

## The Complete DNA Sequence of the Mitochondrial Genome of a "Living Fossil," the Coelacanth (*Latimeria chalumnae*)

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### ABSTRACT

The complete nucleotide sequence of the 16,407-bp mitochondrial genome of the coelacanth (*Latimeria chalumnae*) was determined. The coelacanth mitochondrial genome order is identical to the consensus vertebrate gene order which is also found in all ray-finned fishes, the lungfish, and most tetrapods. Base composition and codon usage also conform to typical vertebrate patterns. The entire mitochondrial genome was PCR-amplified with 24 sets of primers that are expected to amplify homologous regions in other related vertebrate species. Analyses of the control region of the coelacanth mitochondrial genome revealed the existence of four 22-bp tandem repeats close to its 3' end. The phylogenetic analyses of a large data set combining genes coding for rRNAs, tRNAs, and proteins (16,140 characters) confirmed the phylogenetic position of the coelacanth as a lobe-finned fish; it is more closely related to tetrapods than to ray-finned fishes. However, different phylogenetic methods applied to this largest available molecular data set were unable to resolve unambiguously the relationship of the coelacanth to the two other groups of extant lobe-finned fishes, the lungfishes and the tetrapods. Maximum parsimony favored a lungfish/coelacanth or a lungfish/tetrapod sistergroup relationship depending on which transversion:transition weighting is assumed. Neighbor-joining and maximum likelihood supported a lungfish/tetrapod sistergroup relationship.

COELACANTHS were for the first time recognized as a distinct taxonomic group >150 years ago by Louis Agassiz in his renowned book "*Recherches sur les Poissons Fossiles*" (1844). Since then, many more extinct coelacanth species have been discovered (CLOUTIER and FOREY 1991). These fossils range in age from the Early Devonian to the Late Cretaceous (HUXLEY 1861; WOODWARD 1891; STENSIÖ 1921). No fossil coelacanths are known from more recent times, and, hence, this group was believed to have gone extinct ~70–80 mya. However, in 1938, a living coelacanth (*Latimeria chalumnae*) was trawled off the estuary of the Chalumna river, near East London, South Africa (SMITH 1939, 1956; COURTENAY-LATIMER 1979). Since the sensational discovery of this living fossil, a total of only ~200 specimens have been caught off the Comoro archipelago near the eastern coast of Africa in the Indian ocean. It proved to be the only known coelacanth population (SCHLIEWEN *et al.* 1993; FRICKE *et al.* 1995).

Coelacanths are large gray-blue marine fishes whose morphology has remained seemingly unchanged during the last 400 million years (but see FOREY 1991). During the last 50 years, detailed anatomic and functional morphological analysis of the circulatory, reproductive, locomotive, and respiratory systems of the coelacanth have been conducted (*e.g.*, MILLOT 1954, 1955;

SMITH 1956; THOMSON 1966; BEMIS and NORTH CUTT 1991; MANGUM 1991; SCHULTZE and CLOUTIER 1991; TAMAI *et al.* 1994). These fishes are characterized by the possession of several distinct morphological, physiological, and life history traits, such as lobed paired fins, an heterocercal tail with an apical lobe, an intracranial joint, a rostral electroreceptor organ, a urea retention mechanism and very large eggs (and give birth to living young) (*e.g.*, FOREY 1988; LONG 1995; MAISEY 1996). Unlike other sarcopterygians, the group to which they belong (see below), coelacanths lack choanae (internal nostril), maxillae and functional lungs.

Unfortunately, little is known about the coelacanth at a molecular level. To date, only the triosephosphate isomerase (KOLB *et al.* 1974), the  $\alpha$  and  $\beta$  parvalbumin (PECHERE *et al.* 1978), and the  $\alpha$  and  $\beta$  hemoglobin amino acid sequences (GORR *et al.* 1991), as well as the 18S rRNA (STOCK *et al.* 1991), 28S rRNA (HILLIS *et al.* 1991; ZARDOYA and MEYER 1996b), cytochrome *b* (MEYER and WILSON 1990), 12S rRNA (MEYER and WILSON 1990; MEYER and DOLVEN 1992; HEDGES *et al.* 1993), 16S rRNA (HEDGES *et al.* 1993), cytochrome oxidase subunit 1 (YOKOBORI *et al.* 1994), and Mhc class I (BETZ *et al.* 1994) nucleotide sequences have been reported from the coelacanth (reviewed by MEYER 1995).

Despite much prior effort, the phylogenetic position of the coelacanth still remains controversial (reviewed by MEYER 1995). It is well established that the coelacanth, together with lungfishes, and the extinct rhipidistians form the subclass Sarcopterygii, the lobe-

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**TABLE 1**  
**PCR and sequencing primers used in the analysis of the coelacanth mitochondrial genome**

PCR primers	Sequence (5' → 3')	Approximate product length (bp)
1. 16S F	AGT TCA GAC CGG AGC AAT CCA GG	600
2. LATI ND1 R	ACT TCG TAG GAA ATA GTT TGT GC	
3. LATI ND1 F1	CCT TGA TCG GGG CCC TTC GAG CA	600
4. LATI MET R	TCG GGG TAT GGG CCC GAA AGC TT	
5. LATI GLN F	TGA ACC TAT ACT AAA GAG ATC AA	750
6. LATI ND2 R	ATT TTT CGT AGC TGC GTT TGA TT	
7. LATI ND2 F	GGA CTT ATC CTG TCG ACT TGA CA	700
8. LATI TRP R	TTA AAG CTT TGA AGG CTC TTA GT	
9. LATI TRP F	AAG CAT CAT AAC CCT CAT AGC ACT	700
10. LATI COI R	GGC ATC ACT ATA AAG AAG ATT AT	
11. LATI COI F1	GAT GAC CAA ATT TAT AAT GTA GT	700
12. LATI COI R1	ATT GCC ATT ATC GCT CAG ACT AT	
13. LATI COI F2	AAG AAA GAA CCA TTC GGG TAT AT	1200
14. H7886 <sup>a</sup>	TAN SWY CAR TAY CAY TGR TGN CC	
15. LATI COII F	TAA TTG ATG AAG TCG AAA ACC CTC A	650
16. LATI AT8 R	TTA GGC TCA TGG TCA GGT TCA	
17. LATI AT8 F	AGT GAA TGC CTC ARY TAA AYC C	1000
18. LATI COIII R	TTT TGT ACA GGT AGT GTG TGG TG	
19. LATI COIII F	ATA TAT CAA TGA TGA CGA GA	600
20. LATI COIII R1	ACA TCA ACG AAA TGT CAG TAC CA	
21. LATI COIII F1	TAC CAC TTC ACA TCA AAC CAC CA	1600
22. LATI ND4 R	TAG TAG GAC GGC GGC TAG TAC TAT	
23. LATI ND4 F1	CCT AAR GCC CAY GTA GAR GC	900
24. LATI LEU R	TTT GCA CCA AGA GTT TTT GGT TCC TA	
25. LATI LEU F	CTA AAG GAT AAT AGC TCA TCC ATT	600
26. LATI ND5 R	CAY CAG CCR ATT ART ARR AAT GAY AT	
27. LATI ND5 F	CAR YTA TTY ATC GGN TGR GAR GG	650
28. LATI ND5 R1	CCY ATY TTT CKG ATR TCY TGY TC	
29. LATI ND5 F2	TAA AGC AAT GCT GTT CCT ATG CT	850
30. LATI ND5 R3	AAN AGN GTN AGR TAN GTY TTR AT	
31. LATI ND5 F3	ATC AAC AAC TCG CAA CAA GGA C	1100
32. H15149 <sup>b</sup>	AAA CTG CAG CCC CTC AGA ATG ATA TTT GTC CTC A	
33. LATI CYTB F	TAC TTA CAA AAA GAA ACC TG	850
34. LATI THR R	CGG CTT ACA AGG GCC GAT GCT TT	
35. LATI THR-F	AGC CTT AGT AGC TTA AAC CC	1000
36. LATI12S-R	AGC AAG GCT GGG ACC AAA CCT TT	
37. L-Phe	TAA AGC ATA GCA CTG AAA ATG	850
38. 12Sb	GAG GGT GAC GGG CGG TGT GT	
39. 12Sa	TGG GAT TAG ATA CCC CAC TAT	800
40. 16S-1H	KTA GCT CRC CYA GTT TCG GG	
41. 16S-1L	AGT ACC GCA AGG GAA ARC TGA AA	800
42. 16S-2H	GAT TRY GCT ACC TTY GCA CGG TCA	
43. 16Sar <sup>c</sup>	CGC CTG TTT ATC AAA AAC AT	500
44. 16Sbr <sup>c</sup>	CCG GTC TGA ACT CAG ATC ACG T	

TABLE 1  
Continued

PCR primers	Sequence (5' → 3')	Approximate product length (bp)
45. LATI ND6-F 32. HI5149 <sup>b</sup>	ATC CCC CCY AAA TAR ATC AAA AA AAA CTG CAG CCC CTC AGA ATG ATA TTT GTC CTC A	650
46. LATI 16S-F 47. LATI 16S-R	CAA TCC TTT CCA AGA GTC CAA AT GTA CTG GCT CCT TAA TGA ATA GT	450
Sequencing		
LATI COI F3	CCC TGC TAT GAG CAC TAG GAT TT	
LATI ND3 F	AAT GTG GCT TTG ATC CTC TAG GA	
LATI ND3 R	TTT AGT CGC TCT GTT TGA TTA CC	

IUB code: R, A/G; Y, C/T; W, A/T; S, G/C; and K, G/T.

<sup>a</sup>YOKOBORI *et al.* (1994).

<sup>b</sup>KOCHER *et al.* (1989).

<sup>c</sup>PALUMBI *et al.* (1991).

finned fishes (ROMER 1966; CARROLL 1988; AHLBERG 1991; CLOUTIER and AHLBERG 1996) from which tetrapods were originated. The origin of land vertebrates from water-bound lobe-finned fishes involved both morphological and physiological innovations and was one of the most significant events in the history of vertebrates (PANCHEN and SMITHSON 1987). There is recent strong evidence supporting elpistostegids (or panderichthyids), a group of rhipidistians, as the extinct sister group of the earliest tetrapods (VOROBYEVA and SCHULTZE 1991; CLACK 1994; SCHULTZE 1994; AHLBERG *et al.* 1996; CLOUTIER and AHLBERG 1996).

