

The Mitochondrial Genome of the Sipunculid *Phascolopsis gouldii* Supports Its Association with Annelida Rather than Mollusca

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We determined the sequence of about half (7,470 nt) of the mitochondrial genome of the sipunculid *Phascolopsis gouldii*, the first representative of this phylum to be so studied. All of the 19 identified genes are transcribed from the same DNA strand. The arrangement of these genes is remarkably similar to that of the oligochaete annelid *Lumbricus terrestris*. Comparison of both the inferred amino acid sequences and the gene arrangements of a variety of diverse metazoan taxa reveals that the phylum Sipuncula is more closely related to Annelida than to Mollusca. This requires reinterpretation of the homology of several embryological features and of patterns of animal body plan evolution.

Introduction

Complete mitochondrial genome sequences are available for 56 invertebrate animals (Boore 1999; from Genomic Diversity link at <http://www.jgi.doe.gov>). With a few notable exceptions, animal mitochondrial DNAs (mtDNAs) are circular molecules, about 16 kb in size, containing 37 genes: 13 for proteins of electron transport (cytochrome oxidase subunits I–III [*cox1–cox3*], cytochrome *b* [*cob*], NADH dehydrogenase subunits 1–6 and 4L [*nad1–nad6* and *nad4L*], ATP synthase subunits 6 and 8 [*atp6* and *atp8*]), 2 for rRNAs (large- and small-subunit rRNAs [*rrnL* and *rrnS*, respectively]), and 22 for tRNAs (abbreviated *trnX*, where “X” is the one-letter code for the corresponding amino acid, with anticodon where necessary for uniquely specifying identity). In a few lineages, the arrangements of these genes vary greatly, such as in bivalve (e.g., Hoffmann, Boore, and Brown 1992) and gastropod (e.g., Yamazaki et al. 1997) mollusks. However, in other lineages, these arrangements have remained unchanged for very long periods. For example, the arrangements of the 37 mitochondrial genes of the horseshoe crab (Staton, Daehler, and Brown 1997; Lavrov, Boore, and Brown 2000) and of the fruit fly (Clary and Wolstenholme 1985) differ by the location of only one tRNA gene.

Comparisons of mitochondrial gene arrangements have been very effective for reconstructing evolutionary relationships (Smith et al. 1993; Boore et al. 1995; Boore and Brown 1998, 2000; Boore, Lavrov, and Brown 1998; Downton 1999; Stechmann and Schlegel 1999; Kurabayashi and Ueshima 2000). Despite the occasional finding of mtDNAs with “scrambled” gene arrangements, the data set remains nearly free of homoplasy (Boore 1999, 2000). Since the rate of rearrangement varies among lineages, these gene order characters may provide resolution at differing taxonomic levels.

Key words: *Phascolopsis*, mitochondria, evolution, phylogeny, genome, Annelida.

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Within rapidly changing lineages, the signal may be best at lower levels, with anciently shared arrangements being eroded. Conversely, in more slowly evolving lineages the more ancient signal may be retained, but without rearrangements to resolve more recent branching. The merits of using gene rearrangements for recovering phylogenetic patterns lie not in their infrequency, as has been suggested (e.g., Saccone et al. 1999; Le et al. 2000), but in their complexity (i.e., there are a very large number of potential arrangements) and their near irreversibility (as judged by the infrequency of identified homoplasy). Thus, the ability to resolve any particular phylogenetic relationship may depend on the rate of gene rearrangement, but the confidence in a clade that is defined by a rearrangement, as analyzed by phylogenetic methods, does not.

In addition to providing gene order characters for inferring phylogeny, this data set also allows comparison of sequences as an independent estimator of evolutionary relationships. Furthermore, mitochondrial genome comparisons serve as a model system for genome evolution, where we can examine such factors as the following: What determines whether genes are all encoded by the same DNA strand versus being distributed between the two strands? How does mutation bias influence A+T-richness, dinucleotide frequency, codon usage, strand-skew between purines and pyrimidines, and amino acid composition of proteins? How do the structures of tRNAs evolve?

We describe here the partial mitochondrial genome of *Phascolopsis gouldii*, the first representative of the phylum Sipuncula to be so examined. The placement of this phylum within metazoan evolution has varied since its recognition as a taxon. Lamarck (1816) confused sipunculans with sea cucumbers (Holothuroidea), and they were later considered to be a derived group of annelids (Delle Chiaje 1823). In his reorganization of metazoans, de Quatrefages (1847) created the Gephyrea, or “bridge group,” which contained sipunculans, echiurans, sternapsid annelids, and priapulans. With their lack of segmentation and simple internal anatomies, de Quatrefages envisioned these groups as the transitional forms from “lower” to “higher” metazoan archetypes. Sipunculans were later elevated to the phy-

lum level (Sedgwick 1898) and associated with other spiralian either as a sister taxon to Annelida on the basis of biochemical properties (Florkin 1976; Henry 1987) or as sister to mollusks based on similarities in development (Scheltema 1993, 1996).

The developmental lynchpin that supports a sipunculan/molluscan affiliation is a shared “molluscan cross” cleavage pattern in the blastomeres (Heath 1899; Gerould 1907; Baba 1951; van Dongen and Geilenkirchen 1974; Rice 1975, 1985; Verdonk and van den Biggelaar 1983), which has been interpreted as a synapomorphy uniting these two groups (Scheltema 1993, 1996). Annelids and echiurans possess a different arrangement of the blastomeres, termed an “annelid cross” (Gerould 1907). As an extension of this proposed relationship, Scheltema (1996) hypothesized further homology of sipunculan larval characteristics to those of larval and adult mollusks, including the ventral buccal organ of sipunculan larvae with the odontophore of mollusks, the ciliated lip below the mouth of sipunculan pelagosphera stage larvae with the foot of a molluscan pediveliger larva, and the lip glands of sipunculan larvae with the pedal glands of larval chitons and adult neomeniid aplacophorans.

