Cloned Ribosomal Ribonucleic Acid Genes from *Pseudomonas aeruginosa* as Probes for Conserved Deoxyribonucleic Acid Sequences

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Ribosomal ribonucleic acid (rRNA) genes were isolated from a *Pst*I digest of *Pseudomonas aeruginosa* chromosomal deoxyribonucleic acid (DNA), cloned in *Escherichia coli*, and used as probes for conserved gene sequences. Recombinant plasmid pHF1 contained an 8,800-base pair insertion containing 5S, 16S, and 23S rRNA genes. We constructed subclones of pHF1 containing parts of the 16S and 23S rRNA genes (pHF1.1) and parts of the 23S and 5S rRNA genes (pHF1.2). DNA-DNA hybridization experiments in which we used filter-bound chromosomal DNA from various bacteria and 35S-labeled plasmid rRNA genes (rDNA) indicated that the homology values reflected the actual phylogenetic distances to *P. aeruginosa*. Compared with oligonucleotide sequence analysis of 16S rRNA, a good correlation was found between DNA-rDNA homology values and SAB (similarity coefficient of 16S rRNAs) values above 0.4. The use of rDNA instead of rRNA in hybridization experiments offers several advantages; e.g., rDNA can easily be labeled in vitro, and the degree of relatedness can be expressed in terms of percent homology and does not have to be determined by laborious measurement of thermal stability, as in the case of rRNA.

Deoxyribonucleic acid (DNA)-DNA hybridization is a powerful tool for elucidation of close relationships among bacteria. However, it falls above the intragenic level. Lower levels of relationships can be detected by comparing the primary structures of ribosomal ribonucleic acids (rRNA) either by sequencing or by hybridization. Comparative sequence analysis of 16S rRNA has proved to be the best approach to determine lower levels of relationships (20). Moreover, this approach requires the use of specialized techniques which are not readily applicable to large numbers of strains. Determination of rRNA cistron similarities by hybridization is certainly not as accurate as the sequencing approach, but it is a simpler and more rapid technique. The similarities of rRNAs have usually been measured by the differences between the thermal stabilities of the homologous and heterologous DNA-rRNA duplexes (2). The percentage of RNA homology to DNA is not a reliable measure of rRNA homology, because the amount of RNA bound does not depend only on the actual rRNA homology (3). Therefore, it is not possible to express the degree of relatedness in terms of percent homology, and one has to use the more laborious measurement of thermal stability. The similarities of rRNAs have also been measured by adding competitor rRNA to the homologous hybridization reaction (15). By using this method, it is possible to determine competition homology values and to avoid determination of the thermal stability of the hybrids. The disadvantage of the latter method are the large amounts of unlabeled rRNA that are needed from the competitor strain. However, isolation of pure rRNA is one of the major difficulties of the DNA-rRNA hybridization method (16). Therefore, we decided to clone rRNA cistrons and to use the rRNA genes (rDNA) as a probe for studying distant phylogenetic relationships.

MATERIALS AND METHODS

Organisms and growth conditions. Most of the strains which we used are listed in Table 1; also used were *Escherichia coli* RR28 and vector pHE3, which were received as gifts from H. Hennecke (ETH, Zürich, Switzerland). Pseudomonads and *E. coli* were cultivated aerobically in 0.5% peptone-0.3% yeast extract-bouillon (pH 7.0), and staphylococci, micrococci, and bacilli were cultivated in 1% peptone-0.5% yeast extract-0.5% glucose-0.8% sodium chloride-bouillon (pH 7.2). Streptococci were grown without aeration in CASO bouillon (E. Merck AG, Darmstadt, Federal Republic of Germany). The incubation temperature was 30°C for pseudomonads and micrococci and 35°C for streptococci, staphylococci, bacilli, and *E. coli*. Cells were harvested in the late logarithmic growth phase.

Isolation of rRNA and DNA. rRNAs were isolated and purified as described elsewhere (18). Extraction of chromosomal DNA was carried out by using the method of Marmur (12), with the modifications described by Meyer and Schleifer (13, 14). Plasmid DNA was isolated by the method of Clewell and Helinski (1) or by the rapid preparation procedure described by Klein et al. (11).

Labeling of rRNA and DNA. The rRNAs were partially hydrolyzed by mild alkali treatment in 500 mM sodium borate buffer (pH 9.0) at 70°C for 35 min, using the method of Engel and Davidson (5). The resulting fragments were 5'-labeled by using [γ-32P]adenosine triphosphate (New England Nuclear Corp., Dréeich, Federal Republic of Germany) and T4 polynucleotide kinase (Bethesda Research Laboratories, Neu Isenburg, Federal Republic of Germany). The conditions used were those described by Stackebrandt et al. (18, 19). Excess [γ-32P]adenosine triphosphate was removed by thin-layer chromatography on diethylamino-ethyl-cellulose glass plates (Macherey and Nagel, Düren, Federal Republic of Germany) which were developed with 0.45 M ammonium formate-0.1 M ethylenediaminetetraacetate in 9 M urea at 60°C.

