little from their ancient ancestors. Their common name, the chambered nautilus, comes from the chambers that develop in their spiral-shaped shells as they grow, beginning with about four in the young animal and progressing to 30 in the adult, and which are filled with gas to control buoyancy. The animal lives only in the largest, outermost chamber of the shell. They are carnivorous, using their many grooved tentacles to grasp prey and pass it to their mouth, where a beak-like jaw tears it and passes it to the shredding radula, and they move about by squirting jets of water. They live throughout the Southwest Pacific Ocean, at depths as great as 610 meters, and traverse a great range, as shallow as 90 meters, apparently in search of prey.

## **Results and Discussion**

### **Gene content and organization**

The *Nautilus* sp. (Mollusca: Cephalopoda) mitochondrial genome is 16,258 bp in length (GenBank accession number NNNNNNNN) and contains the set of 37 genes most commonly found for animal mtDNAs [5]. Fifteen genes are located on one strand and 22 on the other (Fig. 1). There are several substantial non-coding regions (see below), the largest of which is 972 nts long and between *trnQ* and *trnT*. The mitochondrial gene arrangement of *Nautilus* sp. differs from that of the distantly related *Katharina tunicata* [28] (the only sampled representative of the

Polyplacophora, an early diverging class of the Mollusca) by only transpositions of four tRNA genes and the switch in position of two large blocks of genes (Fig. 2). By comparison with an outgroup, the phoronid *Phoronis architecta* [11] (and confirmed by others not shown), we can see that one of these tRNAs, *trnD*, remains in the ancestral condition in these two cephalopods shown in Fig. 2, with a transposition having occurred in the polyplacophoran, whereas all other changes are derived for *Nautilus* sp. The mtDNA of *Octopus vulgaris* [29] is nearly identical in arrangement to that of *K. tunicata*, differing otherwise only in the inversion of *trnP*, which it shares with *Nautilus* and many other mollusks.

The mtDNA sequence is available for two other cephalopod mollusks, the squids *Loligo bleekeri* [30] and *Todarodes pacificus* [29], each of which is more highly rearranged (Supplemental Table 1). There are only three gene blocks that *L. bleekeri* shares with *Nautilus* sp. mtDNA (minus symbol indicates opposite transcriptional orientation, and oriented here to correspond with Fig. 2): (1) *-trnS2*, *-cob*, *-nad6*, *-trnP*, *-nad1*; (2) *-rrnL*, *-trnV*, *-rrnS*; and (3) *trnS1*, *nad2*, *cox1*. *L. bleekeri* mtDNA also shares the two blocks *-nad4*, *-nad4L*, *trnT* and *trnD*, *atp8*, *atp6* with *K. tunicata* and *O. vulgaris*. The mtDNA of *T. pacificus* also has several blocks of genes conserved with one or more mollusks: (1) *trnN*, *trnI*, *nad3*; (2) *cox1*, *cox2*, *trnD*, *atp8*, *atp6*, *-trnF*; (3) *-trnV*, *-rrnS*; (4) *cox3*, *trnK*; (5) *trnS1*, *nad2*, *cox1*, *cox2*, *trnD*, *atp8*, *atp6*; (6) *-nad5*, *-trnH*, *-nad4*, *-nad4L*, *trnT*, *-trnS2*, *-cob*, *-nad6*, *-trnP*, *-nad1*, *-trnL2*, *-trnL1*, *-rrnL*; (7) *-trnG*, *-trnE*. However, there are six genes that appear in duplicate, accounting for the listing of some genes more than once in differing arrangements here.



## Gene initiation and termination

Mitochondrial genomes often use a variety of non-standard initiation codons [19], but *Nautilus* mtDNA has only one type of deviation; three genes (*nad3*, *nad4*, and *nad5*) start with GTG and all others use the standard ATG (Fig. 3). Seven genes have unambiguous termination codons, either TAG (*atp6*, *cox1*, *nad5*) or TAA (*atp8*, *cox3*, *nad1*, *nad2*). In four cases (*cox2*, *cob*, *nad3*, *nad4*) genes are probably abbreviated to a single T or to TA such that the excision of the adjacent, downstream tRNA from the polycistronic message leaves an mRNA that is polyadenylated to complete a TAA stop codon. However, in each of these cases, a complete stop codon is available if there is, alternatively, overlap of only one or two nucleotides with the downstream tRNA. Perhaps these act as a “backup” for cases where translation precedes message cleavage. The other two cases are more ambiguous. *nad4L* could have an abbreviated stop codon, but is inferred to overlap *nad4* by seven nucleotides to the first legitimate stop codon, since overlap of this pair has been commonly observed for other mtDNAs, where they are thought to be translated as a bicistron. *nad6* is inferred to overlap *cob* by eight nucleotides, perhaps suggesting that these are processed also as a bicistron, but could instead end on an abbreviated stop codon if there were some signal for message cleavage (i.e., other than a tRNA) that we do not recognize. Inferred in this way, all protein-encoding genes have lengths nearly identical to those of *K. tunicata* mtDNA (Supplemental Table 2).

## Base composition and codon usage

The *Nautilus* sp. mtDNA is 59.5% A+T. The strand that includes *cox1*, which we will arbitrarily designate as the plus strand for the purpose of discussion, is 33.7% A, 25.8% T, 11.9% G, and 28.5% C. This strand is strongly skewed (as calculated in [20]) away from both T

(T-skew = -013) and G (G-skew = -041) in favor of A and C. As can be seen in Table 1, this is strongly reflected in the use of synonymous codons. For example, while TTT and TTC are used with approximately equal frequency to specify phenylalanine in plus strand genes, the bias is 158 to 3 for their usage in minus strand genes. The use of G vs. A in UUR (leucine) codons is in the ratio of 16 to 89 for plus strand genes but, even though the mtDNA is A+T-rich, it is 195 to 60 for minus strand genes. Presumably the biased use of synonymous codons is driven by strand-specific mutational propensity.