We report the sequence of about half (7,470 nt) of the mtDNA of the first representative of the phylum Sipuncula, *P. gouldii*. We analyze genomic features in comparison with those of other animal mtDNAs and compare both gene arrangement and inferred amino acid sequences to resolve the phylogenetic placement of this phylum. These comparisons strongly support the closer relationship of Sipuncula with Annelida to the exclusion of Mollusca and other taxa.

Materials and Methods

Molecular Analysis

Live specimens of *P. gouldii* (Fisher 1950) were purchased from the Marine Biological Laboratory at Woods Hole, Mass. We isolated total DNA from excised retractor muscles using a CTAB method described in Collins et al. (1996). We also isolated total DNA from the upper “nuclear” fraction of the CsCl purification method for mitochondrial DNA (Dowling et al. 1996). The upper layer of that preparation contains both nuclear and nonsupercoiled (i.e., nicked) mitochondrial DNA.

Initially, primers designed to match generally conserved regions of the mtDNA were used in the PCR to amplify four short (450–710 nt) fragments from *cox1* (primers LCO1490 and HCO2198; Folmer et al. 1994), *cox3* (primers COIIF and COIIB; Boore and Brown 2000), *cob* (primers CytbF and CytbR; Boore and Brown 2000), and *rrnL* (primers 16SARL and 16SBRH; Palumbi 1996). DNA sequences obtained from these fragments were used to design oligonucleotides that were then employed in “long-PCR” (Barnes 1994) to amplify the portions of the mtDNA spanning *cox1–cox3*, *cox3–cob*, and *cob–rrnL*. Reaction conditions and results, fragment purification, DNA sequence determination and assembly, and gene identifications were as in Boore and Brown (2000). All sequence was determined

for both strands using synthetic oligonucleotides to “primer-walk” through the long-PCR-amplified fragments. This 7,470-nt sequence was deposited in GenBank under accession number AF374337.

Phylogenetic Analysis of Sequences and Gene Arrangements

The invertebrate mitochondrial genetic code was used to infer the amino acid sequences of the six protein-encoding genes identified in the studied portion of *P. gouldii* mtDNA. Those of Cob, Cox1, Cox2, and Cox3 were aligned to the homologs of 16 other, phylogenetically diverse animals (listed, along with citations, in table 1) using ClustalW as incorporated in MacVector 6.5 (Accelrys). The other two inferred protein sequences, Atp8 and Nad6, were judged to be too divergent to align with confidence and thus were not used in this phylogenetic analysis. The BLOSUM matrix was used to weight shared amino acids, with gap and extension penalties of 10 and 1, respectively. Amino acid alignments were then adjusted manually; these alignments can be viewed at EMBL accessions ALIGN_000119 and ALIGN_000121–ALIGN_000123.

Protein alignments were analyzed using three methods: (1) maximum likelihood (ML) quartet sampling with Tree-Puzzle 5.0 (Strimmer and von Haeseler 1996) using the mtREV24 model of substitution (Adachi and Hasegawa 1996), with gamma-distributed rates (eight categories) and parameters estimated from the data set and with no clock assumed; (2) equal-weighted parsimony (MP) (heuristic search of 1,000 random sequence additions); and (3) neighbor-joining (NJ) analysis of total pairwise distance of amino acids (Saitou and Nei 1987). The two latter methods were each implemented using PAUP*, version 4.0b4a (Swofford 2000). Gaps were treated as missing data. Data were also bootstrapped using the MP and NJ methods (1,000 replicates each). The single shortest tree for the heuristic parsimony analysis was used to calculate branch support values (Bremer 1994) using the program AutoDecay (Eriksson 1998) in combination with PAUP*.

Additional analyses specifically tested the effect of grouping *P. gouldii* with the two annelids versus with the mollusk/brachiopod group, first using a Kishino-Hasegawa test (Kishino and Hasegawa 1989) as implemented in Tree-Puzzle (<http://www.tree-puzzle.de/>) and second using a nonparametric Wilcoxon signed-ranks test (Templeton 1983) as implemented in PAUP*. For the partially determined mtDNA sequences of two other annelids, *Galathealinum brachiosum* and *Helobdella robusta* (Boore and Brown 2000), only the partial sequences of the *cob* genes are known, along with the complete sequences of *cox1*, *cox2*, and *cox3*. These were included in a separate analysis to verify that the phylogenetic placement of *P. gouldii* was not affected.

We also compared the gene arrangement of this portion of *P. gouldii* mtDNA with those completely determined for 13 other animals (table 1). These taxa were the same as those included in the sequence analysis ex-

Table 1
Complete Mitochondrial Gene Arrangements for the Taxa Considered in this Phylogenetic Analysis, with Those Matching Partial *Phascolopsis gouldii* Arrangements Shown in Boldface