However, there is no general agreement about the relationships among the coelacanth, lungfishes and the rhipidistian+tetrapod clade (*e.g.*, SCHULTZE 1994; for review, see MEYER 1995). Since molecular phylogenetic data can only be collected from extant species, they can contribute only in limited ways to this debate (MEYER and WILSON 1990; MEYER and DOLVEN 1992; HEDGES *et al.* 1993; YOKOBORI *et al.* 1994; ZARDOYA and MEYER 1996b; see MEYER 1995 for a review). However, the establishment of the phylogenetic relationships among

the three extant sarcopterygian groups, *i.e.*, lungfishes, the coelacanth and tetrapods, can be accomplished by the evolutionary analyses of molecular data sets and would help to recreate the sequence of evolutionary events that might have preadapted for the extinct sarcopterygian conquest of land. All three possible competing hypotheses that explain the relationships among living representatives of sarcopterygians, have been supported at various times by either morphological and/or molecular data: (1) the lungfishes as the closest living relatives of tetrapods (ROSEN *et al.* 1981; FOREY 1987, 1988; MEYER and WILSON 1990). (2) The coelacanth as the sister group of tetrapods among living vertebrates (FRITZSCH 1987; SCHULTZE 1987; GORR *et al.* 1991). And (3) lungfishes and the coelacanth form a monophyletic group and, hence, are equally closely related to tetrapods (NORTHCUTT 1987; FOREY 1988; CHANG 1991; SCHULTZE 1994; ZARDOYA and MEYER 1996b). The apparently rapid successive origin and subsequent diversification of the three remaining lineages of sarcopterygians back in the Devonian makes the determination of their phylogenetic relationships difficult. Many mor-

### *Latimeria chalumnae* mtDNA

16,407 bp

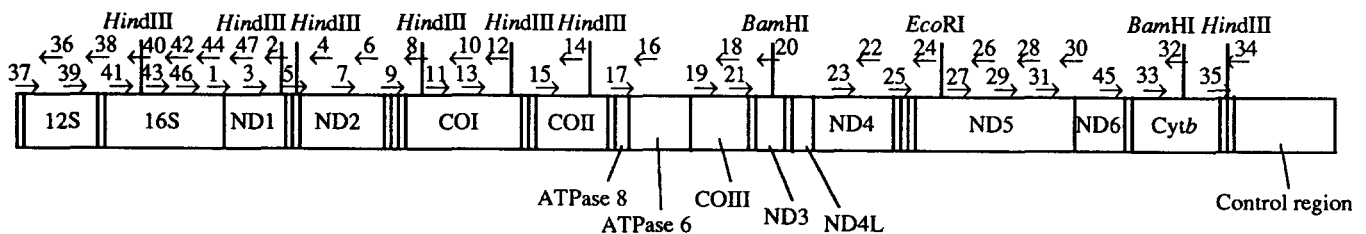


FIGURE 1.—Gene organization, restriction map, and cloning/sequencing strategy for the coelacanth mitochondrial genome. All protein coding genes are encoded by the H strand with the exception of ND6, which is coded by the L strand. Localization and direction of the primers used in the PCR amplification are denoted by arrows (see Table 1 for the primer DNA sequence associated with each number).