There are eight non-coding regions of this mtDNA that are larger than 20 nucleotides, several of which are associated with shifts in these skew values (Fig. 4, Table 2, Supplemental Table 3). **ADD MORE HERE.**

### **Transfer RNAs**

Sequences were identified whose potential secondary structures indicate that they encode the 22 tRNAs typically found for animal mtDNA (Fig. 5). In general, these appear well paired with only a few mismatches.

There are three cases where tRNA genes appear to overlap, and these potential structures suggest how this is resolved. *trnL1* appears to overlap *trnL2* by only the former's discriminator nucleotide (A). *trnQ* appears to overlap *trnW* by two nucleotides. *trnK* appears to overlap *trnA* by four nucleotides, GGCT. These are well-paired in the potential structure of tRNA(A), but these four correspond to two G-T pairs, one mismatch, and the discriminator nucleotide of tRNA(K). It appears for each case that cleavage to form a complete downstream tRNA followed by (poly)adenylation of the upstream tRNA (as has been demonstrated for some mitochondrial tRNAs [Yokobori and Paabo 1995]) would yield fully formed, well-paired structures for all. This

is illustrated in Fig. 5 by lower case, parenthetical letter “a” appended to the genome-encoded nucleotide to indicate likely nucleotides in the actual transcript.

Usually T is in the first anticodon position for tRNAs that recognize either four-fold degenerate codon families or to specifically recognize NNR codons; G is usually in this position only to specifically recognize NNY codons. (Due to the convention of always drawing RNAs from 5' to 3' in orientation, the first nucleotide listed for an anticodon pairs with the last nucleotide of a codon.) All but two of the *Nautilus* mitochondrial tRNAs follow this pattern. One exception is tRNA(M), which has the anticodon CAT (to recognize both ATA and ATA), as is almost universally the case for all animal mitochondrial systems. However, it is unusual that the tRNA(S-AGN) has a GCT anticodon, since this requires the G to pair with all four nucleotides in the wobble position of AGN codons. It is clear the AGA and AGG codons are being used and are not stop codons (as is the case in vertebrate mtDNAs), since they appear in the reading frames of protein encoding genes 117 times. It is possible that some anticodon nucleotides are post-transcriptionally modified to alter their base pairing properties.

### **Non-coding regions**

The mtDNA of *Nautilus* sp. has 1,416 nucleotides that are not assigned to genes. This is not an unusually large number, but it is atypical that they are distributed among so many regions of the genome (Table 2 and Supplemental Table 3). It is particularly unusual to find this in a mitochondrial genome that has not undergone significant rearrangements, since intergenic non-coding regions appear in some cases to be vestiges of pseudogenes generated by the gene duplication-random loss process of rearrangement [Mueller and Boore 2005; Boore 2000; Macey et al 2004; Arendt and Smith 1996].

In the largest non-coding region, between *trnQ* and *trnT*, and beginning adjacent to a (CA)<sub>13</sub> run (see below), there are six repeats of a 62 nucleotide element followed by a partial repeat of 39 nucleotides. Within this are five overlapping regions that have potential for forming tRNA-like structures (Fig. 5). The anticodon portion of these structures is AGT, which would pair with codon ACT (or perhaps ACN) to specify threonine. However, having A in this anticodon position would be very unusual and there is little sequence similarity to *trnT* (or any other tRNA).

Tandem repeats of CA are common, with (CA)<sub>3</sub> in each of the intergenic regions of *trnA-trnR* and *trnG-atp6* and an especially noteworthy (CA)<sub>13</sub> in the region between *trnQ* and *trnT*. Homopolymer runs of T<sub>10</sub>, nine C<sub>9</sub>, and A<sub>20</sub> are in the regions *trnQ-trnT*, *trnG-atp6*, and *trnE-cox3*, respectively. Non-coding, non-functional portions of mtDNA are generally eliminated rapidly (Ashley et al. 1989), presumably due to selection for small size at the point of entry into the primordial germ plasm during embryogenesis (Mignotte et al. 1987), but whether these or any particular motif plays any role in regulating replication of transcription awaits experimentation

## **Conclusions**

**ELABORATE ON THESE ASPECTS FROM THE ABSTRACT: *Nautilus* sp.**  
**mtDNA contains an expected gene content that has experienced few rearrangements since the evolutionary split between cephalopods and polyplacophorans. It contains an unusual number of non-coding regions, especially considering that these otherwise often are generated by the same processes that produce gene rearrangements. This appears to be yet**