Taxon ^a (classification)	Gene Arrangement	Reference
<i>Alligator</i> ^b (Chordata: Vertebrata)	cox1, ^c - S2, ^d D, cox2, K, atp8, atp6, cox3, G, nad3, R, nad4L, nad4, S1, H, L1, nad5, -nad6, -E, cob, T, -P, F, rrnS, V, rrnL , L2, nad1, I, -Q, M, nad2, W, -A, -N, -C, -Y	Janke and Arnason (1997)
<i>Branchiostoma</i> (Chordata: Cephalochordata)	cox1, -S2, D, cox2, K, atp8, atp6, cox3, nad3, R, nad4L, nad4, H, S1, L1, nad5, G, -nad6, -E, cob, T, -P, rrnS, F, V, rrnL , L2, nad1, I, M, -Q, nad2, -N, W, -A, -C, -Y	Boore, Daehler, and Brown (1999)
<i>Balanoglossus</i> (Hemichordata)	cox1, -S2, -D, cox2, K, atp8, atp6, cox3, G, nad3, R, nad4L, nad4, H, S1, nad5, cob, E, T, -P, -nad6, F, rrnS, V, rrnL , L1, L2, nad1, -Q, I, M, nad2, -N, W, -A, -C, -Y	Castresana et al. (1998)
<i>Asterina</i> (Echinodermata: Asteroidea)	cox1, R, nad4L, cox2, K, atp8, atp6, cox3, -S2, nad3, nad4, H, S1, nad5, -nad6, cob, F, rrnS, E, T, -rrnL, -nad2, -I, -nad1, -L2, -G, -Y, D, -M, V, -C, -W, A, -L1, -N, Q, -P	Asakawa et al. (1995)
<i>Paracentrotus</i> (Echinodermata: Echinoidea)	cox1, R, nad4L, cox2, K, atp8, atp6, cox3, -S2, nad3, nad4, H, S1, nad5, -nad6, cob, F, rrnS, E, T, P, -Q, N, L1, -A, W, C, -V, M, -D, Y, G , L2, nad1, I, nad2, rrnL	Cantatore et al. (1989)
<i>Locusta</i> (Arthropoda: Hexapoda)	cox1, L2, cox2, D, K , atp8, atp6, cox3, G, nad3, A, R, N, S1, E, -F, -nad5, -H, -nad4, -nad4L, T, -P, nad6, cob , S2, -nad1, -L1, -rrnL, -V, -rrnS, I, -Q, M, nad2, W, -C, -Y	Flook, Rowell, and Gellissen (1995)
<i>Drosophila</i> (Hexapoda), <i>Daphnia</i> (Crustacea)	cox1, L2, cox2, K, D, atp8 , atp6, cox3, G, nad3, A, R, N, S1, E, -F, -nad5, -H, -nad4, -nad4L, T, -P, nad6, cob , S2, -nad1, -L1, -rrnL, -V, -rrnS, I, -Q, M, nad2, W, -C, -Y	Clary and Wolstenholme (1985); Crease (1999)
<i>Artemia</i> (Arthropoda: Crustacea)	cox1, L2, cox2, K, D, atp8 , atp6, cox3, G, nad3, A, R, N, S1, E, -F, -nad5, -H, -nad4, -nad4L, T, -P, nad6, cob , S2, -nad1, -L1, -rrnL, -V, -rrnS, I, -Q, M, nad2, W, -C, -Y	Valverde et al. (1994)
<i>Limulus, Ixodes</i> (Arthropoda: Chelicerata)	cox1, cox2, K, D, atp8 , atp6, cox3, G, nad3, A, R, N, S1, E, -F, -nad5, -H, -nad4, -nad4L, T, -P, nad6, cob , S2, -nad1, -L2, -L1, -rrnL, -V, -rrnS, I, -Q, M, nad2, W, -C, -Y	Staton, Daehler and Brown (1997); Black and Roehrdanz (1998)
<i>Katharina</i> (Mollusca: Polyplacophora)	cox1, D, cox2, atp8, atp6, -F, -nad5, -H, -nad4, -nad4L, T, -S2, -cob, -nad6, P, -nad1, -L2, -L1, -rrnL, -V, -rrnS, -M, -C, -Y, -W, -Q, -G, -E, cox3, K, A, R, N, I, nad3, S1, nad2	Boore and Brown (1994)
<i>Loligo</i> (Mollusca: Cephalopoda)	cox1, -C, -Y, -E, N, cox2 , -M, R, -F, -nad5, -nad4, -nad4L, T, -L2, -G, A, D, atp8 , atp6, -H, -L1, cox3, nad3, -S2, -cob, -nad6, -P, -nad1, -Q, I, -rrnL, -V, -rrnS, -W, K, S1, nad2	Sasuga et al. (1999); GenBank AB009838
<i>Terebratulina</i> (Brachiopoda)	cox1, cox2, D, atp8 , atp6, Y, C, M, rrnS, V, rrnL , L1, A, L2, nad1, nad6, P, cob, K, N, S2, nad4L, nad4, Q, W, H, nad5, F, E, G, cox3 , T, R, I, nad3, S1, nad2	Stechmann and Schlegel (1999)
<i>Lumbricus</i> (Annelida: Oligochaeta)	cox1, N, cox2, D, atp8, Y, G, cox3, Q, nad6, cob , W, atp6, R, H, nad5, F, E, P, T, nad4L, nad4, C, M, rrnS, V, rrnL , L1, A, S2, L2, nad1, I, K, nad3, S1, nad2	Boore and Brown (1995)
<i>Platynereis</i> (Annelida: Polychaeta)	cox1, N, cox2 , G, Y, atp8, M, D, cox3, Q, nad6, cob , W, atp6, R, H, nad5, F, E, P, T, nad4L, nad4, rrnS, V, rrnL , L1, S2, A, L2, nad1, I, K, nad3, S1, nad2, C	Boore and Brown (2000); GenBank AF178678

^a Only the genus is named here for brevity. Full binomens appear in figures 4 and 5.

^b *Alligator* mtDNA differs from the gene arrangement typical of vertebrates (see Boore 1999) by only the position of *trnH*, which is not considered in this analysis since it is not in the sequenced portion of *P. gouldii* mtDNA.

^c All of these genomes are circular but have been graphically linearized at the arbitrarily chosen *cox1* gene.

^d Transfer RNA genes are abbreviated by the one-letter code for the corresponding amino acid, with the two each for leucine and serine being differentiated by numerals. A minus sign indicates opposite (i.e., right-to-left as shown) transcriptional orientation.

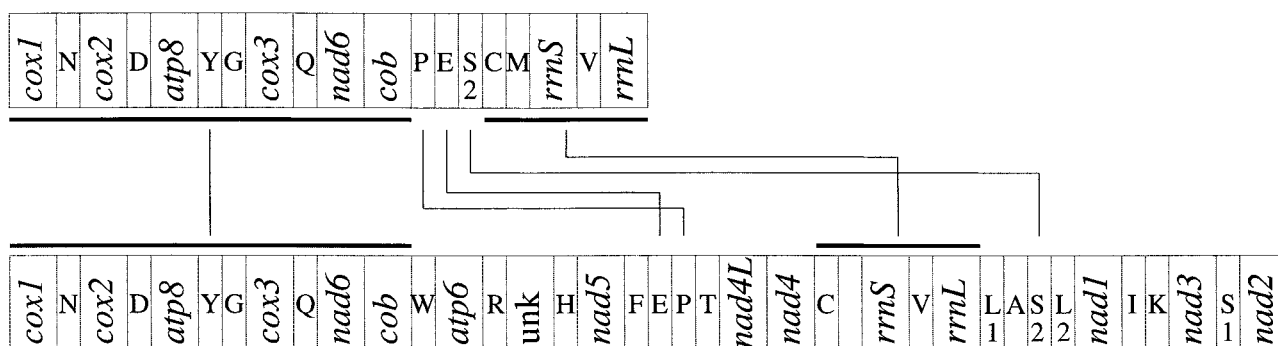
Phascolopsis gouldii (Sipuncula)*Lumbricus terrestris* (Annelida, Oligochaeta)