tRNA-Phe→ 12s rRNA→

1 GTGAGTGTAGCTTAAATAAAAAGCATAGCACTGAAGATGCTAAGATGAATTACAAAATATCCACTGACACAAGGTTTGGTCCCAGCCTTGCTATCAATT 100

101 TTAACCAGGATTACACATGCAAGCATCAACTCCCCAGTGAGAATGCCCTGACTTATCCGTCGAAAGATAACAGGGAGTAGGTATCAGGCACACAACATTA 200

201 ACGCTAGCCCAAGACACCTTGTCCAGCCACACCCCCAAGGGAAGTACAGCAGTATAGACATTAATAATAAGTGAAAACCTTGACTCAGCCATGGTTACAA 300

301 GGGCCCGTCAACTCCGTGCCAGCCACCGCGGTTACACGGAAGACCCAAAATGATAACACTACCGGCGTAAAGCGTGATTAAAGGACACCCACCATAATGG 400

401 AGCCACAAATAACTAAAGCTGTTATACGCACTTAAAAAATATGCTCATCACAGAAAGTAACCCAGCACCCAAAGGAACCTGAACCCACGAAAGCTA 500

501 AGAAACAAACTGGGATTAGATACCCCACTATGCTCAGCCCTAAACACAAACAATTCAAACACACACTGTTCCGCAAGGGAACACAGCGCCAGCTTCAA 600

601 ACCCAAAGGACTTGGCGGCACCTCAAACCCACCTAGAGGAGCCTGTTCTAAAAGTACAACCCCACTAACCTCACCATCCCTAGCCATTAAACAGCC 700

701 TATATACCGCCGTCGCGACCCACCCTGTGAAGGAAATACAATGGGCAAAAATAAAAAATTAAAAACGTCAGGTGAGGTAGCAAATGAGATGGGAA 800

801 GAAATGGGCTACATTTTCTAAATATAGAATATTACGAAAAATACAGCGAAACCTGTACTTTGAAGGAGGATTTAGCAGTAAAAGGGGAATAGAGAGCCC 900

901 CTCTGAAACCGGCCCTGAAATGCGCACACACCGCCCGTCACTCTCCTCACCCAAAATCGGCCCATCTTTTAATAAACAAAAACCAAGCATACAAGTAG 1000

tRNA-Val→

1001 AGGAGGCAAGTCGTAACAAGTAAAGTACCGGAAGTGCACCTTGACTAATCAAAATGTAGTTTAAACAAACACCTCCCTTACACTGAGAAGACACCCA 1100

16S rRNA→

1101 CGAAAGCGGGTCACTTTTGGAGCTATATAGCTAGGCCAACAAACCATAACCACTACACCTTAACCAAAAACCTTGAACAAACAAAACCTAAACCATTTACC 1200

1201 ATCCAAGTATAGGCGATAGAAAAGACACCAGGCACAATAGTAAAAGTACCGCAAGGGAACGCTGAAAAAGAAATGAAACAACCTCGTAAAAGCAATAAAAA 1300

1301 CCAAGACTAACCCCTGTACCTTTTGCATTATGATCTAGTTAGACCCACCGGGCAAAATGAATTTAAGTCCAACCCCGAAACTAAGTGAGCTACTTC 1400

1401 GAAACAGCCTATGAGGGCAACCCCTCTCTGTGGCAAAAGAGTGGGAAGATTCCAAGTAGAGGCGATAAACCTAACGAGCTTAGTGATAGCTGGTTATT 1500

1501 CGAGAAATGATTTTGTAGTTCAGCCCTCGGCCTCAAACCCCTAACACATAAAACCAATGTGAGCCAGAGAGATTATCAAAGGGGTACAGCCCTTT 1600

1601 GAAAAAGGACACAACCTTCTAAGACAGGATAAAGACTATATTAACCTCAAGGATCAGCTGCCCAAGTGACCTAAAAGCAGTCACCTAAAAAGCGTTAAA 1700

1701 GCTTAAGCAGCACTAAACCAACCTATACTAATAAATTTAACCTCATTCCACCATCACTATCGAATTATTCTATATGCATAGAAGAATAAATGTAGAATT 1800

1801 AGTAACAAGAAAGGCCATTAAAGCCTTCTTAACCTGCATAAGTGTACATCAGATTAGATTAACCACTGATAATTAACGACTTCAAAGAGAATACTATGACAT 1900

1901 AAAACAAGAAAAGCACACAGCCACATCGTTAATCCAACACAGGAATGCAACCACGAAAGATTAAAAGAAAGAAAAGGAACCTCGGCAAACTATAAGCCC 2000

2001 CGCCTGTTTACCAAAAACATCGCCTCCCGCCAACAACAGAAATATTGGAGGTCGCGCCTGCCAGTGACAAGATTTAACGGCCGCGGTATCCTGACCGTG 2100

2101 CAAAGGTAGCGAAATCACTTGCTTTTAAATGAAGACCTGTATGAATGGCACCACGAGGGCTTAACGTCTCCTCTTTCCAATCAGTAAAATTGATCTGT 2200

2201 CCGTGCAAGCGGACTTAACCTACATTAGACGAGAAGACCTGTGGAGCTTCAGACACAAAGCCAACCTACAATAAACACTTAGATAATAAATGATAAA 2300

2301 ACCAGTAGCCATAACTGGCCCTATGTCTTTGGTTGGGGCGACCAGGAGAAAAACAATCCTCCAAGCGATTGGTACCACCTGTACTAAAACAAGG 2400

2401 GTAACACCCCAAAGTAATAAAAATTTATCGGACATGACCCAGGACTAAACCTGATCAACGAACCAAGTTACCCAGGGATAACAGCGCAATCCTTTCC 2500

2501 AAGAGTCCAATCGACGAAAGGGTTTACGACCTCGATGTTGGATCAGACACCCCAATGGTGAAGCCGCTATTAAGGTTTGGTTGTTCAACAATTAAG 2600

2601 TCCTACGTGATCTGAGTTCAGACCGGAGAATCCAGGTCAGTTTCTATCTATGATGTTAATTTCTCCAGTACGAAAGGACCGGAGAAGCTTGAGCCAATGC 2700

tRNA-Leu (UUR) →

2701 CACAAGCACGCTCATCTTCAACCTGATGAAAAACAATAAAACAGATAAAGAAGAACACAACCCGCTCCCGTAAACAACGGGAATCGCCGGGTGGCAGAGC 2800

NADH 1→

M T K I I T H L L N P L A V

2801 CTGGCTAAATGCAAGAGGCCTAAACTCTCTCTCTCAGAAGTTCAAACTCTCTCCCGAGCTATGACAAAAATTATTACACACCTGCTAAACCCACTAGCTG 2900

I I P I L L A V A F L T L I E R K V L G Y M Q L R K G P N I V G P

2901 TCATCATTTCAATTTTACTAGCCGTAGCGTTCCTGACACTCATCGAACGAAAAGTACTAGGTTATATGCAACTACGAAAGGGTCTAATAATCGTAGGCC 3000

Y G L L Q P L A D G L K L F I K E P V R P S T S S P L L F I T T P

3001 ATATGGTCTCTACAACCCCTAGCAGATGGACTAAAACCTATTCATTAAGGAGCCAGTACGACCTTCTACATCATCCCGCTACTTCTTACCCTCCA 3100

M L A L T M A L T L W L P L P L P H P M T N L N L G M L F I L A I S

3101 ATACTAGCACTCACCATAGCACTAATCTTACTGACTACCACTACCCCTCCACATCCAATAACAACCTAAACCTAGGAATATTATTTATCCTAGCAATCT 3200

S L T V Y S I L G S G W A S N L K Y A L I G A L R A V A Q T I S Y

3201 CAAGTCTAACAGTATATCAATTTAGGCTCCGGCTGAGCATCAAACCTAAAATATGCCCTAATTTGGGGCCCTCCGAGCAGTCGACACAACAATCTCCTA 3300

E V S L G L I L L A M I I F A G G F T L T T F N T S Q E T I W L L

3301 TGAAGTAAGCCTAGGACTTATCCTACTGGCCATAATCATCTTCGAGGCGGTTTACACTAACAAACATTTAATACATCAGGAAACCATTTGACTTCTA 3400

T P G W P L A A M W Y I S T L A E T N R A P F D L T E G E S E L V S

3401 ACGCCAGGATGACCACTCGCAGCAATATGATACATCTAACCTTAGCAGAAACCAACCGAGCCCATTTGACCTCACAGAAGGAGAATCAGAAGCTTGAT 3500

G F N V E Y A G G P F A L F F L A E Y A N I L L M N T L S T I L F

3501 CGGGGTTAATGTGGAATATGCAGGAGGACATTTGCACATTTCTTTCTAGCAGAATATGCAAAATATTTACTAATAAACACACTATCAACTATTCTATT 3600

M G A M H N P I T P E L T S I N L M I K A S A L S M L F L W V R A

3601 TATAGGAGCCATACACAACCAATCACACCAGAAGTAACTCAATTAACCTAATGATTAAGCCCTCCGCACTATCAATACTCTTCTGATGAGTACGAGCC 3700

FIGURE 2.—Complete L-strand nucleotide sequence of the coelacanth mitochondrial genome. Position 1 corresponds to the first nucleotide of the *tRNA<sup>Phe</sup>* gene. Direction of transcription for each gene is represented by arrows. The deduced amino acid sequence for each gene product is shown above the nucleotide sequence (one-letter amino acid abbreviation is placed above the first nucleotide of each codon). Complete termination codons are indicated (\*). tRNA genes are underlined and the corresponding anticodons are overlined. In the control region, four repeats are shown.

S Y P R F R Y D Q L M H L V W K N F L P I T L A M I L W H T S L P I  
 3701 TCATACCCACGATTCCGATATGACCAGCTAATACACTTAGTATGAAAAAATTCTTACCAATCACCTAGCCATGATCTATGACACACCTCCCTCCAA 3800  
 F T G S L P P Q T \* tRNA-Ile→  
 3801 TTTTACAGGAAGCTTACCACCACAACCTAAAGGAAACGTGCCGAATAAACAGGACCCTTTGATAGAGTGGACTATAGGAGTTAAACCTCCTCGC 3900  
 ←tRNA-Gln  
 tRNA-Met→  
 3901 TTCCTAGGAAAATAGGACTTGAACCTTACTAAAGAGATCABAACCTCTTGGTGCTACCACTACACCATCTCCTAGTAAAGTCAGCTAAAAAGCTTTCGG 4000  
 NADH 2→  
 M S P Y V T M I L I S S L G L G T T I  
 4001 GCCCATACCCGAAACATGTTGGTTAAACCTCCTCTTTACTATGAGCCCTTACGTAACAATAATCCTTATCTCAAGCCTTGGACTCGGGACAACAATT 4100  
 T F T S S S W L M A W M G L E I N T L A I T P L M V K Q H H P R A T  
 4101 ACATTTACAAGCTCATCTGACTGATAGCTTGAATAGGCTAGAAATTAATACCTTAGCCATCACCCCTAATAGTAAACAACATCACCTCGGGCAA 4200  
 E A T T K Y F L T Q A T A S G L L L F A T L N N A W M T G E W N T  
 4201 CTGAAGCCACAACAAAATTTTCTACCCAGGCAACAGCATCAGGACTGCTATTATTCGCAACCTTAACAACGCTTGGATAACAGGAGAATGAAACAC 4300  
 M E L S N N L S A P M I T M A L A L K M G V A P M H F W L P E V L  
 4301 AATAGAATCAACAATTTATCCGCCCAATAATTACAATAGCCCTCGCACTAAAGATAGGAGTAGCACAATACACTTCTGATTACCAGAAGTGCTA 4400  
 Q G L P L L T G L I L S T W Q K L A P F T L L Y M T S H E L N T T T  
 4401 CAAGGACTCCCCTACTAAGTGGACTTATCTGTGCGACTTGACAAAACTAGCCCCCTTACCCTACTATATATAACATCACATGAATTAACACAACAA 4500  
 M T I L G L T S T I I G G L G L N Q T Q L R K V L A Y S I A H  
 4501 CAATAACAATCCTAGGATGACATCAACAATTATCGGTGGCCTTGGTGGATTAACCAAACTCAACTGCGAAAAGTCTAGCTTACTCATCAATGACAA 4600  
 L G W M V I I I Q Y S K T L A L L N L L L Y I T M T S T A F L T L  
 4601 CCTCGGATGAATAGTTATCATTATCCAATACTCCAAACACTAGCCCTACTAAACCTACTGCTATATATACAATAACATCAACAGCATTITTAACACTT 4700  
 M T L S A T K I N T L S T K W A T T P I A T M T A M L A L L A L G G  
 4701 ATGACTCTATCAGCCACAAAATTAATCCCTATCAACAAAATGAGCAACAACCCCTATCGCAACTATAACTGCAATACTAGCTCTACTAGCATTAGGAG 4800  
 L P P L T G F M P K W L I L Q E L T K Q N L P A L A T L M A L S A  
 4801 GTCTCCACCACCTAACAGGATTTATACAAAATGACTAATTTTACAAGAATTTACCAAGCAAAATTTACCCGCTCTAGCCACACTAATAGCCCTATCAGC 4900  
 L L S L F F Y L R M C H T M T L T I S P N T N N N M I T W R K K P  
 4901 CCTATTAAGCCTCTTCTTACCTACGAATATGCCATACAATAACCCCTTACAATCTCACCAACACAAACAATAACATAATTACATGACGAAAGAACCT 5000  
 G Q K A L P L A M L S I M T L M A L P T T P T M V A I M N \* tRNA-Trp  
 5001 GGCCAGAAAGCACTACCCTAGCCATACTAAGCATCAAAACCTCATAGCACTCCCAACAACCCCAACAATAGTAGCCATCATAAACTAATAAGAGACTT 5100  
 →  
 5101 AGGAATAAACCAAGACCAAGAGCCTTCAAGCCCTCAGCAGGAGTGAAAACCTCTAGTCTCTGGATAAGACCTGTAGGATACCTACCACATATTTCTGA 5200  
 ←tRNA-Ala  
 5201 ATGCAACCCAGACACTTTAATTAAGCTAAGGCCTCAGCTAGATGGGTAGGCCTCGATCCTACAAAATATTAGTTAACAGCTAAAAGCCCAAACCGGGCAG 5300  
 ←tRNA-Asn ←tRNA-  
 5301 CATCCATCCTAACATTTCCAAGAAAAAAGGAATGTTTGAAGCCTCAACAAGCACCCTTGTCCCTCCAGGTTTGCACCTGACATGCTAAACACCATGAG 5400  
 COI→  
 Cys ←tRNA-Tyr M M I T R W L F  
 5401 GCTTGGTAAGAGGAGGAATTGAACCTCCCTACACGGGGCTACACACCCCGCCTAAGAGCAATCAGCCATCTTACCTGTGATAATCACTCGTTGACTATTC 5500  
 S T N H K D I G T L Y M I F G A W A G M V G T A L S L I R A E L S  
 5501 TCAACCAACCATAAAGACATTGGTACCCTATACATGATCTTGGTGCCTGAGTGGAATAGTTGGAACCGCCCTAAGCCTGCTTATTGAGCTGAACTCA 5600  
 Q P G A L L G D D Q I Y N V V V T A H A F V M I F F M V M P I M I  
 5601 GCCAACCTGGGGCTCTCCTGGGGATGACCAAAATTTAATGATAGTGCCTACAGCAGATGATTGCTGATAATCTTCTTTATAGTAATACCGATCATAAT 5700  
 G G F G N W L I P L M I G A P D M A F P R M N N M S F W L L P P S  
 5701 CGGCGGTTTGGCAACTGATTAATCCCTGATGATTTGGGACCCGACATAGCATTTCCACGTATAAACAACATAAGCTTCTGACTACTACCACCTCA 5800  
 L L L L L A S S G V E A G A G T G W T V Y P P L A G N L A H A G A S  
 5801 CTCCTACTCTACTAGCATCTTCTGGAGTAGAAGCAGGAGCAGGCACAGGATGGACAGTATACCCTCCACTAGCGGGCAACCTCGCCCATGAGGAGCAT 5900  
 V D L T I F S L H L A G V S S I L G A I N F I T T V I N M K P P T  
 5901 CCGTAGATTTAAACAATTTCTCCTTACATCTAGCCGGTGTATCCTCAATCTTAGGGGCCATCAACTTATCAACAACAGTAATCAACATAAAACCCCAAC 6000  
 M T Q Y Q T P L F I W S V L V T A V L L L S L P V L A G G I T M  
 6001 AATAACACAGTATCAGACACCCTATTTATCTGATCAGTCTTAGTGACCCGCTACTCTCTACTCTCGCTACCGGTGCTAGCTGCCGGAATTACCATA 6100  
 L L T D R N L N T T F F D P A G G G D P I L Y Q H L F W F F G H P E  
 6101 CTACTGACAGATCGAAATCTAAACACAACATTTCTTGACCCTGCTGGAGGAGAGACCCCTATTCTATACCAACCTATTCTGATTCTCGGCCATCTGT 6200  
 V Y I L I L P G F G M I S H I V A Y Y S G K K E P F G Y M G M V W  
 6201 AAGTATACCTTAATTTACCAGGATTTGGTATAATCTCACATTTGTGGCTACTACTCTGGAAAGAAAGAACCATTCGGGTATATAGGTATAGTATG 6300  
 A M M A I G L L G F I V W A H H M F T V G M D V D T R A Y F T S A  
 6301 AGCTATAATGGCAATGGACTTCTAGGCTTATCGTATGAGCCATCATATATTTACCGTAGGAATGGATGTTGACACACGAGCATACTTTACATCAGCA 6400  
 T M I I A I P T G V K V F S W L A T L H G G V T K W D T P L L W A L  
 6401 ACCATAATTATTGCCATCCCAACAGGAGTAAAGTGTTCAGCTGACTAGCGACACTTCACGGAGGAGTGACCAATGAGACACACCCCTGCTATGAGCAC 6500  
 G F I F L F T V G G L T G I V L A N S S L D I I L H D T Y Y V V A  
 6501 TAGGATTTATCTTTCTTTTACAGTAGGAGGCCTAACAGGCATCGTACTGGCAAACTCATCACTAGACATCCTCATACATGACACTATTACGCTAGTAGC 6600  
 H F H Y V L S M G A V F A I M G G L V H W F P L M T G Y T L H N T  
 6601 ACACTTCCACTATGCTCATCAATAGGAGAGTATTTGCAATCATAGGGGACTCGTGCACTGATTTCCACTAATAACAGGATATACCTTACACAACACA 6700

