

Database on the structure of small subunit ribosomal RNA

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ABSTRACT

Over 11 500 complete or nearly complete sequences are now available from the Antwerp database on small subunit ribosomal RNA. All these sequences are aligned with one another on the basis of the adopted secondary structure model, which is corroborated by the observation of compensating substitutions in the alignment. Literature references, accession numbers and taxonomic information are also compiled. The database can be consulted via the World Wide Web at URL <http://rrna.uia.ac.be/ssu/>

CONTENTS OF THE DATABASE

In August 1998, the Antwerp small subunit (SSU) rRNA database contained 3166 eukaryotic, 7336 bacterial, 324 archaeal, 120 plastid and 601 mitochondrial sequences. The database comprises complete or nearly-complete sequences while partial SSU rRNA sequences are included only if the combined length of the sequenced segments amounts to at least 70% of the estimated chain length of the molecule. The chain length of a partially determined sequence is estimated by comparing it to a complete sequence of a presumed close relative. All SSU rRNA sequences are stored in the form of an alignment and contain the postulated secondary structure pattern in encoded form (see the rRNA website at URL <http://rrna.uia.ac.be/> for detailed information).

Table 1 lists the different eukaryotic taxa and the number of representatives in the database for which the SSU rRNA sequence has been determined. The taxonomic classification of the animals is according to Brusca and Brusca (1). For the plants and the Fungi, we have chosen to extend the taxonomic information supplied up to the level of orders, contrary to previous papers describing the contents of the SSU rRNA database. The classification of vascular plants is according to Mabberley (2), while the classification of Bryopsida is according to Crosby and Magill (3). Additional classificatory information for the terrestrial plants was taken from Sitte *et al.* (4) and Farr *et al.* (5). On the rRNA website, taxonomic information regarding the family to which the plant species are classified will also be available. The classification of the 'true' fungi or Eumycota is according to

Hawksworth *et al.* (6), Kurtzman and Fell (7), and de Hoog and Guarro (8). The remaining eukaryotes, viz. the protoctists are classified according to Margulis *et al.* (9). Overall, species are included in the database under the binomial used for the publication of the sequence. We therefore refrained from doing any taxonomic name change, even when obviously needed.

Table 2 covers the prokaryotic SSU rRNA sequences. The classification of prokaryotes is, as before, based on the construction of evolutionary trees. New sequences retrieved from EMBL or GenBank, or from direct submissions, are aligned with their presumed closest relative. Evolutionary trees are then constructed by the neighbor-joining method (10), and according to the phylogenetic position observed, the species are assigned to one of the taxa previously described by Woese and coworkers (11,12) and our research group (13,14).

SECONDARY STRUCTURE AND NUCLEOTIDE VARIABILITY

Our SSU rRNA sequence alignment is based on two different secondary structure models. The first one is the prokaryotic model, which is applicable to Bacteria, Archaea, plastids and mitochondria, while the second one is the eukaryotic model applicable to all Eukaryotes. The two models are slightly different, each containing a number of structural elements specific for the group (see below). The prokaryotic model is essentially identical to those distributed by Gutell (15), but the model followed for eukaryotic SSU rRNAs includes a secondary structure pattern in certain variable areas left undefined in the models of the latter author.

Helices in the SSU rRNA secondary structure model are given a different number if separated by a multibranching loop (e.g., helices 9 and 10), by a pseudoknot loop (e.g., helices 1 and 2), or by a single stranded area that does not form a loop (e.g., helices 2 and 32). A single number is given to 50 universal helices, which are present in all SSU rRNAs from Archaea, Bacteria and plastids known to date. These 50 helices are also present in all known eukaryotic SSU rRNAs except in those of the microsporidians (such as *Vairimorpha*, *Nosema*, and relatives), where some of these helices are missing. Helix 11 is also missing in the

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Table 1. List of eukaryotic taxa represented in the database and number of their representatives (August 1998)