FIG. 1.—Gene map of the sequenced portion of the mtDNA of the sipunculid *Phascolopsis gouldii* compared with the complete map of the oligochaete annelid *Lumbricus terrestris* (Boore and Brown 1995), which has been graphically linearized at *cox1*. All genes are transcribed from left to right as depicted. Transfer RNA genes are designated by a single letter for the corresponding amino acid. There are two tRNAs for each of leucine and serine that are differentiated by numerals: L1 and L2 recognize the codons CUN and UUR, respectively, and S1 and S2 recognize the codons AGN and UCN, respectively. All other genes are designated by standard annotation and are not to scale. Lines connect homologous genes or blocks of genes for each mtDNA.

cept for three: *Daphnia pulex* and *Ixodes hexagonus*, were omitted since they had gene arrangements identical to those of *Drosophila yakuba* and *Limulus polyphemus*, respectively, and *Locusta migratoria* was omitted because it differed from the gene arrangement of *D. yakuba* by only a single tRNA position. A matrix was constructed that scored 36 characters, each as “upstream of X” or “downstream of X,” where “X” refers to each of the 19 genes identified in the studied portion of *P. gouldii* mtDNA. Coded character states were the 5' or 3' end of the adjacent gene. This matrix of scored gene adjacencies was analyzed using parsimony criteria with the programs PAUP*, version 4.0b4a (Swofford 2000), and MacClade (Maddison and Maddison 1992).

Results and Discussion**Gene Content and Organization**

This 7,470-nt portion of *P. gouldii* mtDNA contains six protein-encoding genes (*cox1* is incomplete by only a few nucleotides at the 5' end), 11 tRNA genes, and 2 rRNA genes (*rrnL* is incomplete at the 3' end). Remarkably, the arrangement of these genes is very similar to that of the oligochaete annelid *Lumbricus terrestris* (Boore and Brown 1995; fig. 1). Among animal mtDNAs studied so far (see Boore [1999] and the Genomic Diversity link at <http://www.jgi.doe.gov>), the entire block of 11 genes spanning *cox1* through *cob* is uniquely shared between *P. gouldii* and *L. terrestris* (table 1). Furthermore, the five genes in the block from *trnC* through *rrnL* are arranged identically between these two mtDNAs and also in the mtDNAs of the chiton *Katharina tunicata* (Boore and Brown 1994) and the brachiopod *Terebratulina retusa* (Stechmann and Schlegel 1999). Intervening between these two blocks of genes are *trnP*, *trnE*, and *trnS2(uga)*. This is not similar to the arrangement of these genes in any other animal so far studied (Boore 1999). All of the genes identified

in this portion of *P. gouldii* mtDNA are found in the same transcriptional orientation, as is true of the mtDNAs of *L. terrestris* (Boore and Brown 1995), the polychaete annelid *Platynereis dumerilii* (Boore and Brown 2000; GenBank AF178678), an ascidian (Yokobori et al. 1999), a mussel (Hoffmann, Boore, and Brown 1992), the two sampled brachiopods (Stechmann and Schlegel 1999; Noguchi et al. 2000), the four sampled nematodes (Okimoto et al. 1991, 1992; Keddie, Higazi, and Unnasch 1998), the seven sampled flatworms (Le et al. 2000; GenBank accession number AB018440), and a sea anemone (Beagley, Okimoto, and Wolstenholme 1998).

In many mtDNAs, *atp6* directly follows *atp8*. This is the case even for organisms that are unambiguously outgroup taxa to those considered here, including a sponge (Watkins and Beckenbach 1999) and yeast (e.g., Sekito et al. 1995). The explanation may be that the transcript of these two genes is not cleaved, such that these genes are cotranslated from a single bicistron, as has been shown for mammalian mitochondrial systems (Fearnley and Walker 1986). The gene for *atp6* is not contained in the studied portion of *P. gouldii* mtDNA, but it clearly does not follow *atp8*. So far, these genes have been found separated (see Boore 1999) only for animals that are part of the proposed superphylum Eutrochozoa (Ghiselin 1988; Mollusca, Annelida, and others, but not including Arthropoda), leading to the speculation that the loss of cotranslation of these mRNAs could be a derived feature uniting this group.

Base Composition and Codon Usage

Overall, the 7,470 nt determined for *P. gouldii* mtDNA are 63.1% A+T, similar to the percentage found for the whole mtDNA sequence of *L. terrestris* (61.6%; Boore and Brown 1995). As is common among mtDNAs, CG is the least frequent dinucleotide, occur-

ring at only 53% of expectation given the proportion of G and of C observed in this sequence.

Leucine is inferred to be the most frequent amino acid, present 243 times in these six inferred amino acid sequences, followed by four amino acids in nearly equal numbers, each about half as common as leucine: alanine (124), isoleucine (123), phenylalanine (135), and serine (125). Cysteine is the least frequently used amino acid, occurring only 14 times. These values are all similar to those for the amino acid usage of *L. terrestris* homologs (see table 2).

The DNA strand shown in figure 2 is rich in pyrimidines (C and T). One way to measure this is called skewness, an assessment of whether the G of GC pairs (GC-skew, calculated as $[G - C]/[G + C]$) and the A of AT pairs (AT-skew, calculated as $[A - T]/[A + T]$) are more commonly on the considered strand (Perna and Kocher 1995). GC-skew for the entire 7,470 nt is -0.239 , that for the protein-encoding portion only is -0.265 , and that for third codon positions (see below), which, presumably, are more free to change and therefore better reflect mutational bias, is a remarkable -0.754 . AT-skew values are much less biased: that for the entire region is -0.080 , that for the protein-encoding portion is -0.138 , and that for the third codon positions is -0.028 .