FIGURE 2.—Continued

W T K I H F G V M F T G V N L T F F P Q H F L G L A G M P R R Y S D  
 6701 TGAACAAAATCCACTTTGGTGAATATTCACAGGAGTAAACCTAACATTTTTCCCAACACTTCTCGGACTAGCAGGAATACCACGACGTTACTCAG 6800  
 Y P D A Y T L W N T V S S I G S L I S L I A V I M F M F I L W E A  
 6801 ACTATCCAGATGCCTATACTTTATGAACACAGTATCATCAATGGCTCTCTAATTTCACTAATGCGGTAATCATATTTATTTCTGTGAGAAGC 6900  
 F S A K R E V L I V E M T T T N V E W L H G C P P P H H T Y E E P  
 6901 TTTCTCTGCCAACGAGAAGTACTAATTTGTAGAATAACAACAACAATGTAGAATGGCTGCACGGATGCCACCACCACACCACACATGAAGAACCA 7000  
 A F V Q A P R \* ←tRNA-Ser (UCN)  
 7001 GCATTGCTACAAGTCTCTCGATAAAACACTGAGAAAAGAGGGAATCGAACCCCGTCAGCTAATTTCAAGACAGCCGCATCACCAGCTCTGCCATTTTCTT 7100  
 COII→  
 tRNA-Asp→ M A H P S Q L  
 7101 TATTAAGATTCTAGTAAAACAAATTACACTACCTTGTCAAGATAGAATTATGAGTGAAAACCTCATGAACCTTGACCAATGGCACCCCATCACAGTTA 7200  
 G L Q D A A S P V M E E L L H F H D H A L M I V F L I S T L V F Y I  
 7201 GGATTACAAGATGCAGCTTCTCCCGTTATAGAAGAAGTCCCTCCACTTTCACGATCATGCACTAATAATTTGATTTTTAATTAGCACATTAGTATTTTACA 7300  
 I L A M M T T K M T D K Y I L D A Q E I E I V W T L L P A I V L I  
 7301 TTATTCTAGCCATAATAACAACAAAATAACTGACAATATATCTTAGACGCACAAGAATGAAATTTGTGGAACACTACTCCCAGCAATCGTCTAAT 7400  
 L V A L P S L R I L Y L I D E V E N P H L T I K A M G H Q W Y W S  
 7401 CCTAGTTGCCCTACCCTCGCTACGAATCCTATATCTAATTTGATGAAGTCGAAAACCCCTCACCTAACAAATTAAGCAATAGGCCACCAATGATACTGAAGC 7500  
 Y E Y T D Y E E L S F D S Y M T P L Q D L N P G Q F R L L E T D H R  
 7501 TATGAGTACACGGACTATGAAGAATAAGCTTCGACTCATACATAACACCCTACAAGACCTAAACCCGGGCAATTCGGCTGTGGAACAGACCATC 7600  
 M V I P M E S L I R V L I S A E D V L H S W A V P A L G V K M D A  
 7601 GAATGTTATCCCAATAGAGTCGCTTATCCGAGTACTAATTTCAAGCTGAAGACGCTACTACACTCATGAGCAGTCCCAGCCCTAGGAGTAAAATAGATGC 7700  
 V P C G R G L N Q I T F M I S R P G L Y Y G Q C S E I C G A N H S F M  
 7701 AGTCCAGGCGACTCAACCAATTACATTCATTTCCCGCAGGACTATATTAGGACAATGCTCAGAGATTTGTGGAGCAACACACGCTTTATA 7800  
 P I V L E A I P L D P F E D W S S S M L E E A tRNA-Lys→  
 7801 CCCATCGTACTTGAAGCAATCCCACTAGACCCCTTCGAGACTGATCTTCATCAATGTGGAAGAAGCCTCAGTGAAGCTAAATAGAAAGCGATAGCC 7900  
 ATPase 8→  
 M P Q L N P S P W L L I L L F S W L I  
 7901 TTTTAAGCTAGAGACTGGTGAAAAACAAACCCACCCCTCAGTGCATGCCACAATAAACCCCTCCCCCTGACTACTAATCCTGCTATTTCTCTGACTCATC 8000  
 F L T M L P S K T Q L H T F P N M P S T Q N M C K Q E P E P W T W P  
 8001 TTCTTAATATACTCCCTCTAAGACACAATTACACACCTTCCCAACATGCCATCAACACAAAATATATGCAACAAGAACCAGAACCATGAACCTGAC 8100  
 ATPase 6→  
 M S L N F F D Q F M S P T L L G V P L I A V A M M F P W T L L P T  
 W A \*  
 8101 CATGAGCCTAAACTTCTTTGACCAATTTATGAGCCCAACTATTAGGAGTACCCTCATTTGCTGTAGCAATAATATCCCATGGACCCTATTACCAACA 8200  
 P T N R W L N N R T L T L Q N W F I G R F T N Q L Q P L N T G G G H  
 8201 CCAACCAACCGATGACTTAATAACCGAACACTAACACTACAATACTGATTTATCGGGCGCTCCTAATCAACTACTACAACCAATTAACACJGGAGGAC 8300  
 K W A M I L M S L N L G L L P Y T F T P T T Q L S L N M G L A I  
 8301 ACAAATGAGCAATAATCTTAATATCACTAAACCTCTGGGACTTCTACCGTATACATTACACCAACAACAACAATATCACTAAACATGGGACTTGTCTAT 8400  
 P F W L A T V L L G L R N Q P T A A L G H L L P E G T P T L L I P  
 8401 TCCATTCTGACTAGCAACAGTATTACTGGGACTGCGTAACCAACCCACTGCGCGCTAGGACACCTTCTCCAGAAGGAACCAACCCCTGCTAATCCCA 8500  
 I L I I E T I S L I R P A A C C G V L F W L F T A N L T A N L T A P T P E L G  
 8501 ATCCTAATTTATTAAGCAATACGCCTACTTATCCGCCCTTCGCGCTAGGAGTACGACTAACAGCCAAATCTCACAGCAGGCCACCTCCTAATAACAAT 8600  
 I A T A A F V L L P M M P T V A L L T T L V L F L L T L L E I A V  
 8601 TAATTGCTACCGCGCCTTCTGACTCTACCTATAATACCAACAGTAGCATTATTAACAACATTAGTCCTATTCTCTGACCCCTGCTAGAAATGGCCGT 8700  
 COIII→  
 A M I Q A Y V F V L L L S L Y L Q E N V M A H Q A H A Y H M V D P  
 8701 AGCAATAATCAAGCCTACGTGTTTGTCTATTACTAAGCCTCTATCTACAAGAAAATGTCTAATGGCCACCAAGCACAGCATATCACATAGTTGACC 8800  
 S P W P I T G A T A A L L V T S G L A A W F H F N S M I L I L M G  
 8801 CAAGCCCATGACCCATTACAGGGGCCAGGGCCCTACTTGTAACTCAGGCTAGCAGCGTGATTTCACTTCACTCAATAATCTTAATTTTAAATAGG 8900  
 L T L L L L T M Y Q W W R D I I R E S T F Q G H H T L P V Q K S L  
 8901 ACTAACACTATTGCTACTAATATGATCAATGATGACGAGATATTATTCGAGAAAGCACATTCCAAGGTCACCACACACTACCTGTACAAAAAGCCTA 9000  
 R Y G M I L F I T S E V F F L G F F W L F A P Y H S L A P T P E L G  
 9001 CGATATGGTATAATCCTGTTTCAATTACATCCGAAGTATTCTTCTCCTAGGGTCTTCTGAGCCTTTTACCATTCAAGTCTGGCACCCACTCCTGAACTCG 9100  
 G L W P P T G I T P L D P F E V P L L N T A V L L A S G I T V T W  
 9101 GAGGACTCTGACCTCCCACTGGAATTACACCCTAGATCCATTTGAAGTACCCTATTAACACACAGCAGTTCTACTAGCCTGGGAATTACAGTCACATG 9200  
 A H H S L M E G Q R K E A I Q S L F I T V L L G L Y F T A L Q A T  
 9201 AGCCCATCACGCCTAATAGAGGGCAACGAAAAGAGGCTATCCAATCACTATTTATCAGACTTCTGTTAGGACTAATTTACAGCACTGCAAGCCACA 9300  
 E Y Y E S P F T I A D G A Y G S T F F V A T G F H G L H V I I G S T  
 9301 GAATACTACGAATCCCAATTTACAATCGCTGACGGAGCCATGGCTCAACCTTTTTGTAGCAACCGGATTCACGGTCTACATGTCATTATGGCTCTA 9400  
 F L I V C L V R Q T Q Y H F T S N H H F G F E A A A W Y W H F V D  
 9401 CATTCTAATCGTATGCTAGTACGACAAACAATACCCTTACATCAAACCACCTTTGGCTTGAAGCAGCAGCATGATACTGACATTTCTGTAGA 9500  
 V V W L F L Y V S I Y W W G S \* tRNA-Gly→  
 9501 CGTAGTCTGACTATTCTTATACGTATCAATCTACTGATGAGGCTCATAAACCCCTTGTAGTACAAAATAATACAAATGACTTCCAATCATTTAATCTGGT 9600