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Plasmid DNA and chromosomal DNA were labeled by nick translation, using [α-35S]thiodoxygenadenosine triphosphate (New England Nuclear Corp.) and a reaction kit from Bethesda Research Laboratories. The conditions used were those recommended by the manufacturer. The labeled DNA was separated from free [α-35S]thiodoxygenadenosine triphosphate by filtration through Elutip (Schleicher & Schuell, Dassel, Federal Republic of Germany).

**Molecular cloning of rDNA.** Chromosomal DNA from *Pseudomonas aeruginosa* was digested with restriction enzyme *PstI* (Bethesda Research Laboratories). The resulting fragments were analyzed by Southern hybridization (17) to 32P-labeled homologous 5S, 16S, and 23S rRNAs. Fragments which hybridized to all three rRNAs were recovered after agarose gel electrophoresis (0.7% agarose in 40 mM tris(hydroxymethyl)aminomethane-1mM ethylenediaminetetraacetate-5mM sodium acetate buffer, pH 8.3) and ligated with *PstI*-digested *E. coli* vector pHE3 (8) by using T4 ligase (Bethesda Research Laboratories). After transformation of *E. coli* RR28, positive clones were detected by Southern hybridization of filter-bound plasmid DNA to 32P-labeled homologous 23S rRNA. Subclones were constructed by ligation of digested (*BamHI* or *BstEII* or both) vector pHE3 with restriction fragments (*BamHI* or *BstEII* or both) of the initial recombinant plasmid.

**Hybridization.** Southern hybridizations of filter-bound restriction fragments to 32P-labeled rRNAs were carried out in 3× SSC (0.45 M NaCl plus 0.045 M sodium citrate, pH 7.0) containing 25% formamide at 50°C for 16 h. Fixation of unlabeled DNA on nitrocellulose filters (Schleicher & Schuell, Dassel, Federal Republic of Germany). Molecular cloning of rDNA. Chromosomal DNA from *Pseudomonas aeruginosa* was digested with restriction enzyme *PstI* (Bethesda Research Laboratories). The resulting fragments were analyzed by Southern hybridization (17) to 32P-labeled homologous 5S, 16S, and 23S rRNAs. Fragments which hybridized to all three rRNAs were recovered after agarose gel electrophoresis (0.7% agarose in 40 mM tris(hydroxymethyl)aminomethane-1mM ethylenediaminetetraacetate-5mM sodium acetate buffer, pH 8.3) and ligated with *PstI*-digested *E. coli* vector pHE3 (8) by using T4 ligase (Bethesda Research Laboratories). After transformation of *E. coli* RR28, positive clones were detected by Southern hybridization of filter-bound plasmid DNA to 32P-labeled homologous 23S rRNA. Subclones were constructed by ligation of digested (*BamHI* or *BstEII* or both) vector pHE3 with restriction fragments (*BamHI* or *BstEII* or both) of the initial recombinant plasmid.

**Hybridization.** Southern hybridizations of filter-bound restriction fragments to 32P-labeled rRNAs were carried out in 3× SSC (0.45 M NaCl plus 0.045 M sodium citrate, pH 7.0) containing 25% formamide at 50°C for 16 h. Fixation of unlabeled DNA on nitrocellulose filters (Schleicher & Schuell, Dassel, Federal Republic of Germany).

**RESULTS**

**Cloning of rRNA genes.** The *PstI* digest of *P. aeruginosa* chromosomal DNA was fractionated on 0.7% agarose gels (Fig. 1). Southern hybridizations of the fragments were done to 32P-labeled 5S, 16S, and 23S homologous rRNAs. The autoradiograph in Fig. 1 shows multiple bands. Fragments of about 8,800 base pairs which hybridized to all three rRNAs were isolated and ligated with *PstI*-digested *E. coli* vector pH3. The recombinant vectors were amplified in *E. coli* RR28. Positive clones were detected by Southern hybridization of filter-bound plasmid DNA to homologous 32P-labeled 23S rRNA. One of the recombinant plasmids, which was used in the studies described below, contained an 8,800-base pair insertion and was designated pHF1. This plasmid was characterized by restriction analysis (Fig. 2). The orientation of the rDNA was deduced by selective hybridization of 32P-labeled 5S, 16S, and 23S rRNAs to filter-bound *BstEII*-generated fragments of pHF1. Restriction fragments resulting from digestion of pHF1 by *BstEII* or by *BstEII* and *BamHI* were subcloned in vector pH3. Purified subclone pHF1.1 contained a 2,200-base pair insertion which covered parts of the 16S and 23S rRNAs (Fig. 2), whereas subclone pHF1.2 contained a 1,700-base pair fragment which included the 5S rDNA and part of the 23S rDNA (Fig. 2). The presence of transfer ribonucleic acid genes was not tested in this assay.