**another case where polyadenylation of mitochondrial tRNAs restores what would otherwise be an incomplete structure.**

## **Methods**

### **Molecular techniques**

Mitochondrial DNA was isolated from approximately 1 g of *Nautilus* sp. testis tissue (gift of Wesley Brown) that had been stored at -80° C by first grinding in liquid nitrogen using a mortar and pestle. This powder was dissolved in 14 ml of homogenization buffer (210 mM mannitol, 70 mM sucrose, 50 mM Tris HCl-pH 7.5, 3 mM CaCl<sub>2</sub>) and processed using a TissueMizer T-25 (Tekmar) with three strokes of five seconds each. Membranes were lysed by adding 1/10 volume of 20% SDS and incubating for 20 min at RT. A 1/6 volume of saturated CsCl in water was added and this mixture incubated on ice for 15 min. Debris was pelleted at 17,000 X G for 10 minutes at 4° C. Propidium iodide was added to the collected supernatant to a final concentration of 500 µg/ml and the CsCl concentration was adjusted to a density of 1.57 g/ml. Nuclear and mitochondrial DNA were separated by density gradient centrifugation in a VTi65 rotor at 55,000 X G for 15 hours at 21° C. Although no mitochondrial band was visible in the gradient, the region from about 2-10 mm below the nuclear band was collected using a needle. This was then extracted multiple times with water-saturated butanol to remove the propidium iodide and dialyzed against TE for 24 hours with three buffer changes to remove the CsCl, leaving the sample in a 100 µl volume.

This product was used in PCR as in [Boore and Brown 2000] to amplify first several short fragments of *cox1*, *rrnL*, and *cob* using primers found in [Folmer et al 1994, Palumbi 1996, Boore and Brown 2000]. The fragment of *cox1* was cloned into pBluescript (Stratagene) that

had been digested with *EcoRV*, T-tailed using Taq polymerase, and gel purified using GeneClean (QBiogene). A successful recombinant clone was selected and DNA prepared using standard techniques. The other fragments were purified by three serial passages through an Ultrafree (NMWL 30,000) spin column (Millipore) and sequenced directly. The sequences of these fragments were determined using an ABI377 automated DNA sequencer with BigDye chemistry (Applied Biosystems) according to supplier's instructions.

Primers were designed to known sequences for use in long PCR [Barnes 1994] with rTth-XL polymerase (Applied Biosystems) according to supplier's instructions, sometimes combined with primers to conserved mtDNA regions. Generously overlapping fragments were amplified from *cox1-nad1* (using conserved *nad1* primer CCTGATACTAATTCAGATTCTCCTTC), *nad1-cob*, *cob-rrnL*, and *rrnL-cox1* (using conserved primer 16SARL [Palumbi 1996]), jointly comprising the entire mtDNA. Because there was no information available for the gene arrangement, many combinations of primers were tried, but only these reactions gave bright, singular bands during electrophoretic analysis. Sequence was determined for each as above, then by primer walking through each fragment. To ensure accuracy, all sequence was determined from both strands. Sequencing reads were assembled manually and quality verified by eye using Sequence Navigator (Applied Biosystems).

### **Gene annotation and analysis**

Genes encoding rRNAs and proteins were identified by matching nucleotide or inferred amino acid sequences to those of *Katharina tunicata* mtDNA [Boore and Brown 1994]. Since it is not possible to precisely determine the ends of rRNA genes by sequence data alone, they were assumed to extend to the boundaries of flanking genes. Each protein gene was inferred to begin

at an eligible initiation codon nearest to the beginning of its alignment with homologous genes that does not cause overlap with the preceding gene. In five cases, an abbreviated stop codon was inferred where cleavage of a downstream tRNA from the transcript would leave a partial codon of T or TA, such that subsequent mRNA polyadenylation could generate a TAA stop codon; however, in each of these cases, if the reading frame extended through the first legitimate stop codon there would be only a short overlap with the downstream gene. Genes for tRNAs were identified generically by their ability to fold into a cloverleaf structure and specifically by anticodon sequence. Subsequent sequence analyses were performed using MacVector (Accelrys).

## **Abbreviations**

*cox1*, *cox2*, *cox3*, cytochrome oxidase subunit I, II, and III protein genes; *cob*, cytochrome b gene; *atp6*, *atp8*, ATP synthase subunit 6 and 8 genes; *nad1*, *nad2*, *nad3*, *nad4*, *nad4L*, *nad5*, *nad6*, NADH dehydrogenase subunit 1-6, 4L genes; *trnA*, *trnC*, *trnD*, *trnE*, *trnF*, *trnG*, *trnH*, *trnI*, *trnK*, *trnL1*, *trnL2*, *trnM*, *trnN*, *trnP*, *trnQ*, *trnR*, *trnS1*, *trnS2*, *trnT*, *trnV*, *trnW*, *trnY*, transfer RNA genes designated by the one-letter code for the specified amino acid, with numerals differentiating cases where there are two tRNAs for the same amino acid

## **Acknowledgments**

I am grateful to Wesley Brown for tissue of *Nautilus* sp. This work was supported by DEB-9807100 from the National Science Foundation and by contract DE-AC02-05CH11231 between the US Department of Energy, Office of Biological and Environmental Research, and the University of California, Lawrence Berkeley National Laboratory.



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## Figures

### Figure 1 – Mitochondrial gene map of the cephalopod mollusk *Nautilus* sp.

Genes for proteins and rRNAs are shown with standard abbreviations with an arrow indicating the direction of transcription. Genes for tRNAs are designated by a single letter for the corresponding amino acid, with the two leucine and two serine tRNAs differentiated by numeral (S1, S2, L1, and L2 recognizing codons AGN, UCN, CUN, and UUR, respectively). tRNA genes shown outside the circle are transcribed clockwise and those inside the circle are transcribed counter-clockwise. The largest non-coding region is designated “nc”.