FIGURE 2.—Continued

NADH 3 →

M N L I L A G L L I M S I L S M I L A M I A F W L P N

9601 TATACCCCAAGAATGGGTAATGAACCTGATTCTAGCGGGCTACTTATCATAAGCATCCTCTCTATAATTTTAGCTATAATCGCATTCTGACTACCAAAAC 9700
M T P D T E K L S P Y E C G F D P L G S A R L P F S L R F F L V A I
9701 ATGACCCCTGATACAGAAAACCTATCTCCCTACGAATGTGGCTTTGATCCTCTAGGATCCGCACGACTCCCATTCCTCCCTACGATTTTCCCTAGTAGCAA 9800
L F L L F D L E I A L L L P L P W A D Q L T N P T L A L T W T T S
9801 TCCTATTCCTGCTATTTGACCTAGAAAATGCATTATTATACCCTACCTGGGCAGACCAACTAACAAACCCAACTTGCATTAACTGGACAACAAG 9900
I I A L L T L G L I H E W T Q G G L E W A E tRNA-Arg →
9901 CATCATCGCCCTACTAACACTAGGACTAATCCACGAATGAACCTAAGGAGGCCTCGAATGGGCAGAATAGGTAGTTAGTTTAAAAAACCACTAATTTTCG 10000

NADH 4L →

M T P V Q L S F N T A F T L G L M G V T F

10001 ACTTAGTTAACTGTGGTGAACCCCACTATCTTTCATGACCCGACTACAACCTCAGCTTTAACACTGCATTACACTAGGCTTAATAGGAGTAACATTC 10100
H R A H L L S A L L C L E G M M L S L Y M G L S L W P M Q L E S T T
10101 CACCGAGCCCATCTGCTATCAGCATTACTCTGCCTAGAAGGAATAATATTATCCCTGTATATAGGACTGTCCTATGACCAATGCAACTAGAATCAACTA 10200
Y M T T P L L L L A F S A C E A G A G L A L M V A T S R T H G T D
10201 CATAATAACCCACACCCTACTACTCTGCCTTCTCAGCCTGTGAGGCTGGAGCAGGCCTAGCCCTCATAGTAGCAACATCCCGCACACATGGTACGGA 10300

NADH 4 →

M L K V L M P T I M L I L T T W L T K P A W L W P

H L Q N L N L L Q C \*

10301 CCACCTCCAAAACCTAACTTACTACAATGCTAAAAGTTTAAATACCAACAATTATGCTTATCTTAACCACATGATTAACAAAACCTGCATGACTCTGAC 10400
T M T T N S L L V A T I S L T W L K W D S E S G W K S L N S S M A
10401 CAACAATAACAACCAATAGCCTACTCGTAGCTACCTCAGCTTAACTGACTAAAATGGGACTCAGAGTCAGGATGAAAATCTCTCAACAGCTCAATGGC 10500
T D P L S T P L L I L T C W L L P L M I L A S Q N H M F M E P L N
10501 TACCGACCCCTATCTACACCATTACTAATCTCACATGCTGGCTTCTACCCCTCATAATTTCTCGAAGCCAAAACCATGTTTATAGAACCCTAAAC 10600
R Q R S F I S L L I S L Q T F L I M A F G A T E I I L F Y I M F E A
10601 CGCAACGATCATTTCCTCTCCCTACTCTCCCTACAACATTCCTAATTATAGCATTGGTGCCACTGAAATCATCTATTTTACATTATATTTGAAG 10700
T L I P T L I I I T R W G N Q T E R L N A G T Y F L Y T V M G S
10701 CAACCCCTAACTCCAACTAATTATTATACCAGTGGGTAATCAAACAGCGACTAAACGCAGGAACATACTTTTTTATATACAGTAATAGGGTC 10800
L P L L V A L L M T Q N N L G T L S M P L I Q H M Y Q M K L H T H
10801 ACTACCCTATTAGTTGCACTTTTAAACACAAAATAACCTTGGTACCCTATCAATACCGCTCATCAACACATATACCAAATAAAACTCATACACAT 10900
G D M M W W T A C L L A F L V K M P L Y G V H L W L P K A H V E A P
10901 GGAGACATGATATGATGAACAGCCTGCCTATTAGCCTTCTAGTAAAAATACCCTATACGGAGTCCACCTTGACTCCCAAAGCCCATGTAGAAGCCC 11000
I A G S M V L A A V L L K L G G Y G M M R L I M M L A P M T K T L
11001 CAATTGAGGATCAATAGTACTAGCCGCTCTACTAAAACCTAGGAGGATACCGAATAATACGACTAATCATAATATTAGCTCCAATAACAAAACCCCT 11100
A Y P F I I L A L W G I I M T G S I C L R Q T D L K S L I A Y S S
11101 AGCCTATCATTATCATCTCTCGCCCTATGAGGAATCATTATAACTGGATCAATCTGCTTACGACAAAACAGACCTAAAATCCCTAATCGCCTACTCATCA 11200
V G H M G L V A A G I L T Q T P W G F T G A T V L M I A H G L T S S
11201 GTAGGCCACATAGGACTAGTGGCAGCAGGTATCCTAACACAAACACCATGAGGCTTTACAGGAGCTACTGTTCTAATAATGGCTACGGCTTACATCCT 11300
A L F C L A N T N Y E R T H S R T M I L A R G M Q V I L P L M T F
11301 CAGCCCTATTCTGTCTAGCAACACAACTATGAACGAAACCCATAGCCGAACCATGATCCTAGCAGGAGGATGCAAGTTATCCTCCCACTCATGACATT 11400
W W L M M L A N L A L P P S T N L M G E L L I I T M T F N W S N
11401 CTGACTGACTATAATAAATTTAGCTAAGCTTACCCCTACCCTAATCTAATAGGAGAACTACTTATTATTAACTTCAACTCAACTCAAAAC 11500
W T L T M T G L G M L I T A I Y S L H M F L T T Q R G L M T N H I I
11501 TGACACTAACTATAACAGGACTGGGCATACTAATCACAGCTATCTACTATTACACATGTTCCCTCACACACAACGAGGCCCTGATAACAACACACATTA 11600
S I E P S H T R E H L L M H A L P M L L L I L K P E L I W G W
11601 TCTCAATTGAACCCCTCCACAGCAGCCTACTAATAACAATACATGCCCTCCCAACTACTATTGCTAATCCTTAAACGAGAAGTATGAGGCTG 11700
S Y tRNA-His → tRNA-Ser (AGY) →
11701 ATCCTACTGTAATATAGTTTGAIAAAAACATTAGACTGTGGCTCTAAAATAAGAGTTAAATCTCTTTATTAACCGAGAGGGGCTAGAGCCTACGAA 11800

tRNA-Leu (CUN) →

11801 CTGCTAATCTGTAGTACCATGGTTCAATCCATGGCCACTCAGCCCTAAAGGATAATAGCTCATCCATTGGTCTTAGGAGCCAAAACCTCTTGGTGC 11900

NADH 5 →

M Y T T L I F N S T L M T M F T I L T A P I L T T L N P

11901 AACTCCAAGTAGCGCCATGTACACAACATTAATTTTAACTCAACCTTATAACCATATTTACCATCCTAACAGCCCAATCTTAACCACCCTTAACCC 12000
I K P N K K W T E L W V K T A V Q L A F F T S L I P L F I Y L D Q
12001 TATCAAACCCAAACAAAAGTGAACAGAATTATGAGTAAAAACCCGCTACAGCTAGCCTTCTTACAAGTCTAATCCCGCTCTTCTATCTTACACAA 12100
G I E T I T T N W Q W M N T N T F N I N I S F K F D Q Y S I V F I P
12101 GGAATCCGAGCATTACACAACCTGACAATGAATAAACACCAACACATTAACATTAACATCAGCTTAAATTTGACCAATACTCAATGTATTTCATCC 12200
I A L Y V T W S I L E F A N W Y M H Q D P K M N Q F F K Y L L L F
12201 CAATCGCATGTATGTCACATGCAATCTAGAACTCGCCAAGTATACAGCAACCAAAATAACCAATTTTCAAATACCTGCTCTCTCTATT 12300
L I T M M I L V T A N N M F Q L F I G W E G V G I M S F L I G W
12301 CCTAATCACCATAATAATCTTAGTTACAGCAAAACATGTTCCAATATTTATCGGCTGAGAAGGAGTAGGAATTATATCATTCTTAAATGGCTGA 12400
W H G R A N A N T A A L Q A V I Y N R V G D I G L T L S M V W F A I
12401 TGACATGGCCGAGCTAACGCCAATACCGCAGCCCTACAAGCAGTCTACTACAATGAGTAGGAGACATTGGGTTAACCCCTAAGCATGGTCTGATTTGCCAA 12500
N L N T W E M Q M F I M S Y N T D M T I P L L G L T L A A A G K
12501 TCAACCTAAATACATGAGAAATACAACAATATTTATCATATCTACAACACCCGACATAACCATCCCCCTACTAGGCCTAACTCTGGCAGCAGCAGGAAA 12600