**Hybridization experiments.** Various DNA-DNA hybridization experiments with rDNA and chromosomal DNAs were carried out by using the filter technique to show the usefulness of cloned rDNA for phylogenetic studies. 35S-labeled cloned rDNA from *P. aeruginosa* was hybridized to filter-bound chromosomal DNAs of organisms representing different phylogenetic distances (Table 1). The phylogenetic relationships of the organisms used had been elucidated previously by 16S rRNA cataloging (21) and by DNA-rRNA cistron homology studies (4, 15). The pH3 vector alone showed no homology with the chromosomal DNAs. This was demonstrated by hybridization experiments with 35S-labeled pH3 DNA and unlabeled insertion rDNA. Less than 1% of the radioactivity was bound to chromosomal DNAs, indicating that competition between vector DNA and rDNA did not occur. Table 1 summarizes the data obtained from various hybridization experiments. 35S-labeled DNAs of plasmids pHF1, pHF1.1, and pHF1.2 were used, as well as labeled insertion rDNA containing parts of 16S and 23S rRNA genes. The latter rDNA was excised with *BstEII* from plasmid pHF1.1. Both rDNA binding values and DNA-rDNA homology values are listed in Table 1. The homology values were determined by normalizing the percentage of radioactivity bound in the heterologous reaction to the percentage of radioactivity bound in the homologous reaction.

**FIG. 1.** Gel electrophoresis (0.7% agarose) of *P. aeruginosa* DNA digested with restriction endonuclease *PstI*. Lane a was stained with ethidium bromide and visualized under ultraviolet light. Lane b was a Southern transfer analysis in which [γ-32P]adenosine triphosphate-labeled 23S rRNA from *P. aeruginosa* was used as the radioactive probe. Hybridization was carried out as described in the text.
action. Radioactively labeled probes of plasmid pHFl were hybridized with filter-bound chromosomal DNA from various bacteria under optimal and stringent conditions. The DNA homology values of the closely related pseudomonads were similar in both experiments. However, the less related pseudomonads and the gram-positive bacteria showed significantly lower homology values under stringent conditions. Plasmids pHFl.2 and pHFl.1 and the insertion DNA were hybridized under optimal conditions with filter-bound chromosomal DNAs. pHFl.1, which contained parts of the 16S and 23S rRNA genes, was more specific than pHFl.2, which contained 5S rRNA and parts of the 23S rRNA. The border line of resolution seemed to be Pseudomonas cepacia, whereas in most cases Pseudomonas diminuta yielded homology values which were not significantly different from those of gram-positive bacteria. Separation of P. diminuta from gram-positive bacteria could be obtained only with plasmid pHFl.1 and its insertion rDNA.

Our data indicate that the homology values deduced from the levels of rDNA binding reflect actual phylogenetic relatedness. This is in contrast to DNA-rRNA hybridization studies, in which the thermal stability of the DNA-rRNA hybrids and not the level of rRNA binding is a reliable measure for determining relationships (Table 2).

A graphic representation of the DNA-rDNA homology values is shown in Fig. 3, where they are compared with the phylogenetic relationships deduced from 16S rRNA cataloging data (21).

FIG. 2. Restriction endonuclease map of plasmids derived from E. coli vector pH3. The thick solid line indicates vector DNA. Plasmid pHFl contains the coding region for 5S, 23S, and 16S rRNAs from P. aeruginosa. Plasmids pHFl.1 and pHFl.2 are derived from pHFl. The insertion of plasmid pHFl.1 consists of the 2.2-kilobase pair BstEII fragment of pHFl containing parts of the 23S and 16S rRNA genes. The insertion of plasmid pHFl.2 consists of the 1.7-kilobase pair BamHI-BstEII fragment of pHFl containing 5S rRNA genes and part of the 23S rRNA genes. kb, kilobase.

