RPG: the Ribosomal Protein Gene database

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ABSTRACT

RPG (http://ribosome.miyazaki-med.ac.jp/) is a new database that provides detailed information about ribosomal protein (RP) genes. It contains data from humans and other organisms, including Drosophila melanogaster, Caenorhabditis elegans, Saccharomyces cerevisiae, Methanococcus jannaschii and Escherichia coli. Users can search the database by gene name and organism. Each record includes sequences (genomic, cDNA and amino acid sequences), intron/exon structures, genomic locations and information about orthologs. In addition, users can view and compare the gene structures of the above organisms and make multiple amino acid sequence alignments. RPG also provides information on small nucleolar RNAs (snoRNAs) that are encoded in the introns of RP genes.

INTRODUCTION

The ribosome is a universal and essential catalyst of protein synthesis in all organisms. Because of the fundamental role played by the ribosome, its structure and function have been significantly conserved during evolution. In eukaryotes, the ribosome is composed of four RNA molecules (rRNAs) and about 80 different proteins (RPs) (1), which are each present as a single copy with the exception of two proteins. The genes encoding rRNAs are clustered at a few sites in the eukaryotic genome, whereas the genes encoding RPs are widely dispersed (2). rRNA sequences have been extensively studied and data from thousands of organisms are now available (The Ribosomal Database Project; http://rdp.cme.msu.edu/) (3). On the other hand, little has been done for RP genes despite the acute need for a dedicated database.

In mammalian cells, each RP is encoded by a single gene but this gene generates a large number (10–20 copies) of silent, processed pseudogenes (4). This has hampered the mapping and sequence analysis of the functional RP genes. Through persistent efforts, however, we have completely mapped and sequenced most of these functional genes in the human genome (2,5,6). This has enabled us to compare their gene structures with those of other eukaryotes whose genome sequences have already been sequenced. Here we present a new database containing information about RP genes from various species, which we hope will constitute a valuable resource for biomedical research and a powerful tool for comparative studies of gene evolution.

DATA SOURCES AND FORMATS

Most of the human RP gene sequences in the database were determined in our previous study (6). Others were collected from the DDBJ/EMBL/GenBank databases. Sequences from other eukaryotes (Drosophila melanogaster, Caenorhabditis elegans and Saccharomyces cerevisiae) were obtained primarily by a BLAST search of the following databases using the human cDNA sequences as queries: FlyBase at http://flybase.org/ (7), WormBase at http://www.wormbase.org/ (8) and Saccharomyces Genome Database (SGD) at http://www.yeastgenome.org/ (9), respectively. The archaea (Methanococcus jannaschii) RP gene sequences were collected from the NCBI Entrez FTP site, and the bacterial (Escherichia coli) RP gene sequences were from GenoBase at http://ecoli.aist-nara.ac.jp/. In addition to these sequence data, we have integrated different types of information about RP genes, including chromosomal positions, accession numbers, gene and CDS sizes, orthologs, snoRNAs and links to other public databases. These data were automatically written in RPG unique file format by using a MS-Excel VBA script. Graphical data showing intron/exon structures, translation start and stop sites, and snoRNA gene locations were also formatted by using the VBA script and integrated into the database. Users can access these files through a web site running a CGI program. We employed Clustal W 1.82 to align amino acid sequences among orthologs (10). The Clustal W outputs were formatted into HTML files after color shading of the sequences with a Perl script to show amino acid similarities. The number of current entries from various organisms is summarized in Table 1.

STRUCTURE AND INTERFACE

The data in the RPG database can be accessed in a variety of ways. Each entry can be searched by gene name or organism (Fig. 1A). For human RP genes, each entry is linked to a gene table or a chromosomal map position (5). For other genes, each entry is linked to an orthologous gene classification table, which includes all of the RP gene entries in this database. The main page for each gene provides the following information (Fig. 1B); (i) a schematic view of intron/exon structure, the translation start and stop sites, and positions of snoRNA genes if available, (ii) general information including the source organism, the gene name, the chromosomal localization, the
accession number, the gene and CDS length, and the number
of exons, (iii) a schematic view of human chromosomes with
an indication of map position that links to NCBI Entrez
MapView, (iv) links to orthologous gene entries, comparative
views of orthologous gene structures, and protein sequence
alignments, (v) links to snoRNA gene entries, and (vi) links to
other public databases, e.g., GenBank, LocusLink, and NCBI
Entrez Mapview. For snoRNA genes, especially, the entry
page provides sequence, information about the host gene, the
accession number, the rRNA modification site, and the
modification type, for example ‘Box C/D 2'-O-methylation’
or ‘Box H/ACA pseudouridylation’ (Fig. 1D).

Table 1. Number of current entries from six organisms

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FUTURE DEVELOPMENT
RPG will continue to expand by collecting additional data sets
from other eukaryotes, including mouse, Fugu, Arabidopsis
and Schizosaccharomyces. In addition to current cytoplasmic
RPs, we will also include data from organelles, namely,
mitochondria and chloroplasts, as these organelles possess
their own ribosomes which have evolved from those of ancient
bacteria.

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