### Figure 2 – Reconstruction of mitochondrial genome rearrangements for *Nautilus* sp.

At the top is the nearly complete gene arrangement for *Phoronis architecta* [Helfenbein and Boore 2004], a presumed outgroup to the mollusks, shown to polarize two of the cephalopod rearrangements: Having *trnP* in opposite orientation to *nad6* and *nad1* is the ancestral condition, as is having *trnD* between *cox2* and *atp8*. The only two differences between the chiton *Katharina tunicata* [Boore and Brown 1994] and the octopus is the inversion of *trnP* in the octopus and the transposition of *trnD* in the chiton. (No attempt is being made here to reconstruct all of the rearrangements between the phoronid and the chiton.) The arrangement found in the Chambered Nautilus, then, can be reconstructed by the additional switch in order of two large blocks of genes plus transpositions of *trnF* and *trnT*; whether these changes were sequential or simultaneous is undetermined. Genes are not drawn to scale and are abbreviated as in Fig. 1 except that underlining signifies right-to-left transcriptional orientation. All genomes are circular and only graphically linearized at an arbitrarily chosen point.

### **Figure 3 – Greatly abbreviated sequence of *Nautilus* sp. mtDNA**

To save space the middle portions of many genes are replaced by a numeral indicating the number of omitted nucleotides. Gene orientation is specified by a dart (>). Stop codons are shown by asterisks whether complete or abbreviated, with a plus symbol indicating an alternative that overlaps the downstream gene. Down-facing arrows mark repeats found in the largest non-coding region. When not conforming to the genetic code, the presumed initiator methionine (M) is in parentheses.

### **Figure 4 – Plot of A+C and G+T composition along mtDNAs of *Nautilus* sp. and *Katharina tunicata* using a sliding window of 100 nucleotides**

Numbering of nucleotides begins as the arbitrarily chosen *cox1* (as in Fig. 3 for *Nautilus* sp.). The scaled gene maps are also presented. tRNA genes are pictured but not labeled. Underlining indicates right-to-left transcriptional orientation. Numerals label each non-coding region larger than 20 nts, which are then projected onto the plot by gray highlighting. Several of these correspond to positions where there is a shift in nucleotide bias. Asterisks beside two of the numerals for *K. tunicata* indicate some ambiguity where these may instead be supernumerary tRNA genes (Boore and Brown 1994). Red bars show the major transposition between the two genomes (see Fig. 2).

### **Figure 5 – *Nautilus* sp. mitochondrial tRNA gene sequences folded into typical cloverleaf structures**

Lower case “a” in parentheses indicates likely replacements by (poly)adenylation after transcript cleavage at the downstream tRNA (see text for explanation). Structural features are shown on

tRNA(V). Also shown is the secondary structure possible for the repeats in the large non-coding region that appear to be pseudogenes.



## Tables

**Table 1 – Codon usage for the 13 mitochondrial proteins of *Nautilus* sp.**

The total number of codons is 3711. Stop codons were not included in this count. Here the plus strand refers arbitrarily to the one that contains *coxI*.

AA	Codon	All protein encoding genes	Plus strand only	Minus strand only	AA	Codon	All protein encoding genes	Plus strand only	Minus strand only		
F	TTT	61 %	225	67	158	I	ATT	38 %	141	83	58
	TTC	18 %	65	62	3		ATC	15 %	57	50	7
L	TTA	40 %	149	89	60	M	ATA	19 %	71	49	22
	TTG	57 %	211	16	195		ATG	25 %	92	21	71
S	TCT	22 %	83	35	48	T	ACT	11 %	41	24	17
	TCC	13 %	50	49	1		ACC	12 %	43	39	4
	TCA	15 %	57	49	8		ACA	12 %	44	40	4
	TCG	04 %	16	3	13		ACG	04 %	14	4	10
Y	TAT	25 %	92	14	78	N	AAT	14 %	53	19	34
	TAC	13 %	49	41	8		AAC	12 %	45	40	5
*	TAA	00 %	0	0	0	K	AAA	10 %	36	24	12
*	TAG	00 %	0	0	0		AAG	11 %	42	10	32
C	TGT	18 %	66	6	60	S	AGT	11 %	39	8	31
	TGC	01 %	5	3	2		AGC	05 %	17	12	5
W	TGA	14 %	52	41	11		AGA	12 %	44	24	20

	TGG	17 %	64	16	48		AGG	20 %	73	4	69
L	CTT	22 %	80	53	27	V	GTT	45 %	168	37	131
	CTC	13 %	48	47	1		GTC	08 %	28	26	2
	CTA	21 %	77	72	5		GTA	16 %	61	36	25
	CTG	08 %	28	13	15		GTG	39 %	145	19	126
P	CCT	14 %	53	22	31	A	GCT	24 %	90	31	59
	CCC	08 %	31	26	5		GCC	09 %	35	29	6
	CCA	09 %	33	31	2		GCA	12 %	45	35	10
	CCG	05 %	19	5	14		GCG	06 %	23	0	23
H	CAT	12 %	44	24	20	D	GAT	12 %	45	18	27
	CAC	10 %	36	29	7		GAC	07 %	25	21	4
Q	CAA	12 %	43	34	9	E	GAA	09 %	35	27	8
	CAG	05 %	20	5	15		GAG	15 %	56	14	42
R	CGT	05 %	17	5	12	G	GGT	17 %	62	19	43
	CGC	02 %	9	8	1		GGC	08 %	29	19	10
	CGA	04 %	16	14	2		GGA	28 %	104	54	50
	CGG	05 %	17	4	13		GGG	41 %	153	19	134