Kingdom Animalia ^a				Kingdom Fungi ^a			
Phylum	Class	Number of sequences ^b		Phylum	Order	Number of sequences ^b	
		N	M			N	M
Placozoa		2		Ascomycota			
Orthonectida		2		Archiascomycetes	Pneumocystidales	3	
Rhombozoa	Dicyemida	3			Protomycetales	4	
Porifera	Calcarea	3			Schizosaccharomycetales	5	2
	Demospongiae	3			Taphrinales	18	
Cnidaria	Anthozoa	12		Euascomycetes	Caliciales	14	
	Cubozoa	1			Chaetothyriales	13	
	Hydrozoa	3			Cyttariales	1	
	Uncertain affiliation	7			Diaporthales	6	1
Ctenophora		2			Dothideales	14	
Platyhelminthes	Cestoda	2			Elaphomycetales	3	
	Trematoda	15			Erysiphales	1	
	Turbellaria	34			Eurotiales	58	24
	Uncertain affiliation	1			Hypocreales	4	
Nemertea	Anopla	1			Lecanorales	9	
	Enopla	1			Leotiales	11	
Rotifera		3			Microascales	13	
Gastrotricha		2			Neoclectales	2	
Kinorhyncha		1			Onygenales	18	1
Nematoda	Adenophorea	27			Ophiostomatales	3	
	Secernentea	60			Ostropales	3	
	Uncertain affiliation	10			Peltigerales	2	
Nematomorpha	Gordioida	5			Pezizales	68	
Priapulida		2			Phyllachorales	1	1
Entoprocta		3			Pleosporales	23	
Gnathostomulida		1			Rhizymatales	1	
Acanthocephala	Archiacanthocephala	3			Sordariales	8	1
	Eoacanthocephala	2			Xylariales	2	
	Palaeacanthocephala	7			Uncertain affiliation	23	
Annelida	Hirudinida	6		Hemiascomycetes	Saccharomycetales	120	7
	Oligochaeta	8	1		Uncertain affiliation	1	
	Polychaeta	16		Uncertain affiliation		1	
Sipuncula	Phascolosomida	1		Basidiomycota			
Echiura		1		Heterobasidiomycetes	Agaricostilbales	1	
Pogonophora		1			Atractiellales	1	
Vestimentifera	Basibranchia	1			Auriculariales	3	
Arthropoda	Branchiopoda	5			Filobasidiales	10	
	Chelicerata	50	1		Heterogastridiales	1	
	Insecta	152	23		Tremellales	30	
	Malacostraca	17	2	Homobasidiomycetes	Agaricales	17	1
	Maxillopoda	12			Aphylophorales	81	
	Myriapoda	1			Boletales	1	
Onychophora		1			Bondarzewiales	3	
Tardigrada		5			Cantharellales	9	
Pentastomida	Pentastomata	1			Ceratobasidiales	18	
Mollusca	Caudofoveata	1			Cortinariales	1	
	Bivalvia	34			Dacrymycetales	4	
	Gastropoda	43			Fistulinales	2	
	Polyplacophora	3			Gomphales	2	
	Scaphopoda	1			Hericiales	9	
Phoronida		2			Lycoperidiales	3	
Ectoprocta	Gymnolaemata	1			Nidulariales	2	
	Phylactolaemata	2			Phallales	1	
Brachiopoda	Articulata	8			Schizophyllales	1	
	Inarticulata	7			Sclerodermatales	2	
Echinodermata	Asteroida	4	1		Tulostomatales	1	
	Crinoidea	2	1	Teliomycetes	Uredinales	4	
	Echinoidea	24	4	Ustomycetes	Graphiiales	2	
	Holothuroidea	4	1		Platyglloeales	6	
	Ophiuroidea	4			Sporidiales	31	
Chaetognatha		3			Ustilaginales	10	
Hemichordata	Enteropneusta	2	1		Uncertain affiliation	1	
Chordata	Agnatha	4	1	Uncertain affiliation		13	
	Amphibia	18	62	Zygomycota			
	Aves	3	88	Trichomycetes	Harpellales	5	
	Chondrichthyes	4		Zygomycetes	Entomophthorales	3	
	Mammalia	11	277		Glomales	28	
	Osteichthyes	16	32		Kickxellales	9	
	Reptilia	4	21		Mucorales	5	
	Cephalochordata (Sub.)	1			Uncertain affiliation	1	
	Urochordata (Subphyl.)	4		Uncertain affiliation		1	
Total:		706	516	Total:		775	38