Gene Initiation and Termination

Most of the studied animal mitochondrial genomes contain one or more genes that initiate with some alternative to the standard ATG. Annelids appear as a notable exception: based on the few animals sampled, ATG is used as the initiation codon for all, or nearly all, genes (Boore and Brown 2000). Consistent with this and with the phylogenetic placement of Sipuncula advanced here (see below), each of the *P. gouldii* protein-encoding genes in this portion of the mtDNA appears to start with ATG (the initiator of *cox1* is not present in this sequence).

Mitochondrial genes often terminate with abbreviated stop codons, where a single T or TA is presumably completed by polyadenylation to a TAA stop codon (Ojala, Montoya, and Attardi 1981). This appears to be the case for three genes in *P. gouldii* mtDNA. The least certain of these is for *atp8*, since the next nucleotide (A), otherwise part of *trnY*, would complete the stop codon (fig. 2). If *cob* were to extend to the first complete stop codon, it would overlap *trnP* by 41 nt, and if *nad6* were to do so, it would overlap *cob* by 20 nt. In each case, homologous genes of related animals are similar in predicted amino acid sequences to the abbreviated forms in *P. gouldii* and have no similarity to the overlapping extensions.

All genes are in a compact arrangement, with a total of only 13 noncoding nucleotides. These are distributed in six regions of 1–4 nt each. There is a CC between *trnY* and *trnG*, but otherwise, all are A or T. No genes are inferred to overlap except for the possibility of a single nucleotide shared between *atp8* and *trnY* (see above).

Transfer RNAs

There are 11 sequences identified with the potential for folding into tRNA-like structures (fig. 3). Each has an anticodon matching exactly one tRNA in *L. terrestris* mtDNA. Each has a seven-member amino-acyl acceptor stem (four with a single mismatch each) and a five-member anticodon stem (again, four with a single mismatch each). Two have 5 nt in the extra arm, and all others have 4 nt. All but four have A immediately preceding the anticodon arm. For all except tRNA(N) and tRNA(S2), there are three to five nucleotide pairs in both the DHU and the T Ψ C arms. All but four tRNAs have TA immediately preceding the DHU arm. The nucleotides preceding the anticodon are YT for all tRNAs, and the nucleotide following the anticodon is A for all but tRNA(P).

It is common for tRNA(S) in many mtDNAs to lack a paired DHU arm and, in some cases, to have a six-member anticodon stem (Yokogawa et al. 1991). There is potential for *P. gouldii* tRNA(S2) to have such an extended anticodon stem and also to have a paired DHU arm with an unusual structure, with only one nucleotide separating the amino-acyl acceptor arm from the DHU arm. Several other serine-specifying tRNAs have been identified in animal mtDNAs that have this identical pairing potential (Boore and Brown 2000).

Phylogenetic Reconstruction

Analyses based on maximum likelihood (ML), maximum parsimony (MP), and neighbor-joining of pairwise distances (NJ) of inferred amino acids were in almost complete agreement (fig. 4). In all cases, *P. gouldii* was sister to the two annelids, and the sipunculan, annelids, mollusks, and brachiopods were placed within a well-supported Eutrochozoa (Ghiselin 1988). Furthermore, there was strong support for the monophyly of several traditionally recognized groups (at least as composed by this small sampling), such as Insecta, Chelicerata, Arthropoda, and Echinodermata. Of the 1,396 aligned inferred amino acids, 673 were parsimony-informative. A heuristic search with 1,000 random stepwise additions of taxa yielded a single tree (consistency index = 0.618, retention index = 0.366, and rescaled consistency index = 0.226), 4,552 steps in length, identical to that in figure 4 except for the resolution of two trichotomies.

Discrepancies among the ML, MP, and NJ analyses were in the placement of the brachiopod *T. retusa* and the relationship among the crustaceans *D. pulex* and *Artemia franciscana*. In MP and NJ trees, *T. retusa* was placed as sister to *Loligo bleekeri*, albeit with less than 52% bootstrap support in either case, whereas in the ML tree, *T. retusa* was placed as sister to a clade of the two mollusks (69% of quartets sampled). The two crustaceans were paraphyletic in the MP analysis, albeit with poor (<50%) bootstrap support, but grouped together in both the ML and NJ analyses. This association was found in 95% of the quartets in ML and with 82% of the bootstrap resamplings in NJ.

Table 2
Codon Usage for the Protein-Encoding Genes in the Sequenced Portion of *Phascolopsis gouldii* mtDNA (1,591 codons) Compared with that of the Homologous Portions of *Lumbricus terrestris* mtDNA

AMINO ACID	CODON FAMILY	<i>P. gouldii</i>					<i>L. terrestris</i>				
		All	Third Codon Position				All	Third Codon Position			
			T	C	A	G		T	C	A	G
Phe (F)	TTY	135	93	42	—	—	123	74	49	—	—
Leu (L)	TTR	80	—	—	80	0	61	—	—	56	5
	CTN	163	74	34	53	2	150	39	32	72	7
Total Leu		243	—	—	—	—	211	—	—	—	—
Ile (I)	ATY	123	85	38	—	—	136	93	43	—	—
Met (M)	ATR	66	—	—	57	9	70	—	—	55	15
Val (V)	GTN	83	26	16	38	3	101	19	15	45	22
Ser (S)	TCN	94	46	25	23	0	91	26	30	33	2
	AGN	31	4	5	22	0	39	9	6	19	5
Total Ser		125	—	—	—	—	130	—	—	—	—
Pro (P)	CCN	90	37	15	36	2	85	30	24	25	6
Thr (T)	ACN	90	33	20	36	1	106	31	30	42	3
Ala (A)	GCN	124	49	29	42	4	121	41	37	35	8
Tyr (Y)	TAY	56	36	20	—	—	56	31	25	—	—
His (H)	CAY	64	35	29	—	—	50	18	32	—	—
Gln (Q)	CAR	33	—	—	31	2	27	—	—	24	3
Asn (N)	AAV	61	38	23	—	—	57	26	31	—	—
Lys (K)	AAR	30	—	—	28	2	26	—	—	19	7
Asp (D)	GAY	32	18	14	—	—	39	21	18	—	—
Glu (E)	GAR	28	—	—	25	3	32	—	—	21	11
Cys (C)	TGY	14	8	6	—	—	10	2	8	—	—
Trp (W)	TGR	56	—	—	51	5	56	—	—	44	12
Arg (R)	CGN	36	13	5	18	0	34	9	4	16	5
Gly (G)	GGN	102	22	22	43	15	102	21	22	29	30
Total		1,591	617	343	583	48	1,572	490	406	535	141