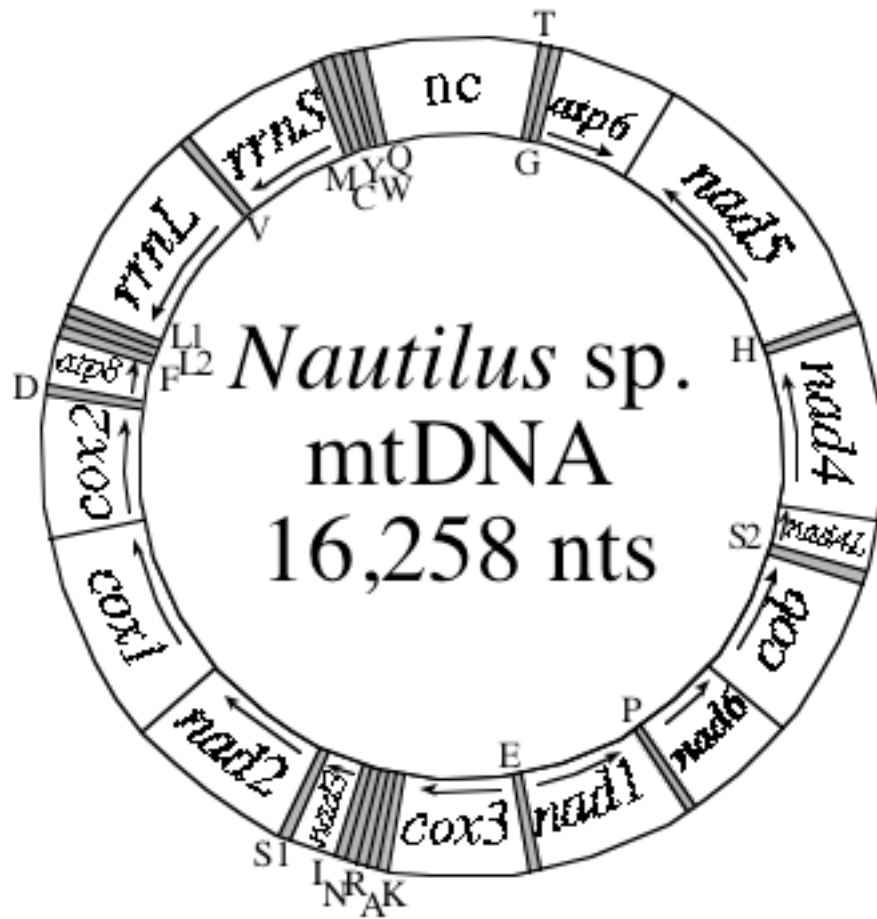
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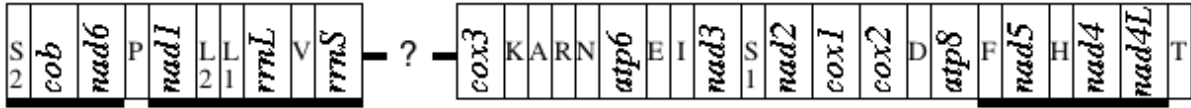
**Table 2 – Number of nucleotides at gene boundaries**

Negative numbers refer to overlapping nucleotides.

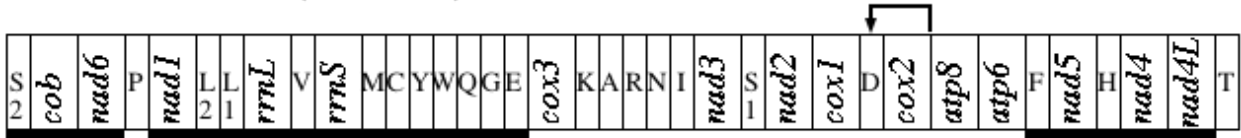
<b>Boundary</b>	<b>nts</b>	<b>Boundary</b>	<b>nts</b>
<i>cox1 – cox2</i>	2	<i>trnH – nad4</i>	0
<i>cox2 – trnD</i>	0	<i>nad4 – nad4L</i>	– 7
<i>trnD – atp8</i>	1	<i>nad4L – trnS2</i>	14
<i>atp8 – trnF</i>	20	<i>trnS2 – cob</i>	0
<i>trnF – trnL2</i>	0	<i>cob – nad6</i>	0
<i>trnL2 – trnL1</i>	– 1	<i>nad6 – trnP</i>	2
<i>trnL1 – rrnL</i>	0	<i>trnP – nad1</i>	5
<i>rrnL – trnV</i>	0	<i>nad1 – trnE</i>	26
<i>trnV – rrnS</i>	0	<i>trnE – cox3</i>	102
<i>rrnS – trnM</i>	0	<i>cox3 – trnK</i>	0
<i>trnM – trnC</i>	3	<i>trnK – trnA</i>	– 4
<i>trnC – trnY</i>	0	<i>trnA – trnR</i>	14
<i>trnY – trnW</i>	0	<i>trnR – trnN</i>	39
<i>trnW – trnQ</i>	– 2	<i>trnN – trnI</i>	4
<i>trnQ – trnT</i>	972	<i>trnI – nad3</i>	0
<i>trnT – trnG</i>	90	<i>nad3 – trnS1</i>	0
<i>trnG – atp6</i>	97	<i>trnS1 – nad2</i>	0
<i>atp6 – nad5</i>	23	<i>nad2 – cox1</i>	2
<i>nad5 – trnH</i>	0		