FIGURE 2.—Continued



15701	CGTTACTCTCAAGTACATTATACATTACATGTTAATTTACCATTAGAACTGTCACACCACATTTAGGGCGAGAAACCAGCAACTTGCATAAATGATTA	15800
15801	ATTAATTTACGATGTGGTTAGACACTTATTTCTTAATCTAAACTTGGCTTTATCCATTACTGGCCACTGGTACTGTGCGATGGAGAATAATAGAAAGATT	15900
15901	TATATGATTATAAATTATCTATTACTGGCATCTGGTTTTGGGTTTAGTGAGGGGAAGGGCTTTTAAACCCGTAACCTCAGTATCACTTTACTATACTGGCC	16000
16001	AGCCGTTATTTTTTTGGGGGGGGGAACGTGAAGTTAGACAGCACTTAACCTTGCCTCAACCGTCGTTTATGATTGGACTTTTAGTCGCACTCAAGTACT	16100
	<hr style="width: 100%; border: 0.5px solid black;"/> <div style="display: flex; justify-content: space-around; font-size: small;"> <span>Repeat 1</span> <span>/</span> <span>Repeat 2</span> <span>/</span> <span>Repeat 3</span> </div> <hr style="width: 100%; border: 0.5px solid black;"/>	
16101	TTTGGATCTATGACAAAGGATAATCAGTTAATGATAGATAGACATATAATGAATGATAGATAGACATATAATGAATGATAAATAGACATATAATGAATGA	16200
	<hr style="width: 100%; border: 0.5px solid black;"/> <div style="display: flex; justify-content: center; font-size: small;"> <span>Repeat 4</span> </div> <hr style="width: 100%; border: 0.5px solid black;"/>	
16201	TAGTTAGATATATAGATTAATGTAAGAAAGATAAATGAACCCATGACAGAGGACATACTTTTAATGATTTCCAGGACATAAACCCATGCACATCAGATTG	16300
16301	TATCAACTAAACCCCTCCCAACCCCTAAAACCCAGGACTCGCTAAACACATCAACCCATATATTTTTTCAGTATATACATTGTTATACATCAAATAAATA	16400
16401	ATGTAAT	16407

FIGURE 2.—Continued

phological and molecular characters that could have evolved at the time of this rapid radiation have been obliterated by subsequent changes and back mutations during the last 400 million years. Based on the currently available data, it had remained difficult to distinguish between competing phylogenetic hypotheses, and larger molecular data sets are needed to address this important phylogenetic issue.

Mitochondrial DNA has been widely and successfully used in the past to infer phylogenetic relationships among many different species and might therefore be expected to be a suitable candidate for the evolutionary question at hand. However, several authors (*e.g.*, CUMMINGS *et al.* 1995; RUSSO *et al.* 1996; ZARDOYA and MEYER 1996c) recently pointed out the high risk of not recovering the exact evolutionary relationships of the taxa under study when individual mitochondrial genes are analyzed and have suggested that larger or more heterogeneous data sets, *i.e.*, complete mitochondrial genomes, are needed to confidently resolve some phylogenetic questions. To address the question of the relationships among living sarcopterygians and to study the evolution of the mitochondrial genome in vertebrates, we previously sequenced the complete mitochondrial genome of an African lungfish (ZARDOYA and MEYER 1996a). Here, we present the complete nucleotide sequence of the mitochondrial genome of the coelacanth, *L. chalumnae*.

#### MATERIALS AND METHODS

**DNA extraction, PCR amplification, cloning and sequencing:** The scarcity of coelacanth samples (BRUTON and COU-TOUVIDIS 1991; FRICKE 1992; FRICKE *et al.* 1995) and their generally poor quality did not allow us to isolate mitochondria directly from tissue and to obtain intact mitochondrial genomes. Therefore, a total DNA extraction was performed (TOWNER 1991) and the isolated DNA was cleaned through a Sephadex G-50 column. A combination of 24 sets of versatile primers was designed (Table 1) based on highly conserved vertebrate mtDNA regions. They were used to amplify via PCR, contiguous and overlapping fragments (averaging ~800 bp) which covered the entire coelacanth mtDNA molecule (Figure 1). Some of these primers are expected to successfully amplify mitochondrial DNA fragments in other related vertebrate species (*e.g.*, the primer pairs 19–20 and 27–28 of Table

1 have successfully amplified in fishes and amphibians, unpublished data). Thirty-five cycles of PCR (denaturing at 94° for 60 sec, annealing at 45–50° for 60 sec and extending at 72° for 60–105 sec) were performed in 25- $\mu$ l reactions containing 67 mM Tris-HCl, pH 8.3, 1.5 mM MgCl<sub>2</sub>, 0.4 mM of each dNTP, 2.5  $\mu$ M of each primer, template DNA (10–100 ng), and AmpliTaq DNA polymerase (1 unit, Perkin-Elmer-Cetus).

PCR products were cloned using the pGEM-T vector (Promega) and sequenced using the FS-Taq Dye Deoxy Terminator cycle-sequencing kit (Applied Biosystems Inc.) with an automated DNA sequencer (Applied Biosystems 373A) as previously described (ZARDOYA and MEYER 1996a). DNA sequences were obtained using both M13 universal sequencing primers, and three specific oligonucleotide primers for those clones containing inserts >1000 bp (see Table 1). Typically, only one clone per PCR product was sequenced. No differences were found in any of the overlapping sequences between contiguous clones. The fidelity of Taq polymerase is  $\sim 5 \times 10^{-6}$  errors per nucleotide incorporated per cycle (GELFAND and WHITE 1990) (one error in 16,400 nucleotides after 35 cycles). However, recently XU and ARNASON (1996) have reported that the sequence of the gorilla mitochondrial genome obtained by direct isolation from mitochondria differed outside the control region by 49 nucleotides with respect to the corresponding mitochondrial genome obtained partially by PCR (HORAI *et al.*, 1995). Obviously, not all these differences are due to Taq polymerase error during amplification but also are due to sequencing errors, so we expect in a more realistic estimate that our sequence might differ maximally (outside the control region) by <20 nucleotides with respect to the hypothetical sequence from a genome obtained by direct isolation from mitochondria. Based on these calculations, we can submit that the sequence reported is representative of the coelacanth mitochondrial genome and that, although we recommend traditional methods for obtaining mitochondrial genomes if possible, our PCR approach to obtain the complete mitochondrial genome sequence will be useful for other rare or endangered species.

**Molecular and phylogenetic analyses:** Sequence data were analyzed with the GCG program package (DEVEREUX *et al.* 1984), MacClade version 3.06 (MADDISON and MADDISON 1992) and PAUP\* version d54 (SWOFFORD 1997). DNA sequences were aligned using CLUSTAL W (THOMPSON *et al.* 1994) followed by refinement by eye. Ambiguous alignments, mainly in 5' and 3' ends of protein-coding genes, in tRNA gene sequences corresponding to the tRNA DHU and T $\psi$ C arms, and in several highly variable regions of the rRNA genes, were excluded from the phylogenetic analyses (aligned sequences are available from the authors upon request). Third codon positions were also excluded from all phylogenetic analyses. Transitions and transversions were given equal weight

**TABLE 2**  
**Localization of features in the mitochondrial genome of the coelacanth**

Feature	From	To	Size (bp)	Codon	
				Start	Stop
tRNA-Phe	1	69	69		
12S rRNA	70	1052	983		
tRNA-Val	1053	1119	67		
16S rRNA	1120	2784	1665		
tRNA-Leu (UUR)	2785	2860	76		
NADH 1	2861	3832	972	ATG	TAA
tRNA-Ile	3833	3906	74		
tRNA-Gln	3974	3904	69 (L)		
tRNA-Met	3974	4043	70		
NADH 2	4044	5090	1047	ATG	TAA
tRNA-Trp	5093	5164	72		
tRNA-Ala	5235	5167	67 (L)		
tRNA-Asn	5310	5238	71 (L)		
tRNA-Cys	5403	5338	65 (L)		
tRNA-Tyr	5475	5404	69 (L)		
CO I	5477	7024	1548	GTG	TAA
tRNA-Ser (UCN)	7100	7030	69 (L)		
tRNA-Asp	7105	7174	70		
CO II	7180	7870	691	ATG	T--
tRNA-Lys	7871	7943	73		
ATPase 8	7944	8111	168	ATG	TAA
ATPase 6	8102	8763	662	ATG	TA-
CO III	8764	9549	786	ATG	TAA
tRNA-Gly	9550	9619	70		
NADH 3	9620	9968	349	ATG	T--
tRNA-Arg	9969	10,036	68		
NADH 4L	10,038	10,334	297	ATG	TAA
NADH 4	10,328	11,708	1381	ATG	T--
tRNA-His	11,709	11,777	69		
tRNA-Ser (AGY)	11,778	11,844	67		
tRNA-Leu (CUN)	11,845	11,917	73		
NADH 5	11,918	13,753	1836	ATG	TAA
NADH 6	14,270	13,750	519 (L)	ATG	AGA
tRNA-Glu	14,340	14,271	70 (L)		
Cyt <i>b</i>	14,343	15,484	1142	ATG	TA-
tRNA-Thr	15,485	15,556	72		
tRNA-Pro	15,626	15,558	67 (L)		
Control region	15,627	16,407	781		

(for exceptions see below). Overlapping positions (two open reading frames) in several genes (*ATPase8/ATPase6*, *ND4L/ND4*, *ND5/ND6*) were duplicated in the analyses. Gaps resulting from the alignment were treated as missing data. The control region was also excluded from the analyses due to its fast rate of evolution which prevented reliable alignment and made it not appropriate for this phylogenetic question.

Three data sets (protein, rRNA, and tRNA coding genes) were combined and subjected to the maximum-parsimony (MP) method (PAUP version 3.1.1, SWOFFORD 1993; and PAUP\* version D54; SWOFFORD 1997), using heuristic searches (TBR branch swapping; MULPARS option in effect) with 10 random stepwise additions of taxa to find the most parsimonious trees. Neighbor-joining (NJ) (SAITOU and NEI 1987) (based on Kimura-corrected distance matrices, jumble option in effect) and maximum likelihood (ML) (transversions were given double the weight of transitions; empirical base frequencies and five random stepwise addition of taxa were used) analyses of the sequences were performed with

PHYLIP (version 3.55) (F84 model, FELSENSTEIN 1989), MOLPHY version 2.2 (ADACHI and HASEGAWA 1992), and PAUP\* version d54 (SWOFFORD 1997). Robustness of the inferred trees was tested by bootstrapping (FELSENSTEIN 1985) (as implemented in PAUP version 3.1.1., PAUP\*, and PHYLIP with 100 pseudoreplications each).