trichomonads and relatives. Helices specific to the eukaryotic model are numbered Ea-b, where a is the number of the preceding universal helix and b sequentially numbers all helices inserted between universal helices a and a+1. Helices specific to the prokaryotic model are similarly given composite numbers of the form Pa-b. Mitochondrial sequences show extreme variability in

length and in the number of helices present. Figure 1 shows the secondary structure model of the plastid SSU rRNA nucleotide sequence of *Zea mays*.

Examples of secondary structure models for eukaryotic and mitochondrial SSU rRNAs have been given in previous papers on our database (13,16,17). Color maps showing the distribution of

Table 1. Continued

Kingdom Plantae ^a				Kingdom Protocista ^a					
Phylum Class	Order	Number of sequences ^b			Phylum	Class	Number of sequences ^b		
		N	M	P			N	M	P
Bryophyta					Actinopoda	Heliozoa	1		
Anthocerotopsida		7			Apicomplexa	Coccidia	52		1
Bryopsida		31		1		Hematozoa	73		3
Junggermanniopsida		12				Uncertain affiliation	2		
Marchantiopsida		11	1	1	Bacillariophyta	Bacillariophyceae	10		3
Pteridophyta						Coscinopiscophyceae	23		
Equisetopsida		2		1	Chlorarachnida		21	1	3
Filicopsida		19		8	Chlorophyta	Charophyceae	30		3
Lycopsida		9		3		Chlorophyceae	167	6	19
Psilotopsida		3		2		Prasinophyceae	7		1
Pinophyta						Ulvoephyceae	39		
Cycadopsida		2				Uncertain affiliation	4		
Ginkgoopsida		1			Chrysophyta	Chrysophyceae	26		2
Gnetopsida		10		1		Dictyochophyceae	3		
Pinopsida		43		2		Uncertain affiliation	9		
Magnoliophyta					Chytridiomycota		9	2	
Liliopsida	Alismatales	2			Ciliophora	Colpodea	2		
	Arales	2				Karyorelictea	4		
	Arecales	2				Litostomatea	16		
	Bromeliales	2				Nassophorea	5	3	
	Commelinales	1				Oligohymenophorea	31	2	
	Cyperales	4	3	3		Phylopharyngea	2		
	Liliales	13				Prostomatea	1		
	Orchidales	8				Spirotrichea	13		
	Poales		2		Conjugaphyta	Conjugatophyceae	9		1
	Typhales	1			Cryptophyta		30		4
	Zingiberales	7			Dictyostelida		1	1	
Magnoliopsida	Apiales	10		1	Dinoflagellata		50		
	Aristolochiales	4			Euglenida		4		7
	Asterales	2			Eustigmatophyta	Eustigmatophyceae	5		
	Batidales	1			Glaucocystophyta	Glaucocystophyceae	1		5
	Campanulales	2			Granuloreticulosa		1		
	Capparidales	6	1		Haplosporida		7		
	Caryophyllales	5			Hyphochytridiomycota		1		
	Casuarinales	1			Labyrinthulomycota		6		
	Celastrales	1			Microspora		60		
	Cornales	10			Myxozoa	Myxosporea	20		
	Daphniphyllales	1			Oomycota		4		
	Dilleniales	4			Phaeophyta		11	1	1
	Dipsacales	3			Plasmodial Slime Molds	Myxomycota	1	2	
	Ebenales	4			Plasmodiophoromycota		1		
	Ericales	6			Prymnesiophyta		17		5
	Eucommiales	1			Rhaphidophyta		4		
	Euphorbiales	7		3	Rhizopoda	Filosea	1		
	Fabales	4	2	4		Lobosea	72	2	
	Fagales	2		1		Uncertain affiliation	1		
	Gentianales	1			Rhodophyta		178	2	6
	Geraniales	4			Xanthophyta		5		
	Haloragidales	3			Zoomastigina	Amebomastigota	6		
	Hamamelidiales	6				Choanomastigotes	2		
	Illiciales	2				Diplomonadida	13		
	Lamiales	3				Kinetoplastida	56	7	
	Laurales	3				Parabasalia	20		
	Linales	1				Proteromonadida	1		
	Magnoliales	9				Zoomastigophora	2		
	Malvales	5			Stramenopiles	Uncertain affiliation	7		
	Myrtales	1	1		Uncertain affiliation		9	1	7
	Nepenthales	3							
	Nymphaeales	6							
	Papaverales	3							
	Piperales	7							
	Plumbaginales	1							
	Polygalales	3							
	Polygonales	1							
	Proteales	4							
	Rafflesiales	5	3	5					
	Ranunculales	21							
	Rhamnales	1							
	Rosales	44							
	Rubiales	1							
	Santalales	72	3	2					
	Sapindales	4							
	Scophulariales	19		3					
	Solanales	6	1	7					
	Theales	2							
	Trochodendrales	2							
	Urticales	4							
	Violales	16		1					
Total:		529	17	49			1156	30	71