The tree shown in figure 4 was specifically compared with the alternative which repositions *P. gouldii* to be more closely related to mollusks than to annelids. Using ML, a Kishino-Hasegawa test (Kishino and Hasegawa 1989) demonstrated that the grouping of *P. gouldii* with the two annelids was significantly better (at the 5% significance level) than the grouping of *P. gouldii* with the brachiopod/mollusk clade. Using MP, nonparametric tests (Templeton 1983) of the single tree with *P. gouldii* as sister to annelids (4,552 steps) and the single most-parsimonious tree with *P. gouldii* sister to mollusks (4,582 steps) demonstrated that the (*P. gouldii*, *P. dumerilii*, *L. terrestris*) tree was significantly shorter ($P < 0.0001$). In case of bias from the inclusion of the ambiguously placed *T. retusa* among the mollusks, a second test omitting *T. retusa* was performed. The resultant sipunculan/annelid tree in this analysis was also significantly shorter ($P < 0.0001$) than (*P. gouldii*, *L. bleekeri*, *K. tunicata*).

Two other annelids, *Galathealinum brachiosum* and *Helobdella robusta*, were included in a separate analysis, since each lacks a complete sequence for *cob* (Boore and Brown 2000). Their inclusion had no effect on the phylogenetic placement of *P. gouldii*, and these two taxa joined *P. dumerilii* and *L. terrestris* in a topology consistent with the results of the earlier study (not shown).

Gene arrangements were also compared for phylogenetic reconstruction by scoring gene adjacencies as characters for cladistic analysis. The strict consensus of 30 equally parsimonious trees is shown in figure 5. The

unusual placement of *Alligator mississippiensis* as basal to the other included deuterostomes is not well supported, since it is based on the sharing of a single gene boundary (*trnI*, *trnM*) between *Branchiostoma floridae* and *Balanoglossus carnosus* in contrast to an alternative arrangement (*trnI*, *-trnQ*, *trnM*) in *A. mississippiensis* and two outgroup taxa (*D. yakuba* and *L. polyphemus*). This shared gene boundary has evidently been created by two independent translocations of *trnQ* from its primitive position (between *trnI* and *trnM*), leading in one case to the arrangement *nad1*, *-trnQ*, *trnI*, *trnM*, *nad2* in *B. carnosus* and in the other to *nad1*, *trnI*, *trnM*, *-trnQ*, *nad2* in *B. floridae*.

Moving *P. gouldii* to make it a sister taxon to an (*L. bleekeri*, *K. tunicata*) clade or to an ((*L. bleekeri*, *K. tunicata*), *T. retusa*) clade increases tree lengths by seven steps in each case for this gene arrangement analysis.

Our findings are in general agreement with several published studies (Winnepenninckx, Bäckeljau, and De Wachter 1995 [parsimony-based analysis]; Giribet et al. 2000 [18S-only tree]; Regier and Shultz 1998) in that sipunculans are closely associated with annelids to the exclusion of mollusks. Other studies, or portions thereof, give results to the contrary. Comparisons of 18S rDNA sequences have placed Sipuncula as basal and sister to an assemblage of worms and mollusks (Field et al. 1988; Winnepenninckx, Bäckeljau, and De Wachter 1995 [neighbor-joining analysis]; Giribet et al. 2000 [when combined with morphological characters]) or as sister to only one of two ectoprocts analyzed (Mackey et al. 1996). The conflicts among these studies indicate that