**Statistical methods:** The statistical confidence of MP analyses was evaluated by calculating the standard deviation of the difference in number of steps between the resulting most parsimonious trees and the two alternative hypotheses using the method of TEMPLETON (1983) as implemented in PHYLIP. Similarly, the statistical confidence of the resulting best tree of the ML analysis with respect to competing hypotheses was assessed by calculating the standard deviation of the difference in log-likelihood between the resulting best tree and the alternative hypotheses using the formula of KISHINO and HASEGAWA (1989) as implemented in PHYLIP, PAUP\*, and MOLPHY. In both cases, competing trees were declared significantly different if the difference in number of steps or

TABLE 3

## Base composition of vertebrate mitochondrial genomes

	A	C	G	T
<b>Proteins</b>				
Tetrapods				
1	30.7	25	21.2	23.1
2	19.3	27.1	12.4	41.2
3	41	31.9	4.8	22.2
Total	30.3	28	12.8	28.9
Lungfish				
1	27.6	25.4	23.6	23.4
2	18.4	27.2	13.3	41.1
3	34.7	28.5	8.4	28.4
Total	26.9	27	15.1	31
Coelacanth				
1	30.2	26	23.2	20.6
2	18.4	27.3	13.3	41
3	47.7	28.3	6.9	17.1
Total	32.4	27.1	14.5	26
Teleosts				
1	26.2	26.6	26.2	21
2	18.5	27.4	13.8	40.3
3	37.8	33.2	8.1	20.9
Total	27.5	29.1	16	27.4
Bichir				
1	29.9	23.7	23	23.4
2	18.9	27	12.8	41.3
3	42.7	27.8	4.4	25.1
Total	30.5	26.2	13.4	29.9
Lamprey				
1	30.4	22.9	22.6	24.1
2	19	26.5	12.9	41.6
3	41.3	21.5	3.8	33.4
Total	30.2	23.7	13.1	33
<b>tRNAs</b>				
Tetrapods	31.9	18.6	19.8	29.7
Lungfish	28.3	20.9	23.6	27.2
Coelacanth	30	20.8	21.8	27.4
Teleosts	28.2	21.3	23.5	27
Bichir	30.7	18.8	20.8	29.7
Lamprey	30.4	18.8	20.4	30.4
<b>rRNAs</b>				
Tetrapods	35.4	23.7	17.7	23.2
Lungfish	33	22.7	20.1	24.2
Coelacanth	36.7	24.9	18.6	19.8
Teleosts	33.4	25.4	21.5	19.7
Bichir	34.5	22	19.9	23.6
Lamprey	36.1	22.6	17.7	23.6

Tetrapods: human, Blue whale, opossum, chicken, and frog; teleosts: Rainbow trout, carp, and loach.

Pag-likelihoods were found to be >1.96 times the standard deviations (FELSENSTEIN 1989).

The complete mtDNA sequence of the coelacanth has been deposited at the EMBL/GenBank data libraries under accession no. U82228.

## RESULTS AND DISCUSSION

**Genome organization and base composition:** The complete L-strand nucleotide sequence of the coela-

canth mitochondrial genome (16,407 bp) is depicted in Figure 2. The organization of the coelacanth mitochondrial genome conforms to the consensus vertebrate mitochondrial gene order (Figures 1 and 2, Table 2). As in other vertebrates, two rRNAs, 22 tRNAs and 13 proteins are encoded by the coelacanth mitochondrial genome. The overall base composition of the L strand is A: 34%; T: 24%; C: 27%; and G: 15%. As in other vertebrate mitochondrial genomes, guanine is the rarest nucleotide whereas adenine is the most frequent (MEYER 1993). A more detailed analysis of the base composition was performed by considering the rRNA, the tRNA and the protein coding genes separately (Table 3). In coelacanth protein-coding genes, there is an anti-G bias in third codon positions and pyrimidines are overrepresented in second codon positions, as was noted before for other vertebrate mitochondrial genomes (NAYLOR *et al.* 1995). Coelacanth tRNAs are A+T rich (57%) whereas the rRNAs have a high adenine content (Table 3). The noncoding intergenic spacer regions (22 bp), which are likely not subjected to strong selection, showed an A+C bias (73%). This indicates that, as is the case in other vertebrate mitochondrial genomes, an asymmetrical directional mutation pressure may be operating in the coelacanth genome (JERMIIN *et al.* 1995).

**Noncoding sequences:** The control region in the coelacanth mitochondrial genome is 781 bp long and 65% A+T rich (Figure 2). The coelacanth mitochondrial control region is characterized by the presence of four 22-bp tandem repeats in the right domain, close to the 3' end. Of these, three are perfect repeats whereas one is imperfect (six nucleotides are different). Even with the presence of these repeats, the coelacanth mitochondrial control region is, so far, the shortest among vertebrates (excluding the atypical control region of the lamprey) (LEE and KOCHER 1995). Neither conserved sequence blocks (CSBs, WALBERG and CLAYTON 1981; SOUTHERN *et al.* 1988; DILLON and WRIGHT 1993) nor termination associated sequences (TASs, DODA *et al.* 1981; FORAN *et al.*, 1988), which are commonly found in other vertebrate mitochondrial control regions, could unambiguously be identified in the coelacanth mitochondrial control region. Putative CSB-II and -III motifs, sharing limited sequence similarity to the human and mouse consensus sequence (WALBERG and CLAYTON 1981), were tentatively identified at positions 16,308 and 16,324, respectively (see Figure 2).

The origin of light strand replication ( $O_L$ ) of the coelacanth mitochondrial genome is, as in most vertebrates, located in a cluster of five tRNA genes (WANCY region) (but see SEUTIN *et al.* 1994) (Figure 2). This region is 26 nucleotides in length and has the potential to fold into a stem-loop secondary structure. The folding of the  $O_L$  does not require the use of part of the adjacent tRNA<sup>CYS</sup> as has been described for the lungfish mitochondrial genome (ZARDOYA and MEYER 1996a).

TABLE 4  
Amino acid composition of vertebrate mitochondrial proteins

	Tetrapods	Lungfish	Coelacanth	Teleost	Bichir	Lamprey	Average
Ala	260 (GCC)	312 (GCC)	286 (GCA)	344 (GCC)	307 (GCC)	297 (GCC)	301 (GCC)
Arg	67 (CGA)	71 (CGA)	73 (CGA)	78 (CGA)	74 (CGA)	68 (CGA)	72 (CGA)
Asn	151 (AAC)	143 (AAY)	147 (AAC)	117 (AAC)	147 (AAC)	143 (AAT)	141 (AAC)
Asp	67 (GAC)	71 (GAC)	71 (GAC)	76 (GAC)	73 (GAC)	67 (GAC)	71 (GAC)
Cys	27 (TGC)	28 (TGY)	25 (TGC)	26 (TGC)	29 (TGC)	40 (TGT)	29 (TGC)
Gln	93 (CAA)	100 (CAA)	103 (CAA)	100 (CAA)	98 (CAA)	98 (CAA)	99 (CAA)
Glu	92 (GAA)	93 (GAA)	102 (GAA)	102 (GAA)	94 (GAA)	93 (GAA)	96 (GAA)
Gly	215 (GGA)	246 (GGA)	239 (GGA)	249 (GGA)	226 (GGA)	213 (GGA)	231 (GGA)
His	100 (CAC)	93 (CAC)	103 (CAC)	105 (CAC)	98 (CAY)	106 (CAC)	101 (CAC)
Ile	333 (ATY)	318 (ATT)	285 (ATC)	281 (ATY)	356 (ATT)	334 (ATT)	318 (ATT)
Leu	626 (CTA)	650 (CTW)	622 (CTA)	637 (CTA)	611 (CTA)	610 (YTA)	626 (CTA)
Lys	93 (AAA)	72 (AAA)	84 (AAA)	70 (AAA)	82 (AAA)	99 (AAA)	83 (AAA)
Met	210 (ATA)	176 (ATA)	224 (ATA)	164 (ATA)	212 (ATA)	217 (ATA)	200 (ATA)
Phe	226 (TTC)	242 (TTT)	214 (TTC)	225 (TTY)	214 (TTY)	235 (TTT)	223 (TTY)
Pro	211 (CCM)	212 (CCA)	206 (CCA)	209 (CCM)	203 (CCA)	202 (CCA)	207 (CCA)
Ser	288 (TCM)	258 (TCM)	226 (TCA)	235 (TCM)	256 (TCA)	263 (TCW)	254 (TCM)
Thr	326 (ACM)	295 (ACA)	359 (ACA)	305 (ACC)	299 (ACA)	306 (ACA)	315 (ACA)
Trp	107 (TGA)	116 (TGA)	121 (TGA)	120 (TGA)	116 (TGA)	109 (TGA)	114 (TGA)
Tyr	125 (TAY)	120 (TAY)	117 (TAY)	114 (TAC)	118 (TAY)	108 (TAT)	117 (TAY)
Val	169 (GTA)	172 (GTA)	183 (GTA)	222 (GTA)	174 (GTA)	192 (GTW)	185 (GTA)
Stop	13 (TAA)	13 (TAA)	13 (TAA)	13 (TAA)	13 (TAA)	13 (AGA/TAA)	13 (TAA)
Total	3799	3801	3803	3792	3800	3813	3801

Tetrapods: human, Blue whale, opossum, chicken, and frog; teleost: Rainbow trout, carp, and loach. The most commonly used codon for each amino acid is shown in parentheses.

The coelacanth  $O_L$  loop, unlike other fish which have a polypyrimidine tract, but similar to tetrapods, contains a stretch of thymines that is needed for the initiation of L-strand synthesis (WONG and CLAYTON 1985).

**Coding sequences:** The coelacanth *12S* and *16S* rRNA genes are 983 and 1665 nucleotides long, respectively (Table 2). The secondary structure of both rRNA genes appears to be reasonably conserved in respect to those of other vertebrates. Most of our rRNA gene sequence showed only minor differences (99.58% similarity) to that previously reported (HEDGES *et al.* 1993) with the exception of the last 332 bp of the *16S* rRNA gene sequence, which differed by 20%. These 332 bp correspond to the last PCR fragment amplified by HEDGES *et al.* (1993) and a search in GenBank revealed that the sequence of this PCR fragment had 96.7% similarity to that sequence of the alligator *16S* rRNA gene sequence (HEDGES 1994). This and other reported cases of apparently scrambled sequences during data preparation can be recognized and prevented by individually checking PCR amplified fragments for compatibility in recovering congruent phylogenetic trees (see EDWARDS and ARCTANDER, 1997).