^aThe Metazoan taxa are listed in the same order as they appear in ref. 1.

^bThe number of sequences listed in the database is larger than the number of species, because for certain species multiple SSU rRNA sequences have been determined, usually by different authors. The sequences are not necessarily identical because they may have been determined for different varieties or strains of a species, or for different genes of the same organism. The number is listed for sequences of nuclear (N), mitochondrial (M) and plastid (P) origin.

^cExcept in the case of the plant phyla, taxa are ordered alphabetically.

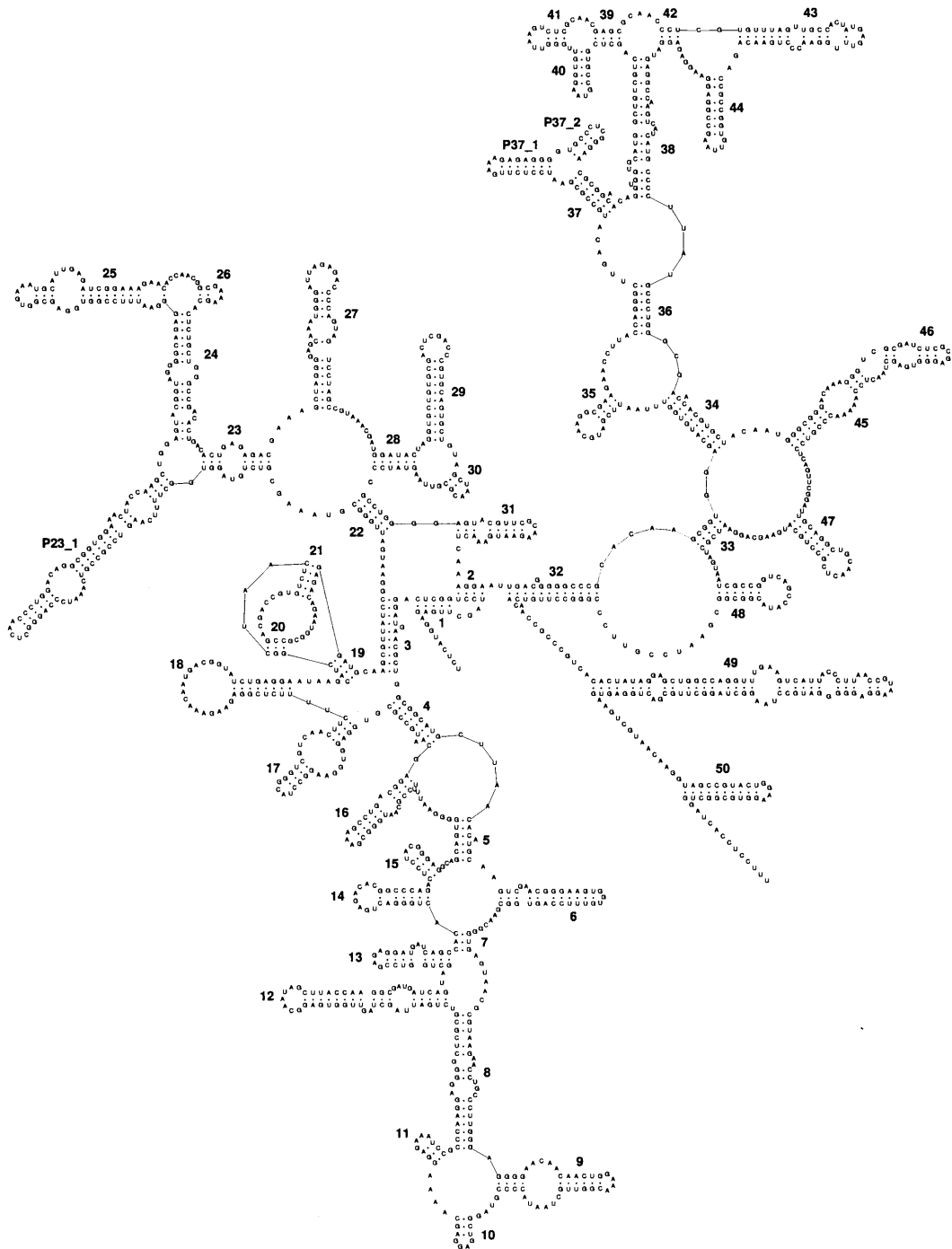


Figure 1. Secondary structure model for the plastid SSU rRNA of *Zea mays*. The sequence is written clockwise from 5' to 3' terminus.

conserved and variable sites in bacterial and eukaryotic SSU rRNAs (18,19) can be consulted via the Internet at URL <http://bioc-www.uia.ac.be/uyvdp/>

AVAILABILITY OF THE DATA

Each SSU rRNA sequence is stored in a separate file in order to simplify access to the data. Each of these files contains primary and secondary structure information, as well as annotations such

as accession number, literature reference, and detailed taxonomic specifications. The SSU rRNA database is made available via the World Wide Web at URL <http://rna.uia.ac.be/ssu/>. Through the WWW, it is very easy to select sequences either one by one, or by taxonomic group, or by a combination of both. Sequences can be retrieved in different formats. On-line information about the database is also available.

Table 2. List of prokaryotic taxa represented in the database and number of their representatives (August 1998)