Keywords: Chambered nautilus, mollusk, mitochondria, mtDNA, evolution, genome

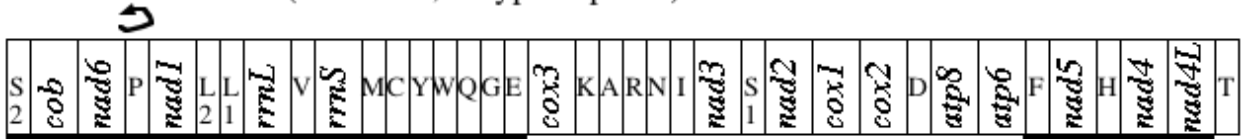




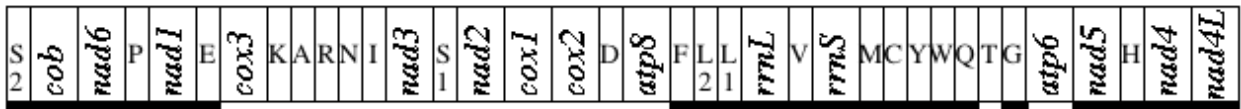
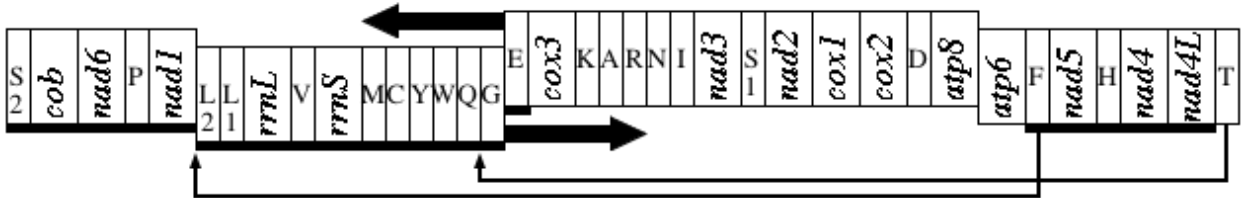
*Phoronis architecta* (Phoronida)



*Katharina tunicata* (Mollusca, Polyplacophora)



*Octopus vulgaris* (Mollusca, Cephalopoda) (and common ancestor with *Nautilus*)



*Nautilus* sp. (Mollusca, Cephalopoda)