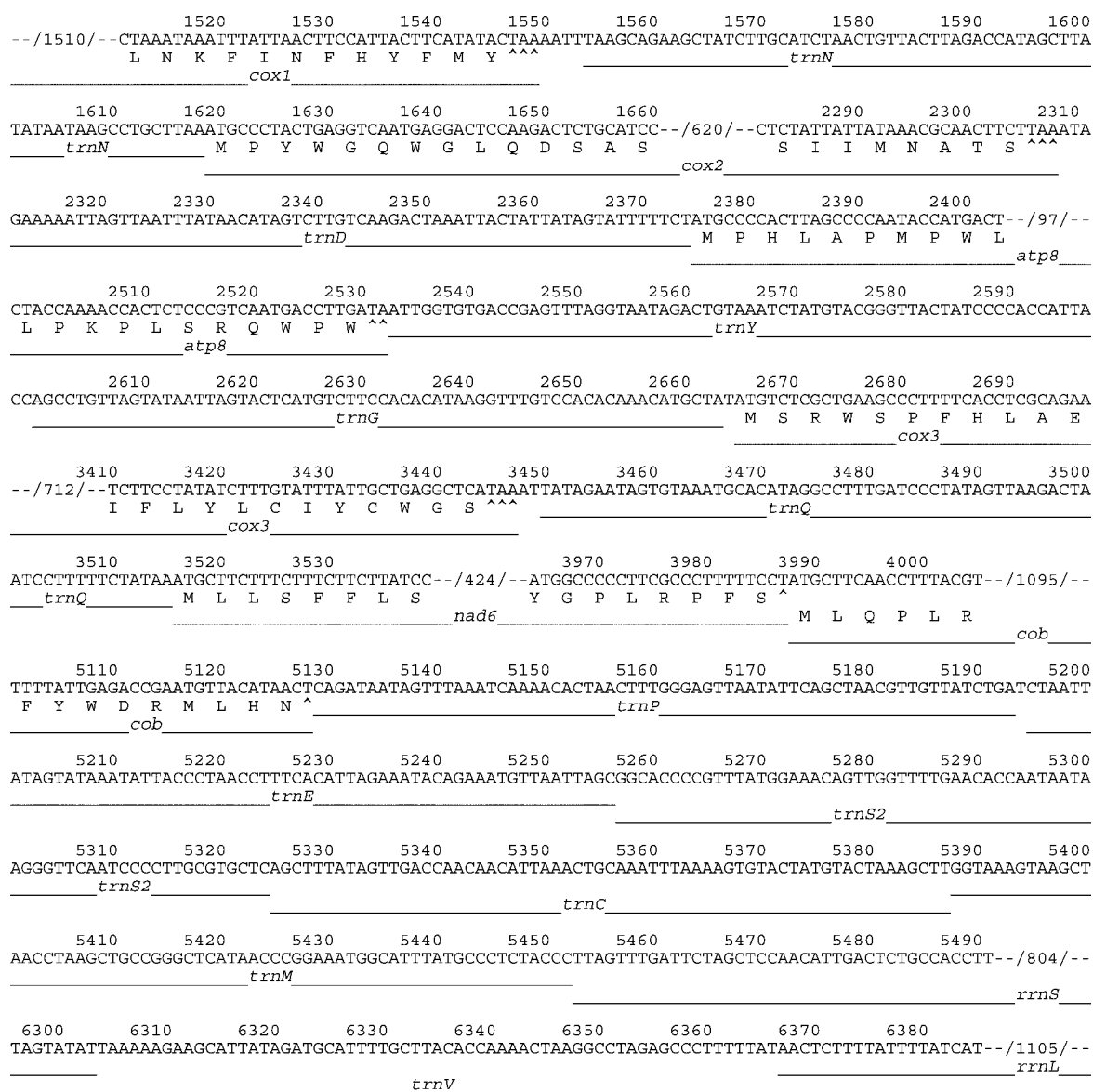


FIG. 2.—Abbreviated representation of the 7,470 nt determined of the mtDNA of the sipunculid *Phascolopsis gouldii*. Center portions of each protein- or rRNA-encoding gene have been replaced with a numeral indicating the number of omitted nucleotides. Nucleotides forming stop codons, partial or complete (see text), are marked with a caret.

comparisons of short DNA sequences lack the necessary resolving power at this level of relationship. Furthermore, some of the associations between sipunculans and annelids found in these past studies were probably discounted based on the problematica associated with these data (Maley and Marshall 1998).

Conclusions

The results of this study constitute the first genomic-level sequence from the phylum Sipuncula and the first molecular evolutionary treatment of the group through comparison of mitochondrial genomes. The shared mitochondrial gene orders suggest a close phylogenetic affinity between sipunculan and annelid taxa, and phylogenetic analysis of their combined amino acid sequences also reflects this relationship. In short, sipun-

culans and annelids form a natural group within the Eutrochozoa separate from that of mollusks. While it might seem controversial from a modern evolutionary perspective, a vermiform bauplan may not represent the simplest ancestral holdover or a failure to develop elaboration beyond the simplest hydrostatic skeleton, but, rather, it may ultimately prove to be a unifying characteristic among eutrochozoan worms.

Likewise, many characteristics that have been hypothesized to link sipunculans with mollusks, e.g., developmental pattern, lack of segmentation, etc., must be reevaluated. The presence of a “molluscan cross” may be ancestral in all eutrochozoans and subsequently lost in the annelids, or possibly is a convergent pattern that has independent origins in the two phyla. Recent reports of segmental development in chitons (Jacobs et al. 2000) suggest that segmentation in some form may be present

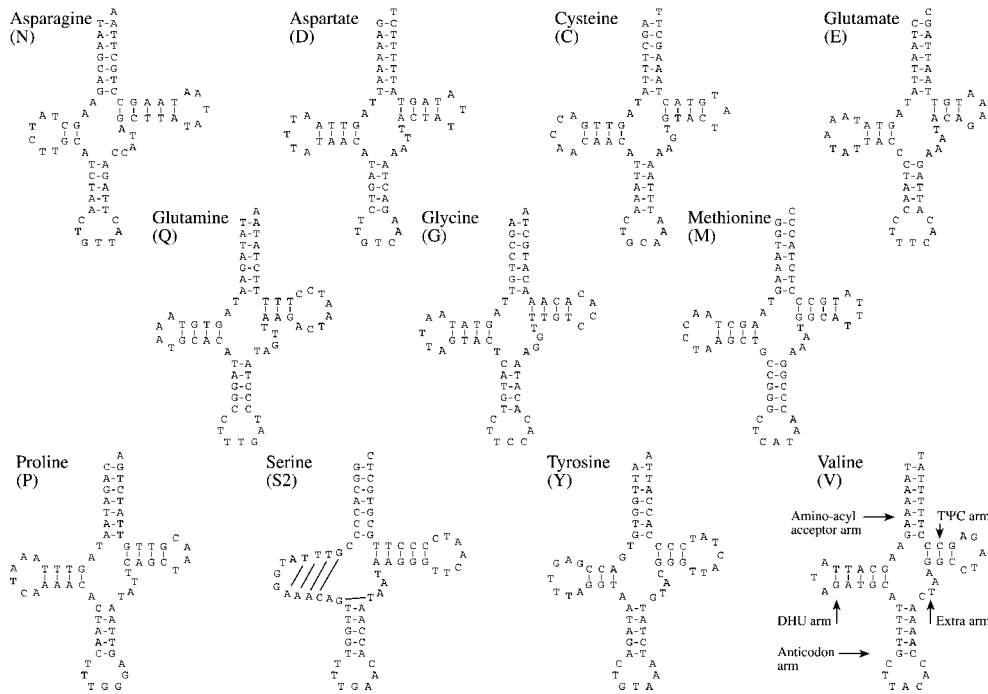


FIG. 3.—DNA sequences of the 11 tRNA genes proposed for *Phascolopsis gouldii* mtDNA folded into the standard cloverleaf structure. Two serine-specifying tRNAs are normally encoded by animal mtDNAs, and it is the one predicted to recognize the codon UCN that is found here; this tRNA is designated S2 for consistency with our earlier work, although there is no universal convention for this. Potential base-pairing for an unusual DHU arm of tRNA(S2) that would have only 1 nt after the amino-acyl acceptor arm and 2 nt before the anticodon arm (see text) is indicated by connecting lines. Structural elements are identified for tRNA(V).