All 22 coelacanth tRNA gene sequences can be folded into a canonical cloverleaf secondary structure with the exception of the tRNA<sup>Ser(AGY)</sup> (data not shown). Two mismatched base pairs in the stems of these putative cloverleaf secondary structures were detected on average for each tRNA. The coelacanth tRNAs ranged in size from 65 to 76 nucleotides (Table 2) and showed

size variability in their DHU and T $\psi$ C arms when compared with other vertebrate mitochondrial tRNAs.

The coelacanth mitochondrial genome contains 13 protein-coding genes (Figures 1 and 2); in two cases there is a reading-frame overlap on the same strand (*ATPases 8* and *6* share 10 nucleotides; *ND4L* and *ND4* overlap by seven nucleotides). All coelacanth mitochondrial protein-coding genes begin with a ATG start codon except *COI*, which starts with GTG (Table 2). This initiation codon usage is also found in all other fish mitochondrial genomes that have been completely sequenced so far (TZENG *et al.* 1992; CHANG *et al.* 1994; LEE and KOCHER 1995; ZARDOYA *et al.* 1995; JOHANSEN and BAKKE 1996; NOACK *et al.* 1996; ZARDOYA and MEYER 1996a). Most coelacanth ORFs end with TAA (*ND1*, *ND2*, *COI*, *ATPase 8*, *COIII*, *ND4L* and *ND5*), one ends with AGA (*ND6*), and the rest have incomplete stop codons, either T (*COII*, *ND3*, *ND4*) or TA (*Cyt b*) (Table 2).

The codon usage of the coelacanth is similar to that of lamprey (LEE and KOCHER 1995), bichir (NOACK *et al.* 1996), carp, trout and loach (TZENG *et al.* 1992; CHANG *et al.* 1994; ZARDOYA *et al.* 1995), lungfish (ZARDOYA and MEYER 1996a), and tetrapods (*e.g.*, ROE *et al.* 1985; DESJARDINS and MORAIS 1990; ARNASON and GULLBERG 1993; HORAI *et al.* 1995) (Table 4). A total of 3803 amino acids are encoded by the coelacanth mitochondrial genome. As in other vertebrates, the most abundant amino acid residue encoded by the coelacanth mitochondrial genome is leucine, whereas the

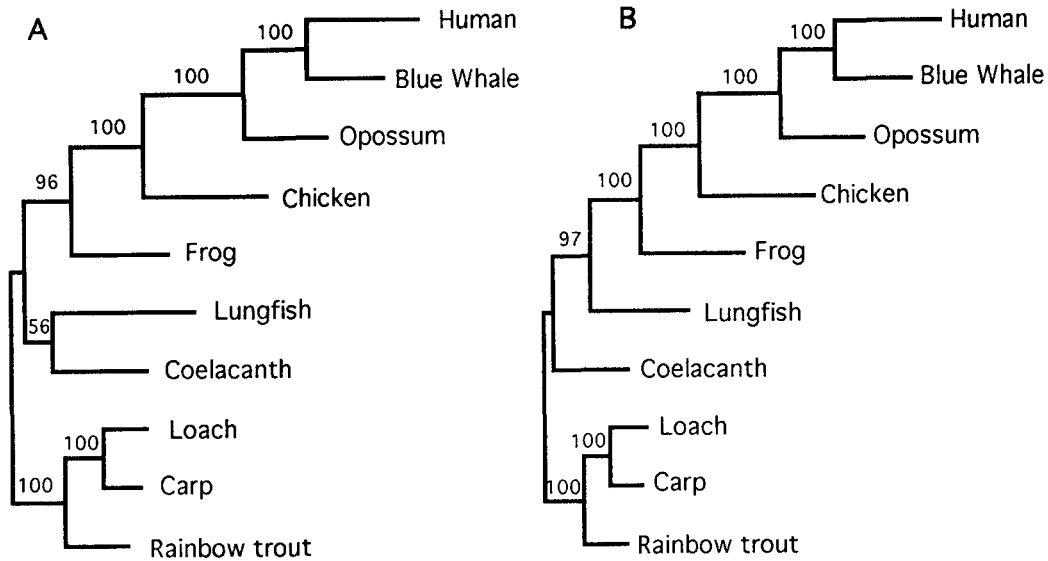


FIGURE 3.—Phylogenetic position of the coelacanth. A data set combining all (protein-coding, rRNA and tRNA) mitochondrial genes was analyzed with maximum parsimony (A), neighbor-joining and maximum likelihood (B) phylogenetic methods. Numbers shown above branches represent bootstrap values from 100 pseudoreplicates for MP (A) and NJ (B). Rainbow trout, carp, and loach were used as outgroup taxa. Third codon positions and transitions in first codon positions of protein coding genes were excluded from the MP phylogenetic analysis. The NJ analysis was performed using Kimura two-parameter distances, an  $\alpha = 0.53$  (gamma distribution), and excluding third codon positions. The ML analysis was performed with a transition: transversion ratio of 2:1, using the F84 model implemented in PHYLIP version 3.55 (FELSENSTEIN 1989), which accounts for different base frequencies, and excluding third codon positions.

rarest is cysteine (Table 4). Adenines and cytosines are preferentially used in third codon positions of the coelacanth mitochondrial protein coding genes, indicating that codon usage is probably influenced by strand-specific base composition mutational bias (FLOOK *et al.* 1995; JERMIIN *et al.* 1994; LEE and KOCHER 1995). Alternatively, third codon position usage may be determined by availability of ribonucleotides in the mitochondria (XIA 1996).

**Phylogenetic position of the coelacanth:** To assess the position of the coelacanth with respect to lungfishes and tetrapods, a combined data set comprising rRNA, tRNA and protein coding nucleotide sequences of the human (HORAI *et al.* 1995), blue whale (ARNASON and GULLBERG 1993), opossum (JANKE *et al.* 1994), chicken (DESJARDINS and MORAIS 1990), frog (ROE *et al.* 1985), carp (CHANG *et al.* 1994), loach (TZENG *et al.* 1992), rainbow trout (ZARDOYA *et al.* 1995), African lungfish (ZARDOYA and MEYER 1996a), and coelacanth mitochondrial genomes was constructed; it is composed of a total of 16,140 characters. This data set was analyzed with the three most commonly used methods of phylogenetic inference, *i.e.*, maximum parsimony (MP; FITCH 1971), neighbor-joining (NJ; SAITOU and NEI 1987), and maximum likelihood (ML; FELSENSTEIN 1989).

When third codon positions and transitions in first codon positions of the protein-coding gene sequences were excluded from the analysis and teleosts (trout, carp and loach) were included as outgroup taxa, MP recovered one most parsimonious tree (8468 steps, C.I. = 0.65) in which a coelacanth/lungfish clade is identi-

fied as the sister group of tetrapods (Figure 3A). This is the same hypothesis as that which is favored by the largest available nuclear DNA data set, the complete *28S rRNA* gene sequences (ZARDOYA and MEYER 1996b). The node grouping the coelacanth/lungfish clade with tetrapods was supported by a 56% bootstrap value (Figure 3A). However, if the same data set was analyzed with MP and transitions in first positions of the protein-coding genes were included in the analysis or a transition:transversion weighting of 2:1 was assumed for the whole data set, a lungfish/tetrapod clade was favored with low or moderate bootstrap support (51 and 68%, respectively).

For the NJ analysis, to account for the variation of substitution rates among sites, the  $\alpha$  shape parameter of the gamma distribution of rate variation was estimated based on the MP tree (Figure 3A) by the method of YANG and KUMAR (1996). A NJ analysis of the same data set, excluding third codon positions of the protein-coding genes, with teleosts as outgroup taxa, using Kimura 2-parameter distances, and an  $\alpha = 0.53$ , arrived at a tree in which lungfishes are placed as the sister group of tetrapods (97% bootstrap value for the lungfish+tetrapod node) (Figure 3B). The same topology was recovered when a ML analysis (F84 model, 2:1 transition:transversion ratio; Ln likelihood = -61,846.08) was performed excluding third codon positions of the protein-coding genes and including teleosts as outgroup taxa. In this case, all branch lengths were found to be significantly greater than zero ( $P < 0.01$ ).

The disagreement between methods of phylogenetic inference was investigated with KISHINO and HASEGAWA (1989) and TEMPLETON (1983) tests. None of the three possible hypotheses, *i.e.*, lungfish as sister group of tetrapods (Ln likelihood = -61,846.08, best ML tree; 10,812 steps, best MP tree), coelacanth as sister group of tetrapods [Ln likelihood = -61,863.43,  $\Delta\text{LnL} = 17.35 \pm 22.77$  (SD); 10,832 steps,  $\Delta\text{steps} = 20 \pm 15.43$ ], or coelacanth as sister group of lungfish (Ln likelihood = -61,859.78,  $\Delta\text{LnL} = 13.70 \pm 22.69$ ; 10,813 steps,  $\Delta\text{steps} = 1 \pm 16.03$ ), could be statistically rejected based on the mitochondrial DNA data set when third codon positions of mitochondrial protein-coding genes were excluded from the analyses and teleosts were used as outgroup taxa. In a separate publication, we will report in more detail on the conflicting phylogenetic signal that is contained in the tRNA, rRNA, and protein mitochondrial data sets and that may be one of the major causes that explain why none of the three hypotheses can be statistically ruled out (R. ZARDOYA, Y. CAO, M. HASEGAWA and A. MEYER, unpublished data).

Our evolutionary analyses confirmed the phylogenetic position of the coelacanth as a closer relative to tetrapods and other sarcopterygians than to the ray-finned fishes (Actinopterygii). These results indicate that the rate of evolution of the mitochondrial genome is appropriate for resolving relationships even among ancient lineages (at least up to the Devonian). However, it seems to fall short of recovering and strongly supporting the relationships among the extant lineage of lobe-finned fishes that originated within a narrow window in time ~400 mya. The rapid origin of the various lobe-finned lineages and their rapid radiation before the origin of tetrapods continues to make it difficult to resolve their relationships with confidence.

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