10 20 1520 1530 1540 1550 2200  
ATGCGATGAGTATTTTCTACA-/1489/-AAACAGGAGCTCTTACATTTAGAAAATGGCCTTATGAGGACAAATAA-/632/-TCTTATTGTGGTTATC  
M R W V F S T E T G A L T L \*\*\* M A L W G Q I K F L L W L S  
\_\_\_\_\_  
cox1 > \_\_\_\_\_ cox2  
2210 2220 2230 2240 2250 2260 2270 2280 2290 2300  
AAAAAACATTAAAAATGAGTTATGTATTAACTAGGACCGTCACTCCCTAAGTCATCACAGAAGTGATATTTTTTAGATGCCTCAACTATCACCCCTCA  
K N I \*\*\* M P Q L S P L  
\_\_\_\_\_  
cox2 > \_\_\_\_\_ trnD > \_\_\_\_\_ atp8  
2430 2440 2450 2460 2470 2480 2490 2500 2510  
-/114/-CTCCCCTATAAGTGATGATAACCTAATCTACTATTTCATTATATACATATTAGTTTACATCAACCTCCTCAGCACCTCAACGCTGCGCTCTT  
P P H Y K W W \*\*\* < \_\_\_\_\_ trnF  
\_\_\_\_\_  
atp8 > \_\_\_\_\_  
2520 2530 2540 2550 2560 2570 2580 2590 2600 2610  
ATAAGCTAAATAAGTTATTAAGAAAGCTCACTACTTATTCCTAGAGCTTAAATCTATCGTACTATTTACCACCTAAATTTGGGTAGGTTAACTTAATC  
trnF < \_\_\_\_\_ trnL1(tag)  
\_\_\_\_\_  
trnL2(taa)  
2610 2620 2630 2640 2650 2660 2670 2680 2690 2700  
GGTTAACTTAATCCTATCCATCGATCCATAATCGATTACACTCATCTGCCAACCCAAATCAAGAACTTAAATTTAAATTTAAACACACATTAACTTCT  
trnL1(tag) < \_\_\_\_\_ trnL  
\_\_\_\_\_  
3990 4000 4010 4020 4030 4040 4050 4060 4070  
-/1280/-TATTCACCTACATAAGACTACCCGCTGTGAAAAGAGTAATATTACTATTAAACCAAGCGTAAAAAGGTCACATTTACTTTATGTTACTTCTCA  
rrnL < \_\_\_\_\_ trnV  
\_\_\_\_\_  
4080 4090 4100 4950 4960 4970 4980 4990 5000 5010  
CTTCAGAAGCAGGTTCCCTACCTCTA-/845/-CAACCAGAACCAAAATTTAGCAAGAAATGAAACAAATTTATTTTGGGGTATGAACCCACTAGC  
rrnS >> \_\_\_\_\_ trnM  
\_\_\_\_\_  
5020 5030 5040 5050 5060 5070 5080 5090 5100 5110  
TTACATTAGCTTATCTTACTACTAAGCTACACCCTAAGTACCTATGAACCTTCAATTCATTGTTCTATTCAAACCTCCTAAGCCTTATGAGAGGCTTA  
trnM < \_\_\_\_\_ trnC < \_\_\_\_\_ trnY  
\_\_\_\_\_  
5120 5130 5140 5150 5160 5170 5180 5190 5200 5210  
TACACCCATTAATGAATCTACAATTCAACACCTAAAAATCAGCCACCTCACACAAGACCTACGCTTTCGACATATCAAGTTTGAAGACTAATAGTT  
trnY < \_\_\_\_\_ trnW  
\_\_\_\_\_  
5220 5230 5240 5250 5260 5270 5280 5290 5300 5310  
TACATTAACCTAAGACCTTGTAGGAGGGTCCGAACCTCCTTAAAAACCCAAATCTTCTCGTGACTCCACACCCACCCACATAAATCAACACCTG  
trnW < \_\_\_\_\_ trnQ  
\_\_\_\_\_  
5320 5330 5340 5350 5360 5370 5380 5390 5400 5410  
CTCTATTAACAAGTAAAGATGACCCCAACAACCTAACAACTTACACAAAGTTTACGTTCTACCTTGACACCCACACTTTTAAACATATACCTAAACATGG  
5420 5430 5440 5450 5460 5470 5480 5490 ↓ 5500 5510  
TACCAACCGATAATCACCTTATACCCCTTACTTCCCCACACACCTAACACACACACACACACACACAGAAAGTTAAAGTACTAACCCGGTAACTA  
5520 5530 5540 5550 ↓5560 5570 5580 5590 5600 5610  
CTCTATACACTGTTTACTACTACTAATACATATACTAGGTTAGGTTAAGCTTAACTCACTCTATACACTGTTTACTCATACTACTACTACTACTA  
↓ 5630 5640 5650 5660 5670 5680 5690 5700 5710  
GGTTAGGTTACTAACCGGTAATCACTCTATACACTGTTTACTACTACTACTAATATACTAGGTTAGGTTACTAACCGGTAATCACTCTATACACTGTT  
5720 5730 5740 ↓ 5750 5760 5770 5780 5790 5800 ↓ 5810  
TACTCATACTACTAATACATATACTAGGTTAGGTTACTAACCGGTAATCACTCTATACACTGTTTACTCATACTACTAATACATATACTAGGTTAGGTTACTAA  
5820 5830 5840 5850 5860 ↓5870 5880 5890 5900 5910  
CCGGTAATCACCTTATACACTGTTTACTCATACTAATACATATACTAGGTTAGGTTACTAACCGGTAATCACTCTACACACTGTTTACTTATCTTAT  
5920 5930 5940 5950 5960 5970 5980 5990 6000 6010  
ACTCATATGGACATAATCTATACATCTTGTCTATACATATGTTCCACCTATATACTGTCTATATACCCTCTATACACCTTCTTTCATTCATTC  
6020 6030 6040 6050 6060 6070 6080 6090 6100 6110  
ATTCAATCTATATCTCTCTTATTCTATATCTTATCTCTATCTTCCATTTCTCGCATCTATATACATCTAGCCCAATGTGGGCTATGCGCGAAAGTT  
6120 6130 6140 6150 6160 6170 6180 6190 6200 6210  
GTTTTTATAACTTTTTTCATAGAAAATCGGCCCTTTTTTTTTTCAGTGCCTTTTTTGAACCTGTAATGCAATCACCTCAAAAACAGGCTAAATAAAAATTTT  
6220 6230 6240 6250 6260 6270 6280 6290 6300 6310  
ATAAACATTACCCTATGTGGTCAAATCCCAATTTTAAAGGAGTTCCCGTAGCAGCCTGGAAGCTTGTTTTAAGTAGCGCCTTGTAACCGGAAGA  
trnT  
\_\_\_\_\_  
6320 6330 6340 6350 6360 6370 6380 6390 6400 6410  
TTGTGATACTAAATCTCTCAGGCGAGTAAATTTTTTCGTAATTTCCATCATCTGTTTCGCTTAAACGCACCTATTTTAAAGTCATTCTCTAGATATTTGGG  
trnT > \_\_\_\_\_  
6420 6430 6440 6450 6460 6470 6480 6490 6500 6510  
CCCCCATACCCCTTATCTACTCTAAACGATTTTAGACCTAATGCTTGGAAGGCACCTATACTTATATATAAAAAGACATCCTCACCCACCTGGAT  
\_\_\_\_\_  
trnG  
6520 6530 6540 6550 6560 6570 6580 6590 6600 6610  
TCGCTGTGGGCTTATCTTTGTCCCCCCCTTCTTCCAGCAGCTCAATAACAAAACCTACAAAAACCCCTCCACACATATGCTATCTGACATCTTCTCA  
M L S D I F S  
\_\_\_\_\_  
atp6  
7270 7280 7290 7300 7310 7320 7330 9000  
-/650/-TATTCAGACGATCAGCTAATAGATAACCTTACCTTTAAACAATTTCAAATAAAAACAACCAACCCCTAC-/1653/-TACTAGCCTTC  
Y S D D H A N \*\*\* \*\*\* I F V V G V G V S A K  
\_\_\_\_\_  
atp6 > \_\_\_\_\_ nad5  
\_\_\_\_\_  
9010 9020 9030 9040 9050 9060 9070 9080 9090 9100  
AAATAAACTTCAAATCAACAACACTATGAAATGGTTACCCCTTCGCAACACCACTTGACATTTTACACATAAATAATCCACTAATACCACCA  
L Y F K L D V F (M) < \_\_\_\_\_ trnH >>> Y W W  
\_\_\_\_\_  
nad5 < \_\_\_\_\_ nad4  
9110 10420 10430 10440 10450 10460 10710 10720  
ATACAGTCCC-/1300/-CCCCTAACGCAAAAAACAACCTAACACTTATACAACTCAAAACCCCTTA-/240/-AAACCAGCGAATATAACAAAACCAT  
Y L G G L A F V C G L (M)  
\_\_\_\_\_  
nad4  
\*\*\* C K Y L S L G S V V L S Y L L V M  
\_\_\_\_\_  
nad4L