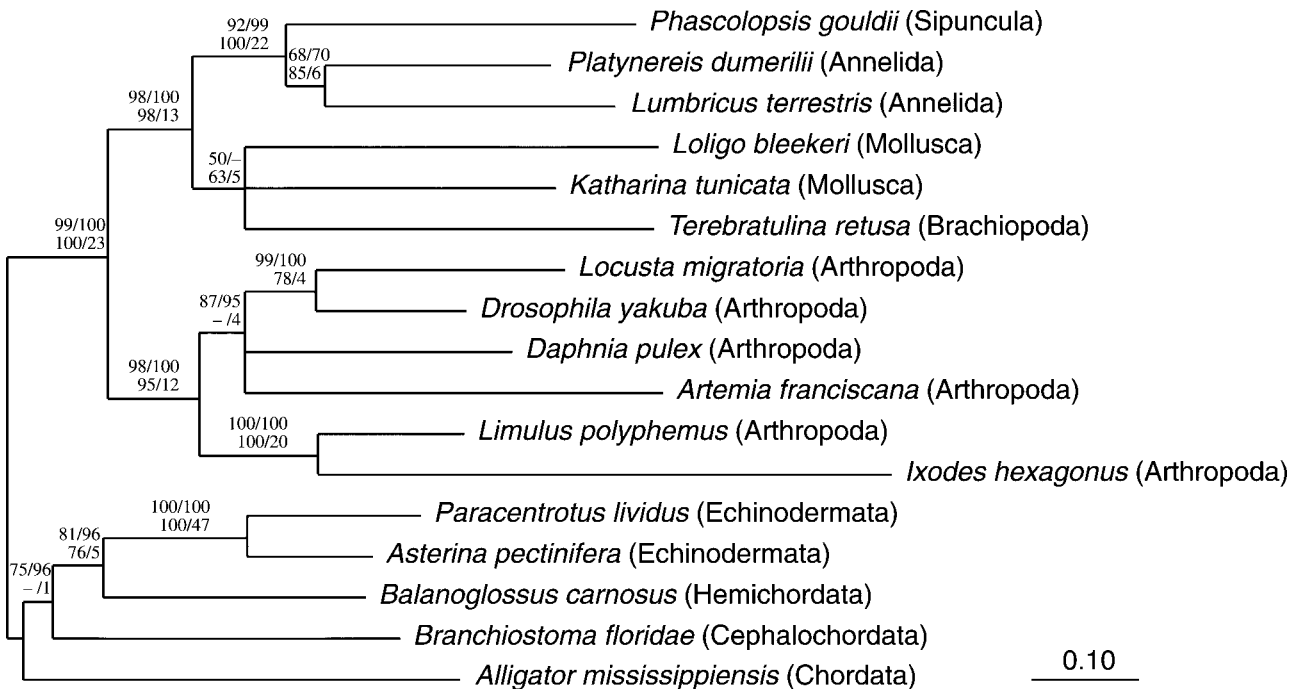


FIG. 4.—Phylogenetic tree based on comparisons of amino acid sequences for Cob, Cox1, Cox2, and Cox3 for 17 taxa. The tree was rooted with the deuterostome sequences, and branch lengths were assigned using Tree-Puzzle (Strimmer and von Haeseler 1996). Numbers above branches are the percentage of puzzle quartets supporting this branch, followed by bootstrap support using distance criteria, then parsimony criteria (a dash denotes <50% bootstrap support), then the parsimony branch support value (Bremer 1994). The two sets of relationships for which the methods give differing results (monophyly vs. paraphyly of the crustaceans *Daphnia pulex* and *Artemia franciscana* and the placement of *Terebratulina retusa* as sister to *Loligo bleekeri* vs. as sister to both mollusks) were subsequently collapsed to polytomies, so this is a strict consensus tree of the three phylogenetic methods employed (maximum likelihood, parsimony, and neighbor joining; see text).

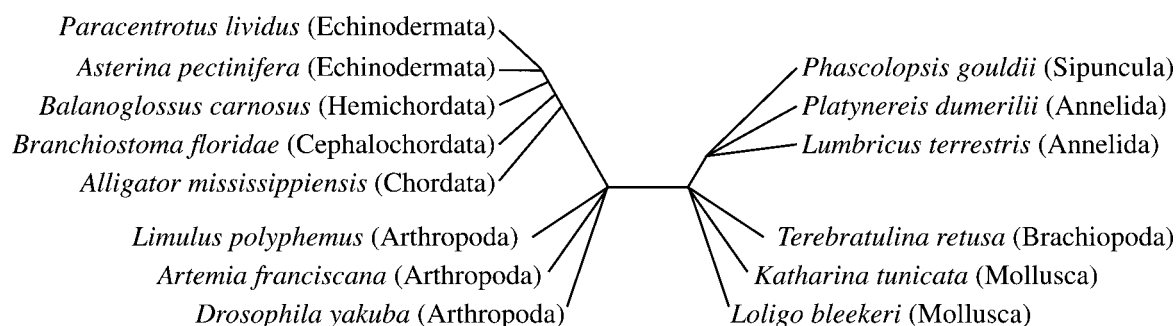


FIG. 5.—Phylogenetic tree based on gene arrangement comparisons using an adjacency matrix. This is the strict consensus of 30 equally parsimonious trees. The unusual placement of *Alligator mississippiensis* is poorly supported (see text).

in all spiralian, so its loss in other mollusks and worm groups may not be surprising. Certainly, the detailed examination of hypothesized synapomorphic larval and adult morphology in the sipunculans and molluscan groups (Scheltema 1993, 1996) will need careful reconsideration to exclude convergence or oversight of annelid larval and adult homologs.

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