DISCUSSION

The conserved character, the ubiquitous distribution, the functional constancy, and the genetic stability of rRNAs make them ideal markers to measure distant phylogenetic relationships (6, 16). The most useful method for determining phylogenetic relationships among bacteria is presently comparative sequence analysis of 16S rRNAs. However, this is a rather sophisticated, expensive, and laborious technique which is not suitable for routine studies of large groups of strains. Determination of RNA rRNA cistron similarity via hybridization is certainly less accurate and more restricted than the sequencing approach. The hybridization method is not useful for detecting deep branchings in the bacterial kingdom, but it can be used to determine phylogenetic relationships among related genera. Independent studies by Schleifer and Stackebrandt (6) and Johnson and Harich (9) have shown that the correlation between rRNA cistron similarity studies and 16S rRNA oligonucleotide analysis is very good for SAH (similarity coefficient of 16S rRNAs) values above 0.45.

DNA-rRNA hybridization studies are not as easily and as reliably performed as DNA-DNA hybridization studies. The main disadvantages have been discussed previously (16) (see above). Since DNA-DNA hybridization studies are easier to perform and also are more accurate than comparisons of rRNA hybrids, we cloned rRNA cistrons and used the cloned rDNA as a probe for conserved gene sequences.
Radioactively labeled plasmids containing rDNA fragments from *P. aeruginosa* were used for hybridization experiments with filter-bound chromosomal DNAs from various bacteria. *P. aeruginosa* was chosen for this investigation because pseudomonads are the best-studied group of organisms with regard to comparative rRNA analyses (4, 15, 21).

To establish the usefulness of the method, DNAs from bacteria belonging to different phylogenetic groups were used. The DNA-rDNA hybridization values resulted in a dendrogram of relatedness very similar to the results obtained from competition (15) or thermal stability experiments (4). Compared with oligonucleotide cataloging of 16S rRNA, a good correlation was found with *SAB* values above 0.4 (Fig. 3). One of the main advantages of the DNA-rDNA hybridization technique was that, at least with regard to the organisms studied, the measurement of DNA binding homology was sufficient to determine phylogenetic relationships (Tables 1 and 2).

**TABLE 2.** Comparison of DNA-rDNA and DNA-rRNA hybridization studies

<table>
<thead>
<tr>
<th>Source of filter-bound DNA</th>
<th><em>P. aeruginosa</em></th>
<th><em>P. fluorescens</em></th>
<th><em>P. putida</em></th>
<th><em>P. alcaligenes</em></th>
<th><em>P. maltophilia</em></th>
<th><em>P. cepacia</em></th>
<th><em>P. acidovorans</em></th>
<th><em>P. diminuta</em></th>
<th><em>B. megaterium</em></th>
<th><em>B. subtilis</em></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T&lt;sub&gt;max&lt;/sub&gt; (°C)</strong></td>
<td>81.0</td>
<td>76.5</td>
<td>77.5</td>
<td>76.5</td>
<td>67.5</td>
<td>64.5</td>
<td>61.0</td>
<td>61.0</td>
<td>56.0</td>
<td>56.0</td>
</tr>
<tr>
<td>% rRNA binding from <em>P. fluorescens</em></td>
<td>0.14</td>
<td>0.12</td>
<td>0.16</td>
<td>0.11</td>
<td>0.09</td>
<td>0.07</td>
<td>0.07</td>
<td>0.05</td>
<td>0.11</td>
<td>0.10</td>
</tr>
<tr>
<td>% rRNA binding from <em>P. aeruginosa</em></td>
<td>0.24</td>
<td>0.13</td>
<td>0.13</td>
<td>0.12</td>
<td>0.08</td>
<td>0.06</td>
<td>0.06</td>
<td>0.05</td>
<td>0.047</td>
<td>0.028</td>
</tr>
</tbody>
</table>

* Data from reference 4.  
* <sup>T<sub>max</sub></sup> Temperature at which 50% of the bound RNA or DNA was eluted from the filters.  
* Plasmid pHF1 was used as the source of rDNA. Hybridization was carried out under optimal conditions.
above about 0.4 (Fig. 3). Thus, cloning of rRNAs from a limited number of eubacteria would be sufficient to use them as probes for the major lines of descent. Further studies are planned to check whether the results obtained with cloned rDNA of *P. aeruginosa* can be confirmed with cloned rDNAs of other bacteria. Experiments with cloned rDNAs of *Bacillus subtilis* and *Micrococcus luteus* are in progress.

The use of rDNA instead of rRNA also offers additional advantages. rRNA is usually labeled in vivo, whereas DNA can also be easily labeled in vitro by the nick-translation method. The cloned rDNA fragments can be reduced in size, and fragments of different conserved characters can be isolated and recloned. These fragments may be used as probes for determining even lower degrees of phylogenetic relatedness within procaryotes.

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LITERATURE CITED