Bacteria		
Division		Number of sequences ^a
Chlamydiae		86
Cyanobacteria		107
Fibrobacter		18
Flavobacteria and relatives		442
Fusobacterium and relatives		32
Gram Positives and relatives, Low G+C		1645
Gram Positives and relatives, High G+C		1308
Green Sulfur		29
Green non sulfur		25
Planctomyces and relatives		60
Proteobacteria α		877
Proteobacteria β		385
Proteobacteria γ		1417
Proteobacteria δ		181
Proteobacteria ϵ		167
Proteobacteria, uncertain affiliation		9
Radioresistant micrococci and relatives		51
Spirochetes		279
Thermophilic oxygen reducers		8
Thermotogales		18
Uncertain affiliation ^b		192
Total:		7336

Archaea		
Division	Subdivision	Number of sequences ^a
Crenarchaeota		93
Euryarchaeota	Archaeoglobales	4
	Halobacteria	70
	Methanobacteriales	34
	Methanococcales	19
	Methanomicrobium group	63
	Methanopyrales	1
	Thermococcales	29
	Thermoplasma	2
	Uncertain affiliation	9
Total:		324

^aThe number of sequences listed in the database is larger than the number of species (cf. Table 1).

^bIn some cases, it cannot be decided to which taxonomic group a species should be ascribed, since the clustering of its SSU rRNA sequence is unstable and depends on the tree construction method used and on the set of sequences included in the analysis.

If problems occur in connecting to the server or in retrieving data, the authors can be contacted by electronic mail to yvdp@uia.ua.ac.be or dwachter@uia.ua.ac.be. Users publishing results based on data retrieved from our database are requested to cite this paper.

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