10730 10740 10750 10760 10770 10780 10790 10800 10810 10820  
 TTCCTAATATGCTAAACCATTAAGCGACTCGAAGACCTCCACATCTGCCTTCAAACAACCCCTAACCTTTAGTAATGGCTACCCCAACAAAACAACACT  
 < \_\_\_\_\_ *trnS2(tga)* \_\_\_\_\_ > \*\*\* G L L V V S  
 < \_\_\_\_\_ *cob* \_\_\_\_\_ >

10830 11920 11930 11940 11950 119600 12430 12440  
 AATATACCC-/1077/-ATGCTCTTTTCGAATAGACCTAAGCATAATACATACATAAAAGCCGAGAGG-/456/-ACAAAACACCACCTATTAACTCTCAT  
 I Y G H S K R I S S L M Y M S F P R L P C F V V M L S M  
 < \_\_\_\_\_ *cob* \_\_\_\_\_ > < \_\_\_\_\_ *nad6* \_\_\_\_\_ >

12450 12460 12470 12480 12490 12500 12510 12520 12530 12540  
 ATTGAGATAAGGCCTTCCTCAATCACTAATTTCCAATATAAAATTTTACATAAACTACCTTCTGCACCCCTAGCCCACTATAACCCCTACCCCT  
 < \_\_\_\_\_ *trnP* \_\_\_\_\_ > \*\*\* G W G M V S V S  
 < \_\_\_\_\_ *nad1* \_\_\_\_\_ >

13430 13440 13450 13460 13470 13480 13490 13500 13510  
 -/877/-GTAACCAACACACAAAACCTAACCATTACGCTACCCCAACAAAACCTCTAACCTAGGTGTACATCACACTCCCTTAAACTGAAA  
 T V L V C L V S V M < \_\_\_\_\_ *trnE* \_\_\_\_\_ >

13520 13530 13540 13550 13560 13570 13580 13590 13600 13610  
 CTTTAACGTGCAACCTATACACCCTAAGCCCAATCCACCCCATAGCTTTCAACAGCTAATTTCTCCACCCCAAAAAAAAAAAAAAAAAAAAA  
 < \_\_\_\_\_ *trnE* \_\_\_\_\_ >

13620 13630 13640 13650 13660 13670 14410 14420  
 TAACAAAACCTATAAATCTATAAATTTCTCAATGATCCGAAATCCCTTTTCAATTTAGTGA-/721/-TACACATGTATATTTGATGAGGATCCTAA  
 M I R N P F H L V E Y T C V Y W W G S \*\*\*  
 < \_\_\_\_\_ *cox3* \_\_\_\_\_ >

14440 14450 14460 14470 14480 14490 14500 14510 14520  
 TTTAAGGTAACCTAATTTAGTGTAACTTTTAAATTTAAAAATGGTCGTACACCCCTTGGCTTTATACCTTAAGAGCTAGAAGCCCTGATTTGCATT  
 < \_\_\_\_\_ *trnK* \_\_\_\_\_ > < \_\_\_\_\_ *trnA* \_\_\_\_\_ >

14540 14550 14560 14570 14580 14590 14600 14610 14620  
 TAGGTAGTAGGATCACCCCTTAAAGCCCTTTACACACAAGGAGAAATAAAGTGAAGTATCACATGCGGTTTCGGCCCGCAAGTTGGAAACCTATTCTCTAA  
 < \_\_\_\_\_ *trnA* \_\_\_\_\_ > < \_\_\_\_\_ *trnR* \_\_\_\_\_ >

14640 14650 14660 14670 14680 14690 14700 14710 14720  
 TTCCTTTAGCCATTCCTTTACTTCACCTCGCTTCTCCTCCCGAAGAGAGCTAATAATAGCATTAAATTTGTAATTAATAAATAGTAATGCATTTT  
 < \_\_\_\_\_ *trnN* \_\_\_\_\_ >

14740 14750 14760 14770 14780 14790 14800 14810 14820  
 ACCATTCCGGCTTGATATTGAGCCGGAATAACGGATTACATTGATGTTGTAATCACGGACATATTATGTACCCAATATCCGTGTCAATATTAATAGTTG  
 < \_\_\_\_\_ *trnN* \_\_\_\_\_ > (M) S M L M V  
 < \_\_\_\_\_ *trnI* \_\_\_\_\_ > < \_\_\_\_\_ *nad3* \_\_\_\_\_ >

14840 15130 15140 15150 15160 15170 15180 15190 15200  
 TTCCTCTATCCT-/280/-CTCCATGAATGGTCTCAAGGTCCTTGAGTGAGTCTCCTAAGAATGATGAGAATGAAGTGGGCTGCTAACCTTACTT  
 V S S I L L H E W S Q G S L E W V S \*\*\* < \_\_\_\_\_ *trnS1(gct)* \_\_\_\_\_ >  
 < \_\_\_\_\_ *nad3* \_\_\_\_\_ >

15210 15220 15230 15240 15250 15260 16230 16240 16250  
 CGGATGGTTCAAAACCTCCCTTCTTTATGTATACTAAAAGATTCCCAATTTACGTTCCCTA-/960/-AGGTATTGCTATTCTTACTTTTGGGTCTAATT  
 < \_\_\_\_\_ *trnS1(gct)* \_\_\_\_\_ > M Y T K S F P F T F L G I A I L T F V V \*\*\*  
 < \_\_\_\_\_ *nad2* \_\_\_\_\_ >



